1	Predicting the deleterious effects of mutation load
2	in fragmented populations
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ABSTRACT

23 Human-induced habitat fragmentation constitutes a major threat to biodiversity. Both genetic 24 and demographic factors combine to drive small and isolated populations into extinction 25 vortices. However, the deleterious effects of inbreeding and drift load may depend on 26 population structure, migration patterns and mating systems, and are difficult to predict in the 27 absence of crossing experiments. We performed stochastic individual-based simulations 28 aimed at predicting the effects of deleterious mutations on population fitness (offspring 29 viability and median time to extinction) under a variety of settings (landscape configurations, 30 migration models, and mating systems) on the basis of easy-to-collect demographic and 31 genetic information. Pooling all simulations, a large part (70%) of variance in offspring 32 viability was explained by a combination of genetic structure (F_{ST}) and within-deme 33 heterozygosity (H_S). A similar part of variance in median time to extinction was explained by 34 a combination of local population size (N) and heterozygosity (H_S). In both cases, the 35 predictive power increased above 80% when information on mating systems was available. 36 These results provide robust predictive models to evaluate the viability prospects of fragmented populations. 37

INTRODUCTION

40 Human-induced habitat fragmentation constitutes a major threat for biodiversity (Frankham 41 1995). Consequences are, at first, demographic. Small and isolated populations suffer from 42 increased stochasticity and limited rescue effects, which may suffice to cause local extinctions 43 (Lande 1993; Hanski & Ovaskainen 2000). But fragmentation also has genetic consequences, 44 which are likely to contribute significantly to extinction risks. Increased genetic drift reduces 45 the effectiveness of selection against deleterious mutations (Kimura et al. 1963), leading to their progressive accumulation (e.g., Lynch et al. 1995), and decreases both the standing 46 47 genetic variation and the rate of fixation of beneficial mutations (Whitlock 2003), limiting the 48 evolutionary potential of isolated populations. Although the importance of genetic relative to demographic factors is still debated (e.g., Frankham 1995; Lande 1995; Spielman et al. 2004), 49 50 the 2 factors are expected to interact and feed back on each other, progressively driving 51 fragmented populations into "extinction vortices" (Lacy & Lindenmayer 1995) or "mutational 52 melt-downs" (Lynch et al. 1995; Higgins & Lynch 2001).

53 The potential effects of deleterious mutations on population fitness are often estimated 54 from the level of inbreeding load. Assuming a negative exponential relationship between 55 fitness and inbreeding coefficient of individuals within a population (Morton et al. 1956; 56 Kalinowski & Hedrick 1998), the slope of the regression of log(fitness) against inbreeding coefficient provides an estimate of inbreeding load, or number of lethal equivalents (reviewed 57 58 in Keller & Waller 2002). Inbreeding load in wild populations is commonly high (e.g., Ralls 59 et al. 1988; Kruuk et al. 2002; Reed et al. 2007) although exceptions exist (e.g. Duarte et al. 60 2003).

61 Viability losses, however, may also come from drift load, i.e. the local fixation of mild
62 deleterious mutations hidden from selection by drift (Keller & Waller 2002). Small

63 populations are actually expected to harbour more drift load (Whitlock et al. 2000) and less 64 inbreeding load than large ones (because individuals are genetically more similar locally; e.g., Bataillon & Kirkpatrick 2000). Local drift load is not revealed by regression of fitness on 65 66 inbreeding coefficient but by heterosis effects (i.e., fitness increase of offspring from crosses among compared to within populations), and has also received wide empirical support (e.g., 67 68 Coulson et al. 1998; Marr et al. 2002; Bush 2006). Since local populations may exhibit low 69 inbreeding load but high drift loads, management decisions made on the basis of inbreeding 70 depression only may be misleading.

71 Although the dramatic consequences of both inbreeding and drift loads have been 72 recognized, there is no simple way to incorporate them in the toolbox of conservation 73 geneticists without turning to heavy experimental designs (within and between population 74 crosses). Keller and Waller (2002) suggest use of F_{ST} as "an index of the susceptibility of a 75 population to the deleterious effects of drift load". Whitlock (2002) showed that the local drift load caused by mild deleterious mutations may indeed increase with F_{ST} in an infinitely large 76 77 metapopulation, depending on the mode of population regulation and mutation parameters. 78 Using stochastic individual-based simulations, Higgins and Lynch (2001) showed 79 metapopulation viability increases with the number, size, and connectivity of local demes. 80 Theodorou and Couvet (2006) further showed that, for a given metapopulation size, fitness is 81 higher with a few large populations than with several small ones, and that for small, isolated 82 populations the increase in local population size has a much greater positive effect on 83 population fitness than other parameters, such as migration or number of demes. 84 The effect of connectivity was formalized by the one-migrant-per-generation (OMPG)

rule, according to which one migrant per generation should suffice to protect local
populations from the accumulation of deleterious mutations (e.g., Mills & Allendorf 1996;
Couvet 2002; Wang 2004). However, migration rate is notoriously difficult to assess in the

field (Whitlock & McCauley 1999), and its effect may depend on other parameters such as
migration model, total metapopulation size, and mating system, which affects the purging of
deleterious mutations (Glémin 2003).

In the present study, we used stochastic individual-based simulations to investigate population fitness (offspring viability and time to extinction) under various metapopulation settings. The aims were (1) to derive robust predictive models of population fitness from easy-to-collect genetic and demographic data that may account for both inbreeding and drift loads, and (2) to test the validity of the OMPG rule under different migration models and mating systems.

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METHODS

99 Life cycle

100 We performed simulations in Nemo, a stochastic, individual-based, genetically-explicit 101 framework (Guillaume & Rougemont 2006). Model organisms were diploid, with separate 102 genders or not depending on the mating system, and lived in a structured metapopulation of d103 demes with local carrying capacity, N. A series of loci were subject to deleterious mutations, 104 whereas others were neutral. We implemented the following semelparous life cycle: (1) 105 viability selection on newly-born offspring that survived with a probability derived from their 106 deleterious mutation genotype; (2) dispersal of surviving offspring according to a specific 107 migration model (see below); (3) random regulation of local populations, which reduced the pool of competing individuals to the local carrying capacity (with equal sex ratios in case of 108 109 separate genders); (4) reproduction during which females were assigned a fecundity value 110 drawn from a Poisson distribution with constant mean f and were mated as many times as 111 indicated by their fecundity (one offspring per mating). Males were chosen according to 1 of

the 3 mating systems described below (random mating, selfing, or polygyny). Offspring
alleles at neutral and selected loci were inherited randomly (i.e., no linkage), baring
mutations. Sex was set randomly (with equal sex ratio) when genders were distinct. Adults
were removed after reproduction, and the cycle started again.

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117 Population structure, dispersal, and mating system

118 We ran simulations under different metapopulation configurations to cover a large range of 119 fragmentation levels. We varied independently local (*N*=4, 8, 16, 25, 50, and 100 individuals) 120 and total metapopulation sizes (N_t =200, 400, 800, and 1600). The number of demes (d) was 121 set by the ratio $d=N_t/N$. For each metapopulation configuration, we used 4 different migration 122 rates (m=0.001, 0.003, 0.01, and 0.1) and 2 different migration models (island and linear 123 stepping stone) that represented the 2 extremes of a continuum of isolation by distance. Most 124 realistic cases are likely to fall in-between. The effect of systematic inbreeding induced by 125 mating patterns was explored with 3 systems: random mating, selfing, and polygyny. Selfing 126 rate was set to 50%, the other 50% resulted from random mating within the deme. Under 127 polygyny, only one-quarter of the males present in each population were allowed to 128 reproduce, so successful males mated on average with 4 females.

We thus obtained a fully-factorial core set of simulations exploring 576 parameter combinations (6 local population sizes, 4 total population sizes, 4 migration rates, 2 migration models, 3 mating systems). The 6 combinations of mating systems and migration models will be referred to as datasets 1 to 6. For this core set, average fecundity *f* was set to 15 for females (random mating and polygyny) and 7.5 for hermaphrodites (selfing) in order to keep the same reproductive output per population. The effect of lowered fecundity (*f*=6) was investigated under random mating and island migration in an additional set of simulations (dataset 7).

137 Mutation models

The neutral markers, used to assess the level of neutral genetic diversity within and among populations, followed a *k*-allele mutation model (KAM), with *k*=256 possible allelic states over each of 24 loci and a mutation rate u=0.0001.

141 Fitness was controlled by a set of L (fixed to 1000) independent loci carrying deleterious alleles of various strength and dominance effect. We drew the number of new 142 143 mutations occurring in a particular genome from a Poisson distribution with mean equal to the 144 diploid genomic mutation rate (U). Mutations affected only nondeleterious alleles, turning 145 them into the deleterious form (reverse mutations were neglected), and acted independently 146 on fitness so that offspring viability (v) was computed as the product of fitness at each locus i: 147 $v=\prod_L v_i$, where v_i is 1, $1 - s_i$ or $1 - h_i s_i$ if the locus was homozygous wild-type, homozygous 148 mutant, or heterozygous, respectively. The values used for the mean mutation effect ($\bar{s} = 0.05$) and average dominance (\overline{h} =0.36) were derived from *Drosophila* studies (reviewed in Lynch 149 150 et al. 1999) and are commonly used in simulations (Wang et al. 1999; Higgins & Lynch 2001; 151 Theodorou & Couvet 2006).

152 For the core set of simulations, the genomic mutation rate was fixed to U=0.5, and the 153 mutant effects s were exponentially distributed among the L loci. Following Wang et al. 154 (1999), the dominance coefficient h of a mutation with effect s was set to satisfy the 155 relationship $h=\exp(-ks)/2$, where k is a constant chosen so that the average dominance of all mutations in the genome equals \overline{h} (Caballero & Keightley 1994). This induced an inverse 156 157 relationship between the magnitude of effect of a mutation and its degree of dominance, as 158 expected from biochemical arguments and supported by mutation accumulation experiments (Simmons & Crow 1977; Phadnis & Fry 2005). 159

160 We also performed additional simulations under random mating and island migration

to further explore the effects of genomic mutation rate (U=1, dataset 8) and the distributions of deleterious effects, assuming either a truncated log-normal (dataset 9) or a gamma distribution (dataset 10). The log-normal was parameterized according to Loewe and Charlesworth (2006) with log-mean = -6.4 and log-stdev = 5.3. The distribution was truncated to the right to have no s >1 (and \bar{s} =0.05). The shape parameter for the gamma distribution (α =1.69374) was taken from Keightley's (1994) estimation on the Mukai et al. (1972) *Drosophila* dataset, and its scale adjusted to have \bar{s} =0.05.

168

169 Simulations

170 For each of the 960 combinations of parameters (576 for the core set plus 384 for the 4 171 additional sets), we first performed 30 replicates over 50,000 generations with neutral markers 172 only (in order to get the required statistics for parameter values that would lead to population crashes in the presence of deleterious mutations), then 15 replicates over 5000 generations 173 174 after adding deleterious-mutations effects. At the start of simulations, neutral markers were 175 assigned random allelic values to insure a maximal initial variance, and loci under selection 176 were fixed to the fit allele. Statistics were recorded every 10 generations and measured from 177 the offspring that survived the viability selection episode.

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179 Statistical analyses

We computed mean F_{ST} , H_O , H_S , and H_T (Nei & Chesser 1983), first over the 30 neutral replicates (averaged over generations 20,000 to 50,000), second over the 15 replicates with deleterious mutations (generations 4,500 to 5,000), together with offspring viability v (for simulations in which all 15 replicates survived) and median time to extinction MTE (for simulations in which more than 50% of the replicates crashed before 5000 generations).

185 To find the best predictors of offspring viability from the core dataset we proceeded in 186 3 steps. First, we transformed the potential predictors $(F_{ST}, H_O, H_S, H_T, d, N, \text{ and } N_t)$ with the 187 functions x, log(x), 1/x, and 1/log(x), as well as log(1-x) for predictors ranging from 0 to 1, 188 and selected the transformations providing the best linear relationship with logit(viability) $(=\log \frac{v}{1-v})$. These turned out to be $\log(1-F_{ST})$, $\log(H_O)$, $\log(H_S)$, H_T , $\log(N_t)$, $\log(N)$, and 189 190 $\log(d)$. 191 Second, we performed linear regressions of logit(viability) on the transformed 192 predictors for each mating system and migration model independently (hence, 6 partitions).

193 The same analyses were performed on data pooled by mating system (3 partitions), migration

194 models (2 partitions) and then on the entire data set.

Third, we combined predictors 2 by 2 ($y \sim ax1 + bx2 + c$) to find the best bivariate prediction of logit (viability) on the same partitions as above. The different models were then ranked on the basis of the amount of explained variance, and ranks were averaged to select the best overall model.

We used the same procedure to predict median time to extinction from the core dataset (with 1/MTE as the dependent variable), and to analyze the additional simulations (datasets 7 to 10). We did not perform multiple stepwise regressions, which, owing to the high power of simulation studies, tend to retain too many variables, and usually different ones depending on the settings (data not shown).

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RESULTS

- 206 Genetic parameters calculated on neutral markers (F_{ST} , H_O , H_S , and H_T) did not differ
- 207 whether calculated in the presence or absence of deleterious mutations (correlation
- 208 coefficients ranging 0.97 to 0.99); thus, we consider only values in absence of mutation load

209 hereafter. The F_{ST} averaged 0.68 (range 0.007 to 0.995), $H_0=0.07$ (range 0.003 to 0.38),

210 H_S =0.08 (range 0.004 to 0.38), and H_T =0.49 (range 0.03 to 0.98). Spearman rank correlation

211 between the predictors ranged from -0.87 to 0.97 (Table 1).

212

213 Offspring viability

Over the surviving populations from the core dataset, offspring viability averaged 43%, depending greatly on the mating system and slightly on the migration model. It was highest under self-fertilization (averaging 49% and 45% for the island- and stepping-stone models of migration) but had a wide range (19% to 68%). Values were slightly lower under random mating (45% and 43% respectively, range 23% to 59%) and lowest under polygyny (37% and 33% respectively, range 18% to 56%).

Offspring viability was well predicted by $log(H_S)$, $log(1-F_{ST})$, and $log(H_O)$, with 41% to 67% of the variance explained depending on the partition used, but none of them ranking systematically higher (average ranking 1.7, 2.0 and 2.3, respectively). All 3 descriptors still explained at least 46% of variance when pooling data by mating system, but $log(H_O)$ lost explanatory power when data were pooled by migration model ($R^2 < 31\%$). When pooling all 6 partitions, $log(H_S)$ and $log(1-F_{ST})$ remained the best predictors ($R^2 = 46\%$ and 42%, respectively).

Both were also included in the best bivariate combination (Table 2), with an average
rank of 1.0 (i.e., best in all cases). When pooling all 6 partitions, 70% of variance (Table 2)
was explained by:

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$$\log \frac{v}{1-v} = a \log(1-F_{sT}) + b \log(H_s) + c \quad , \tag{1}$$

with a=0.39, b=0.52, and c=1.19. The explained variance increased to 71-74% when splitting data by migration models (2 partitions), 87-92% when splitting by mating system (3 partitions), and 90-97% when simultaneously splitting by migration model and mating system
(6 partitions; Table 2). Regression coefficients were positive in all cases, but viability
displayed a sharper transition from high to low values under selfing than under polygyny or
random mating (Fig.1). Simulations that collapsed because of mutational meltdown fell well
within the predicted low-viability area, even though these data were not used for model
fitting.

Decreasing fecundity (f=6, dataset 7) had no effect on offspring viability, and model (1) explained 85% of variance (Table 2). Increasing mutation rate (U=1, dataset 8) lowered offspring viability, but model (1) remained excellent (rank 2; $R^2=93\%$). The log-normal and gamma distributions of deleterious effects (datasets 9 and 10) had only marginal effects on offspring viability, and model (1) also remained the best (rank 1 in both cases), with 95% and 87% of variance explained respectively (Table 2).

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246 *Time to extinction*

Extinction rate averaged 51% over the core dataset (294 out of 576 simulations were extinct
before 5000 generations) and was higher under polygyny (70%) than under random mating
(45%) or selfing (38%). It was also higher under the stepping-stone dispersal (76%, 50%, and
43% for polygyny, random mating, and selfing respectively) than under island model (67%,
40% and 32% respectively).

Median time to extinction (MTE) did not differ much among mating systems (averages 1079, 1118, and 1137 generations under polygyny, random mating, and selfing respectively), with similar ranges (200 to 4000 generations). The 1/MTE correlated mainly with $log(1-F_{ST})$, $log(H_S)$, $log(H_O)$, and log(N) (with R^2 ranging from 22% to 83%), but none of these variables showed consistency among the different models. Accordingly, model ranks were very similar (2.2, 2.3, 3, and 3.2 for $log[1-F_{ST}]$, $log[H_S]$, log[N], and $log[H_O]$

respectively). After pooling all data, $\log(1-F_{ST})$ and $\log(N)$ remained the best candidates $(R^2=50\% \text{ and } 49\%, \text{ respectively})$, followed by $\log(H_S)$ ($R^2=43\%$).

260 When pooling all 6 partitions, the best bivariate model for predicting extinction time 261 combined $log(H_S)$ and log(N):

$$1/\text{MTE} = a\log(H_s) + b\log(N) + c, \qquad (2)$$

with $a = -8.36 \ 10^{-4}$, $b = -8.30 \ 10^{-4}$, and $c = 7.65 \ 10^{-4}$ ($R^2 = 68\%$; Table 3). The same model 263 emerged when considering the average ranking over the different partitions (rank 2.2; the 264 265 second-best model was $1/\text{MTE} \sim \log[N_t] + \log[H_0]$ with rank 2.8). The explained variance 266 reached 66-81% when splitting data by migration models (2 partitions), 81-86% when 267 splitting by mating system (3 partitions), and 79-97% when simultaneously splitting for 268 migration model and mating system (6 partitions; Table 3). Regression coefficients were 269 negative in all cases (Table 3 and Fig.2). Metapopulations still viable at generation 5000 fell 270 well within the predicted viable area, even though these data were not used for model fitting. 271 The model (2) was also good to predict median time to extinction in simulation runs 272 with f=6, U=1, and log-normal or gamma distribution of deleterious effects (average rank about 3), with 73-94% of the variance explained (Table 3). 273

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275 Inbreeding and purge of the genetic load

The ratio of offspring viability under selfing or polygyny relative to random mating was used to assess the extent of the purge in either of these mating systems. The ratio consistently exceeded unity in selfing populations (1.167 ± 0.136 SD), which had thus purged part of their mutational load. Polygynous populations underwent a higher rate of accumulation of deleterious mutations than under random mating (ratio 0.656 ± 0.168), inducing the higher extinction rates noted above. Note however that F_{IS} was not retained as a good predictor for 282 offspring viability or time to extinction in the regression analyses (data not shown).

283

284 One-migrant-per-generation rule

285 For both random mating and selfing, one migrant per population per generation was enough 286 to allow metapopulation persistence under our core settings (Fig.3b and c). None of the 287 simulations where effective migration rate (N_m) exceeded 1 went extinct, and there were only 288 a handful for N_m values between 0.1 and 1 (4 under selfing and 9 under random mating), 289 occurring under small local populations sizes (N=4 to 16) and low connectivity (stepping-290 stone dispersal). Under polygyny by contrast, extinctions occurred for N_m values exceeding 1, 291 but only at small metapopulation sizes (N_t =200). Lower fecundity values (f=6, Fig.3d) did not 292 affect offspring viability, but increased the threshold value below which populations are at 293 risk. Extinctions occurred for N_m values exceeding 1, but only when both total and local 294 populations sizes were small (N_t =200 and N<100). A higher genomic mutation rate (U=1, 295 Fig.3e) decreased offspring viability, so that extinctions occurred for N_m values exceeding 1, 296 but only for small metapopulation sizes (N_t =200).

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DISCUSSION

299 On the basis of our results, the effects of deleterious mutations on population fitness can be 300 largely accounted for by a few basic genetic and demographic measurements. Offspring 301 viability increased with genetic diversity within demes (H_s) and decreased with differentiation 302 among them (F_{ST}) , in line with both analytical treatments (Kimura et al. 1963; Whitlock et al. 303 2000; Whitlock 2002) and empirical observations (e.g., Madsen et al. 1996; Newman & 304 Pilson 1997; Saccheri et al. 1998). On its own, F_{ST} explained 42% of the variance in offspring 305 viability over our core dataset, corroborating Whitlock's (2002) analytical results under 306 infinite-island settings and supporting Keller and Waller's (2002) suggestion that

307 F_{ST} be used as "an index of the susceptibility of a population to the deleterious effects of drift 308 load". The positive role of diversity (H_S), on the other hand, more likely resulted from the 309 deleterious effect of inbreeding load. A combination of both H_S and F_{ST} accounted for both 310 loads and thus explained a large part of the variance in offspring viability (R^2 >85% for a 311 given mating system).

312 The median time to extinction also increased with diversity within demes (H_S) , but the 313 best bivariate regression included local population size (N), rather than F_{ST} , in addition to H_S (with $R^2 > 80\%$ for a given mating system). This point underlines the importance of both 314 315 demographic and genetic effects during the process of mutational meltdown, in line with both 316 analytical models (Lande 1994; Lynch et al. 1995) and empirical observations (e.g., Saccheri 317 et al. 1988). Small population sizes are known to enhance both demographic and genetic 318 stochasticity, with positive feedbacks. Populations collapsed at offspring viability values 319 below 0.2 for f=15 (and below 0.4 for f=6; Fig.3), corresponding to effective reproductive 320 rates exceeding unity, which, in absence of stochasticity, should allow positive growth rate 321 and population persistence. This illustrates the initiation of extinction vortices by the interplay 322 between demographic and genetic factors as soon as the system enters a critical state in terms 323 of population size and mutation load.

324 Local population size was not retained in offspring viability models, contrasting with 325 empirical support for a positive correlation between population size and fitness (reviewed in 326 Reed 2005; see also Reed et al. 2007). Local size certainly affects a population's ability to 327 resist drift, but our simulations also included other factors that drastically affect local genetic 328 diversity (mating system, migration rate, and total metapopulation size). Population fitness 329 and genetic diversity should depend more on local *effective* size, which may present only 330 weak correlations with *census* size when such interacting factors are varied. Mating systems 331 also had an effect of their own in lowering the relationship between population size and

offspring viability, since selfing, which reduces effective size, increased fitness by purging the
genetic load. Selection is more efficient at removing deleterious mutations under such mixed
systems than under random mating, due to increased variance in individual fitness and
inbreeding coefficients (see Glémin 2003 for an analytical treatment). Under polygyny, by
contrast, both the effective population size and variance in inbreeding were reduced, leading
to a greater rate of mutation accumulation and population extinction.

338 Total population size (N_t) also played a significant role in our simulations because the 339 dynamics of local genetic diversity within demes also depends on inputs from the 340 metapopulation reservoir. Depending on the mating system, 53% to 78% of the variance in H_S 341 was explained by $log(N_t)$. In small metapopulations ($N_t \leq 800$), furthermore, deleterious 342 mutations may get fixed at the global scale, with long-lasting consequences on population 343 fitness via drift load, but without contributing to inbreeding depression or heterosis anymore 344 (see also Whitlock 2002). Metapopulations of 200 individuals were often too small to persist, 345 owing to dramatically low offspring viabilities, whatever the connectivity (Fig.1 and 3). 346 These effects have been poorly investigated until now, mainly because analytical treatments 347 usually assume infinite or very large metapopulations (e.g., Whitlock 2002) and that previous 348 simulations studies have not addressed variance in this parameter (Higgins & Lynch 2001; 349 Theodorou & Couvet 2006).

Our results rejoin these 2 latter studies with respect to the effects of fragmentation. For the same total number of individuals (and no environmental stochasticity), one big population was better than several small. Doubling the number of populations was much less efficient than doubling the size of local populations. Increasing connectivity was quite efficient, provided that the total population size was not too small (>500) and that local populations were smaller than 100 individuals. We thus emphasize the importance for persistence of connecting isolated populations to a reservoir of genetic diversity.

357 Our results also provide some validation for the OMPG rule, with some caveats 358 however. As shown in Fig.3, populations did not collapse for effective numbers of immigrant 359 exceeding one under most parameter values. Exceptions occurred only for very small 360 metapopulation sizes (N_t =200), and only in conjunction with other negative effects such as 361 polygyny (Fig.3a), low fecundity (Fig.3d) or high genomic deleterious mutation rate (Fig.3e). 362 Migration rates and population sizes were deliberately set to low values in order to 363 simulate endangered populations, usually characterized by small global sizes (<2500 364 individuals, World Conservation Union 2001) and reduced connectivity. As a result, about 365 half of the simulations collapsed due to mutational meltdown (whereas the persisting ones 366 presented a large range of viability values), and population structure (F_{ST}) sometimes reached 367 values close to unity. This obviously exceeds the values usually documented in natural 368 situations because most situations fall within the range of 0 to 0.2 (Morjan & Rieseberg 369 2004). However, endangered populations frequently display F_{ST} values exceeding 20% (e.g., Rowe et al. 2000; Eckstein et al. 2006; Kawamura et al. 2007). Moreover, the F_{ST} values 370 371 measured for most of recently fragmented and/or bottlenecked populations are likely to be 372 underestimates because these populations usually have not had enough time to reach genetic 373 equilibrium (Whitlock 1992; Wang 2004). We thus covered a wide panel of population 374 genetic structures within which most endangered species are expected to fall. 375 One important point to emphasize is that our results must not be considered in

quantitative (absolute) terms, but only in qualitative (relative) terms, owing to the specificity of several assumptions underlying our simulations. This caveat obviously applies to our lifehistory assumptions. Lower fecundities, in particular, increase the viability threshold under which extinctions occur (Fig.3d), even though models 1 and 2 still hold qualitatively. Also, the genetic variance at neutral markers depends not only on effective sizes, but also on mutation rates, which might be species-specific and difficult to estimate precisely.

382 Similarly, the mutation model and parameters values used in our core simulations come from 383 accumulation experiments performed on a single model organism, Drosophila (Simmons & 384 Crow 1977; Lynch et al. 1999). Although our conclusions seem quite robust regarding the 385 distribution of deleterious mutations (Fig.3c), parameter values may vary among species. The 386 genomic mutation rate in particular quantitatively affected expectations (Fig.3e), even though 387 models 1 and 2 still hold qualitatively. Our results will be therefore best used in a comparative 388 context, e.g. to rank the effects of different management strategies for a given endangered 389 species. The conservation value of different scenarios can be evaluated on the basis of their effect on a set of very few key genetic and demographic parameters (H_S , F_{ST} , and N). More 390 391 specific questions might also be addressed with the same simulation framework (Nemo; 392 Guillaume & Rougemont 2006), or, if empirical data are available, by directly evaluating 393 regression coefficients of offspring viability on the relevant variables (i.e., H_S , F_{ST} , and N). 394 Our approach also bears a series of important advantages. First, it provides robust 395 predictive models with which to assess the viability prospects of fragmented populations, 396 which might usefully complement the OMPG rule (e.g., Frankel & Soulé 1981; Mills & 397 Allendorf 1996; Wang 2004) since assessing migration rates in nature is still a major 398 challenge in ecology (Whitlock & McCauley 1999), which often precludes its effective use. 399 Second, predictive power is large even without specific information on the mating system or dispersal model ($R^2 \approx 70\%$ on pooled data) and increases with additional information on the 400 mating system ($R^2 > 80\%$). Third, no lab breeding or controlled crosses are needed to estimate 401 402 inbreeding or drift load, which are both difficult or impossible to carry out on threatened 403 species. The genetic information required is readily obtained from neutral loci (e.g., 404 microsatellites, now easily available for many species) and can be sampled noninvasively 405 (e.g., from shed hair or faces, Taberlet & Luikart 1999; Broquet et al. 2007). The only 406 demographic information required is the size of local populations, which can be obtained

407	from basic field (e.g., mark-recapture) observations. Finally, the negative effects of both drift
408	load (fixed deleterious mutations) and inbreeding load (segregating deleterious mutations) are
409	accounted for, whereas empirical methods relying on the estimation of the number of lethal
410	equivalents (Morton et al. 1956) only consider the later. Given that small populations might
411	already be inbred to some degree, they are likely to have lost part of their standing variation
412	and fixed part of their mutation load. We hope our study will help clarify the effects of spatial
413	structure and connectivity on viability prospects of fragmented populations and provide
414	additional tools to evaluate extinction threats for endangered populations.
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547 TABLES

548 **Table 1**. Spearman rank correlations between the variables used as predictor of offspring549 viability and median time to extinction (MTE).

						550	
	$Log(N)^{a}$	$Log(d)^{b}$	$Log(1-F_{ST})^{c}$	$Log(H_O)^d$	$Log(H_S)^e$	H_T^{fr}	-
$Log(N_t)^g$	0.003	0.58	-0.12	0.35	0.35	0.53	-
Log(N)		-0.80	0.66	0.52	0.55	-0.55	
Log(d)			-0.60	-0.22	-0.24	0.76	
$Log(1-F_{ST})$				0.75	0.77	-0.87	
$Log(H_O)$					0.97	-0.43	
$Log(H_S)$						-0.44	

^a N = local population size, ^b d = number of demes, ^c $F_{ST} = \text{genetic structure}$, ^d $H_O = \text{observed heterozygosity}$, ^e $H_S = \text{within deme expected heterozygosity}$, ^f $H_T = \text{expected heterozygosity}$, ^g $N_t = \text{total metapopulation size}$.

- **Table 2**. Regression models for offspring viability as a function of F_{ST} and H_S for the
- 552 different data sets.^{*a*}

Detectb	Parameters ^c	Mating system and migration model ^d	Intercept	$Log(1-F_{ST})$		$Log(H_S)$		Total P^2	Donk
Dataset				Coef.	R^2	Coef.	R^2	I Otal K	Kalik
1	f=15, U=0.5, exponential	IM-random mating	0.903	0.543	0.41-0.54	0.442	0.36-0.49	0.90	1
2	<i>f</i> =15, <i>U</i> =0.5, exponential	IM-polygyny	1.411	0.598	0.46-0.51	0.678	0.43-0.48	0.94	1
3	f=15, U=0.5, exponential	IM-selfing	1.548	0.606	0.45-0.63	0.518	0.28-0.46	0.91	1
4	<i>f</i> =15, <i>U</i> =0.5, exponential	SSM-random mating	1.133	0.319	0.27-0.41	0.573	0.51-0.65	0.92	1
5	f=15, U=0.5, exponential	SSM-polygyny	1.877	0.380	0.31-0.40	0.903	0.57-0.65	0.97	1
6	f=15, U=0.5, exponential	SSM-selfing	1.976	0.381	0.26-0.57	0.726	0.36-0.67	0.93	1
1-3	f=15, U=0.5, exponential	IM	1.114	0.529	0.32-0.45	0.468	0.25-0.39	0.71	1
4-6	f=15, U=0.5, exponential	SSM	1.414	0.322	0.19-0.40	0.629	0.34-0.54	0.74	1
1, 4	f=15, U=0.5, exponential	Random mating	0.948	0.384	0.30-0.46	0.485	0.41-0.56	0.87	1
2, 5	f=15, U=0.5, exponential	Polygyny	1.494	0.460	0.36-0.47	0.737	0.50-0.56	0.92	1
3, 6	f=15, U=0.5, exponential	Selfing	1.645	0.454	0.33-0.59	0.585	0.30-0.56	0.88	1
1-6	f=15, U=0.5, exponential	All data pooled	1.185	0.391	0.24-0.42	0.523	0.28-0.46	0.70	1
7	f=6, U=0.5, exponential	IM-random mating	0.689	0.342	0.20-0.25	0.326	0.59-0.64	0.85	5
8	f=15, U=1, exponential	IM-random mating	-0.395	0.483	0.46-0.50	0.358	0.43-0.48	0.93	2
9	<i>f</i> =15, <i>U</i> =0.5, log-normal	IM-random mating	0.905	0.480	0.41-0.59	0.419	0.36-0.54	0.95	1
10	<i>f</i> =15, <i>U</i> =0.5, gamma	IM-random mating	0.798	0.516	0.49-0.61	0.323	0.26-0.38	0.87	1

^a Also shown are the intercept, the regression coefficients and the variance explained by F_{ST} and H_S , the total amount of variance explained, and the model ranking. Because order of introduction of variables in the model affects the amount of explained variance (but not the regression coefficients), partial R^2 is shown for each variable when introduced first and second. Significance levels of all coefficients and R^2 are below 0.0001.

^b Each dataset is assigned a number (1 to 10). When regressions were performed on pooled data, the datasets used are shown.

^c Shown are the fecundity values (per female), the genomic mutation rate and the type of distribution for the deleterious effects.

^d "IM" stands for "Island migration model" and "SSM" for "Stepping-stone migration model". "IM-random mating" means that all simulations run with random mating and island migration model were used in the regression (but with the same parameter values). "IM" means that all simulations run under island migration model with the same parameter values were pooled (independently of the mating system) and "Random mating" that all simulations run under random mating with the same parameter value were pooled (independently of the mating system) and "Random mating" that all simulations run under random mating with the same parameter value were pooled (independently of the migration model). "All data pooled" means that all simulations run with the same parameters values were pooled (independently of the mating system and migration model).

- **Table 3.** Regression models for median time to extinction as a function of *Hs* and local
- 554 population sizes for the different data sets.^a

D ()	Parameters ^c	Mating system and migration model ^d	Intercept	$Log(H_S)$		Log(N)		Total	D 1
Dataset*				Coef.	R^2	Coef.	R^2	R^2	Kank
1	f=15, U=0.5, exponential	IM-random-mating	$2.40 \cdot 10^{-03}$	-8.83·10 ⁻⁰⁴	0.13-0.46	-1.42·10 ⁻⁰³	0.38-0.71	0.84	3
2	f=15, U=0.5, exponential	IM-polygyny	$2.21 \cdot 10^{-03}$	- 8.41·10 ⁻⁰⁴	0.9-0.31	-1.17·10 ⁻⁰³	0.49-0.70	0.79	3
3	f=15, U=0.5, exponential	IM-selfing	-1.08·10 ⁻⁰³	-1.09·10 ⁻⁰³	0.39-0.76	-6.64·10 ⁻⁰⁴	0.17-0.54	0.93	2
4	f=15, U=0.5, exponential	SSM-random-mating	-5.75·10 ⁻⁰⁴ NS	-1.20·10 ⁻⁰³	0.25-0.83	-8.51·10 ⁻⁰⁴	0.12-0.70	0.95	2
5	f=15, U=0.5, exponential	SSM-polygyny	-1.11·10 ⁻⁰³	-1.23·10 ⁻⁰³	0.26-0.79	-6.76·10 ⁻⁰⁴	0.14-0.67	0.93	1
6	f=15, U=0.5, exponential	SSM-selfing	$-2.66 \cdot 10^{-03}$	-1.15·10 ⁻⁰³	0.50-0.94	-2.89·10 ⁻⁰⁴	0.3-0.46	0.97	2
1-3	f=15, U=0.5, exponential	IM	1.30.10-03	-8.73·10 ⁻⁰⁴	0.16-0.30	-9.85·10 ⁻⁰⁴	0.36-0.51	0.66	1
4-6	f=15, U=0.5, exponential	SSM	-1.34·10 ⁻⁰³	-1.15·10 ⁻⁰³	0.32-0.70	-5.61·10 ⁻⁰⁴	0.10-0.48	0.81	2
1,4	f=15, U=0.5, exponential	Random mating	$2.16 \cdot 10^{-03}$	$-7.71 \cdot 10^{-04}$	0.13-0.52	-1.29·10 ⁻⁰³	0.32-0.71	0.84	3
2, 5	f=15, U=0.5, exponential	Polygyny	1.43.10-03	-8.55·10 ⁻⁰⁴	0.13-0.48	-9.83·10 ⁻⁰⁴	0.34-0.68	0.81	2
3, 6	f=15, U=0.5, exponential	Selfing	-1.06·10 ⁻⁰³	-8.95·10 ⁻⁰⁴	0.36-0.73	-5.72·10 ⁻⁰⁴	0.13-0.50	0.86	2
1-6	f=15, U=0.5, exponential	All data pooled	$7.65 \cdot 10^{-04}$	-8.36·10 ⁻⁰⁴	0.19-0.43	-8.30·10 ⁻⁰⁴	0.25-0.49	0.68	1
7	f=6, U=0.5, exponential	IM-random mating	8.53·10 ⁻⁰³	-5.95·10 ⁻⁰⁴	0.04-0.08	-2.49·10 ⁻⁰³	0.65-0.69	0.73	3
8	f=15, U=1, exponential	IM-random mating	$4.47 \cdot 10^{-03}$	-1.43·10 ⁻⁰³	0.12-0.35	-1.91·10 ⁻⁰³	0.38-0.61	0.73	4
9	<i>f</i> =15, <i>U</i> =0.5, log-normal	IM-random mating	$5.56 \cdot 10^{-04}$	$-3.59 \cdot 10^{-03}$	0.21-0.51	$-4.88 \cdot 10^{-04}$	0.38-0.68	0.89	3
10	<i>f</i> =15, <i>U</i> =0.5, gamma	IM-random mating	$3.82 \cdot 10^{-03}$	-7.49·10 ⁻⁰⁴	0.09-0.45	-1.75·10 ⁻⁰³	0.49-0.85	0.94	3

^a Also shown are the intercept, the regression coefficients and the explained variance by H_s and local population size (*N*), the total amount of variance explained, and the model ranking. Because order of introduction of variables in the model affects the amount of explained variance (but not the regression coefficients), partial R^2 is shown for each variable when introduced first and second. Significance levels of slope coefficients and R^2 are below 0.01. For the intercept, p-values < 0.025 but one case noted NS.

^b Each dataset is assigned a number (1 to 10). When regressions were performed on pooled data, the datasets used are shown.

^c Shown are the fecundity values (per female), the genomic mutation rate and the type of distribution for the deleterious effects.

^d "IM" stands for "Island migration model" and "SSM" for "Stepping-stone migration model". "IM-random mating" means that all simulations run with random mating and island migration model were used in the regression (but with the same parameter values). "IM" means that all simulations run under island migration model with the same parameter values were pooled (independently of the mating system) and "Random mating" that all simulations run under random mating with the same parameter value were pooled (independently of the mating system) and "Random mating" that all simulations run under random mating with the same parameter value were pooled (independently of the mating system and migration model).

555 **FIGURE LEGENDS**

Figure 1. Offspring viability as a function of genetic differentiation (F_{ST}) and within-

557 population heterozygosity (H_s) for simulations performed under (a) polygyny (datasets 2 and

558 5), (b) random mating (datasets 1 and 4), and (c) selfing (datasets 3 and 6). Dots indicate

viable metapopulations, lighter symbols signalling higher offspring viability. The expected

isoclines for viability (lines) are calculated from the regression models in Table 2. Crosses
represent simulations that crashed before generation 5000. Data points corresponding to the

same total metapopulation size are aligned on the different curves

Figure 2. Median time to extinction (MTE) as a function of within-population heterozygosity (H_s) and population sizes (N) for simulations performed under (a) polygyny (datasets 2 and 5), (b) random mating (datasets 1 and 4), and (c) selfing (datasets 3 and 6) (x- and y-axes are log transformed for graphical presentation). Dots indicate metapopulations that collapsed, lighter symbols signalling longer time to extinction. The expected isoclines for MTE (lines) are calculated with the coefficients of the regression models in Table 3. Crosses represent simulations that survived at least until generation 5000.

570 **Figure 3.** Offspring viability as a function of effective migration rate N_m (log scale) for (a) 571 polygyny (datasets 2 and 5), (b) selfing (datasets 3 and 6), (c) random mating, pooling all 3 572 distribution models for deleterious effects (datasets 1,4, 9 and 10), (d) random mating with 573 lowered fecundity (f=6, dataset 7), and (e) random mating with increased genomic mutation 574 rate (U=1, dataset 8). A viability of zero is assigned to simulations that crashed, which 575 occurred whenever viability decreased below a threshold value (about 0.2 for f=15, and 0.4 576 for *f*=6). When effective migration rates exceeded 1 (vertical line), extinctions occurred only 577 at low metapopulation sizes (N_t =200, crosses) but never at larger sizes (open circles).

FIGURES

2 Figure 1





