Metapopulation effective size and conservation genetic goals for the Fennoscandian wolf (Canis lupus) population

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The Scandinavian wolf population descends from only five individuals, is isolated, highly inbred and exhibits inbreeding depression. To meet international conservation goals, suggestions include managing subdivided wolf populations over Fennoscandia as a metapopulation; a genetically effective population size of \( N_e \geq 500 \), in line with the widely accepted long-term genetic viability target, might be attainable with gene flow among subpopulations of Scandinavia, Finland and Russian parts of Fennoscandia. Analytical means for modeling \( N_e \) of subdivided populations under such non-idealized situations have been missing, but we recently developed new mathematical methods for exploring inbreeding dynamics and effective population size of complex metapopulations. We apply this theory to the Fennoscandian wolves using empirical estimates of demographic parameters. We suggest that the long-term conservation genetic target for metapopulations should imply that inbreeding rates in the total system and in the separate subpopulations should not exceed \( f = 0.001 \). This implies a meta-\( N_e \) of \( N_{Meta} \geq 500 \) and a realized effective size of each subpopulation of \( N_{E} \geq 500 \). With current local effective population sizes and one migrant per generation, as recommended by management guidelines, the meta-\( N_e \) that can be reached is \( \approx 250 \). Unidirectional gene flow from Finland to Scandinavia reduces meta-\( N_e \) to \( \approx 130 \). Our results indicate that both local subpopulation effective sizes and migration among subpopulations must increase substantially from current levels to meet the conservation target. Alternatively, immigration from a large (\( N_e \geq 500 \)) population in northwestern Russia could support the Fennoscandian metapopulation, but immigration must be substantial (5–10 effective immigrants per generation) and migration among Fennoscandian subpopulations must nevertheless increase.

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INTRODUCTION

The gray wolf (Canis lupus) was hunted to extinction in Scandinavia (Norway and Sweden; Figure 1) during the mid-1900s, and the present population was established in the early 1980s. This population descends from few individuals and has been highly isolated with census size remaining low for decades (Laikre et al., 2013), which has resulted in extreme inbreeding and inbreeding depression (Liberg et al., 2005; Räikkönen et al., 2013). The conservation case of the Scandinavian wolves is well known scientifically and politically (Liberg et al., 2005; Darpö, 2011; Laikre et al., 2013; Chapron, 2014). In this paper we address the topical issue of if and how conservation genetic goals with respect to genetically effective population size can be reached via increased connectivity with wolf populations further east.

The Scandinavian wolf population originates from a founder pair from Finland/Russia that had their first litter in Sweden in 1983. The population has been closely monitored; tracking data combined with molecular genetic analyses of biological material have provided an almost complete pedigree of the population (Vilà et al., 2003; Liberg et al., 2005). After the establishment in 1983 and up to 2012, a total of 18 additional wolves are known to have immigrated into Scandinavia from Finland/Russia, but only three of these have managed to reproduce; one male that first reproduced in 1990, and two males that both first reproduced in 2008. Thus, there is a total of five founders of the present population (Åkesson et al., 2014; Bruford, 2015; Mills and Feltner, 2015).

Population levels remained low during the first decades with substantial poaching, and authorities granting permissions to kill individuals considered to cause problems. Poaching is also viewed as the main reason for the restricted gene flow from populations in Finland and further east (Liberg et al., 2012). This has resulted in very high inbreeding levels, and the current c. 400 Scandinavian wolves are on average as inbred as offspring from full sib matings (\( f \approx 0.25 \); Jansson et al., 2015). Indications of inbreeding depression manifested as reduced litter size and a high frequency of spinal disorders has been reported (Liberg et al., 2005; Räikkönen et al., 2013).

Swedish authorities have issued several licensed wolf hunts during the past few years (SEPA Decisions 2009–2013), which have been strongly criticized by the European Commission (Potočnik, 2011; EC, 2015) because within the European Union (EU), where Sweden is a member, the wolf is listed as a priority species under the conservation legislation of the EU Habitats Directive (Council Directive 92/43/EEC). Similarly, according to national policy, Sweden is committed to maintain naturally occurring species at population sizes large enough to permit harboring genetic variation for long-term persistence (Swedish Environmental Objectives, Government Proposition 2009/10:155). The genetic situation of the Scandinavian wolves has been discussed for decades both
The genetically effective population size ($N_e$) is the most widely used parameter for quantifying and monitoring rates of inbreeding and loss of genetic variation through drift. The concept was developed for single, isolated populations; $N_e$ is defined as the size of an ideal population without the evolutionary forces of mutation, selection, or migration that exhibits the same expected rate of genetic drift or inbreeding per generation as the actual, real life population of interest (Wright, 1931). In conservation genetics the '50/500 rule' has become widely established suggesting that for a single isolated population $N_e > 50$ is needed for short-term conservation and $N_e > 500$ for long-term conservation (Franklin, 1980; Allendorf and Ryman, 2002; Jamieson and Allendorf, 2012).

Considerable research has been devoted to modeling $N_e$ of subdivided populations, but most of these efforts have used simplifying assumptions such as subpopulations of equal size, standard models of migration and drift–migration equilibrium (Wright, 1938; Maruyama and Kimura, 1980; Whitlock and Barton, 1997; Nunney, 1999; Wang and Caballero, 1999; Tufto and Hindar, 2003; Whitley, 2010). Results from that work include demonstrating that under the conditions of Wright’s island model global $N_e$ can increase above the sum of the local $N_e$s, while deviations from these idealized conditions with respect to asymmetrical migration patterns and/or asymmetries in population size can reduce the total effective size below the sum of the $N_e$s of the separate subpopulations (Whitlock and Barton, 1997; Nunney, 1999; Tufto and Hindar, 2003).

We have recently developed a general analytical framework for exploring inbreeding dynamics and effective population sizes in more complex metapopulations than has previously been theoretically or computationally feasible (Hössjer et al., 2014, 2015). Our analytical approach allows modeling systems with any number of subpopulations of arbitrary census and effective size. Migration patterns are also

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**Figure 1** The area of Fennoscandia (medium gray) comprises Norway and Sweden (Scandinavia), Finland and the Russian regions of Kola and Karelia. Detailed wolf distribution data are available for Finland and Scandinavia, and the dark gray areas indicate current wolf occurrence. Such information is missing for Karelia and Kola and it is thus unclear where in those areas wolves occur (marked with stripes). Numbers indicate estimated census sizes (cf. Supplementary Appendices 1 and 2).
optional, as are initial degrees of inbreeding and relatedness (coefficient of kinship; Wright, 1931) within and among populations.

Here we use this new analytical framework to first suggest how to apply the \( N_e \geq 500 \) rule to a subdivided population. We suggest new criteria for long-term conservation goals that take both subpopulations and the metapopulation as a whole into account. In a second step, we model the case of the Fennoscandian wolf to address the following questions: (i) what metapopulation effective size can currently be reached over Fennoscandia based on empirical estimates of parameters of interest? (ii) What is needed in terms of population sizes and migration rates for the Fennoscandian wolf to attain the suggested long-term conservation genetic goal for a subdivided population?

**MATERIALS AND METHODS**

We used the mathematical framework of Hössjer et al. (2014, 2015) that allows modeling the expected change of inbreeding and effective size of arbitrary metapopulations. Key expressions applied here include equation 25 of Hössjer et al. (2014) and equation 48 of Hössjer et al. (2015). Parameter notations used in the present paper are given in Table 1. We considered a diplodiploid organism without selfing where mating occurs after migration. Migration is stochastic, and rates are expressed as the actual number of immigrants per generation (\( m \)) from one subpopulation to another (cf. Figure 2). Thus, one migrant per generation, for example, refers to an average of one immigrant per generation based on binomial sampling. Because we focused on issues relating to rates of inbreeding only, the effective population sizes (rather than census sizes) are of concern. Thus, immigrants are regarded as ‘ideal’ individuals with the same characteristics as the ‘ideal’ individuals of the recipient population (Hössjer et al., 2014, 2015), and we ignored the forces of selection and mutation. We also assumed constant population sizes.

We dealt exclusively with inbreeding effective size (\( N_{eI} \), Wright, 1931,1938) which quantifies the rate at which inbreeding is increasing. There are also several other types of effective sizes (Crandall et al., 1999; Hössjer et al., 2015) but they are beyond the scope of the present focus. We analyzed how expected values of average inbreeding (\( \bar{f} \)) change over time in the separate subpopulations as well as for the system as a whole under various scenarios of subpopulation sizes and migration rates and patterns. Average inbreeding of the metapopulation (\( \bar{f}_{\text{Meta}} \)) corresponds to the probability that the two alleles at a locus are identical by descent in an individual drawn at random from the metapopulation as a whole (here weighting subpopulations according to effective size; Hössjer et al., 2015).

While the short-term conservation genetic rule of thumb \( N_e \geq 500 \) clearly refers to the inbreeding effective size (Franklin, 1980), the \( N_e \geq 500 \) rule is not directly stated in terms of a particular type of \( N_e \). We suggested isolated populations of constant size where inbreeding and variance (\( \sigma^2 \)) effective sizes always are the same (that is, \( N_e = N_e \), where \( N_e \) quantifies the rate of genetic drift, that is, allele frequency change). The \( N_e \geq 500 \) rule relies on the assumption that the loss of additive genetic variation for quantitative traits due to restricted population size is balanced by mutation when \( N_e \geq 500 \), and that additive genetic variation is lost at the same rate as heterozygosity (Franklin, 1980).

We quantified expected inbreeding rates in terms of effective population size. Effective size in generation \( t \) is defined as \( N_e = 1/(2\Delta f) \), where \( \Delta f = (f_0 - f_{t-1})/(1 - f_{t-1}) \) and \( f_t \) is the inbreeding coefficient in generation \( t \). In an isolated population of constant size, where \( \Delta f \) is constant and exclusively determined by drift (in the absence of immigration), this definition of \( N_e \) corresponds to the traditional one. In contrast, in a population receiving immigrants \( \Delta f \) is affected by both immigration and drift, and the corresponding quantity \( 1/(2\Delta f) \) is referred to here as realized effective size (\( N_{eR} \); Table 1).

The realized effective size of the total metapopulation is denoted \( N_{eMeta} \). This quantity can be computed using various schemes for weighting of the separate \( \Delta f_s \) of the different subpopulations (Hössjer et al., 2014, 2015), and here we weighted by local effective size (\( N_{el} \)) such that a large subpopulation contributes more to \( N_{eMeta} \) than a small one. Note that \( N_{eI} \) is the effective size of subpopulation \( x \) if it were completely isolated.

We also computed the eigenvalue effective size \( N_{eE} \) (Ewens, 1982; Hössjer et al., 2014), that is the effective size of the metapopulation as a whole when migration–drift equilibrium has been attained. In a metapopulation where each subpopulation both receives immigrants from, and sends emigrants to, the rest of the system (through one or more subpopulations) the rate of inbreeding will eventually be the same (\( 1/(2N_{eE}) \)) in all subpopulations as well as for the system as a whole (regardless of how the \( \Delta f_s \) of the different subpopulations are weighted; Hössjer et al., 2014, equation 61; Hössjer et al., 2015, equation 49).

The Hössjer et al. (2014, 2015) approach allows for specifying degree of relatedness between subpopulation \( x \) and \( y \) (\( f_{xy} \)) at the starting point (generation \( t = 0 \)). Subpopulation census size (\( N_s \)), and within subpopulation average inbreeding (\( f_t \)) and kinship (\( f_{xy} \)) affect the upper bound for \( f_{xy} \) (Hössjer et al., 2015; equation 2).

**Table 1 Definition/description of symbols used in this study**

| Symbol | Definition/description |
|--------|------------------------|
| \( t \) | Time measured in generations |
| \( m \) | Migration rate, expressed as number of immigrants per generation from a particular subpopulation; migration is stochastic and \( m \) reflects the binomial average. Migrants are regarded as ‘ideal’ individuals with the same characteristics as the ‘ideal’ individuals of the subpopulation into which they immigrate |
| \( f \) | Coefficient of inbreeding |
| \( f_x \) | Average inbreeding in subpopulation \( x \), that is, the expected inbreeding coefficient of an individual selected at random from subpopulation \( x \) |
| \( f_{xy} \) | Average kinship between individuals of subpopulations \( x \) and \( y \) |
| \( x \) | An arbitrary subpopulation that is part of a metapopulation. The subpopulations of the Fennoscandian wolf metapopulation that we analyze here are referred to as Scandinavia (and the subscript \( x \) replaced by Sc), Finland (subscript \( x \) replaced by Fi), Karelia (subscript = Ka), Kola (subscript = Ko) and northwestern Russia (subscript = Ru) |
| \( y \) | See \( x \) |
| \( f_{Meta} \) | Average inbreeding coefficient of the total metapopulation. This quantity corresponds to the expected inbreeding of an individual selected at random from the metapopulation as a whole (when weighting according to local effective size in the absence of immigration, Figure 2). Corresponds to \( f_t \) in Hössjer et al., 2015 |
| \( N_e \) | Inbreeding effective size (in general, note that this paper deals exclusively with inbreeding effective size) |
| \( N_{el} \) | Inbreeding effective size of subpopulation \( x \) (in the absence of immigration; cf. equation 23 in Hössjer et al., 2015) |
| \( N_{eR} \) | Realized effective size; reflects the change of inbreeding under both drift and immigration |
| \( N_{eMeta} \) | Total (global) inbreeding effective size of the metapopulation as a whole. This quantity reflects the change of \( f_{Meta} \) from generation \( t \) to \( t+1 \). It can be viewed as a weighted average of \( N_{el} \) over all subpopulations, or as the realized effective size of the metapopulation as a whole, and it will eventually approach \( N_{eE} \) |
| \( N_{eE} \) | Eigenvalue effective size, i.e. metapopulation effective size when migration–drift equilibrium has been attained |
Parameterization
We reviewed the literature to obtain empirical estimates from Fennoscandian wolf populations of parameters needed for modeling metapopulation-\(N_e\) (Supplementary Appendix 1; Supplementary Table 1). We translated the empirical data into a schematic metapopulation (Figure 2a), with local effective population sizes (\(N_{eK}\)) for Scandinavia (Sc), Finland (Fi), Karelia (Ka) and Kola (Kol) of \(N_{eSc} = 100\), \(N_{eFi} = 50\), \(N_{eKa} = 50\) and \(N_{eKol} = 25\), respectively. All but the \(N_{eKol}\) have an empirical basis; for Kola we used the census estimate of 100 and assumed an \(N_e/N\) ratio of 0.25 as estimated for the Scandinavian population (Supplementary Table 1).

We analyzed scenarios both with and without initial inbreeding in the system (Figure 2). With initial inbreeding we used starting point inbreeding averages of \(f_{Sc} = 0.25\), \(f_{Fi} = 0.10\), \(f_{Ka} = 0\) and \(f_{Kol} = 0\). We used \(f_{Sc} = 0\) rather than 0.07 estimated from \(F_{IS}\) (Supplementary Table 1) because that estimate did not deviate significantly from zero (Jansson et al., 2012). With respect to kinship among populations no direct estimates exist as far as we have been able to find; thus, we used approximations of \(f_{Sc} = 0\) between all populations except between Scandinavia and Finland where we used \(f_{Sc-Fi} = 0.05\) for those scenarios assuming initial inbreeding (Figure 2) because the Scandinavian founders have immigrated from Finland (Supplementary Appendix 1).

With respect to migration we assumed \(m = 1\) as a basis. Immigration from Finland to Scandinavia (\(m_{Fi-Sc}\)) has been monitored intensely and indicates \(m_{Fi-Sc} \approx 1\) (Supplementary Appendix 1; Supplementary Table 1). With respect to migration from Scandinavia to Finland (\(m_{Sc-Fi}\)) only three cases have been documented and two of these migrants were killed before being able to reproduce; the location of the third one—with radio collar—is unknown (Supplementary Appendix 1; Dr Jouini Aspi, University of Oulu, Finland pers. com.) suggesting \(m_{Sc-Fi} = 0\). Empirical data support immigration between Finland and Karelia of \(m_{Fi-Ka} \approx 1\) but with a recent decrease (Supplementary Table 1; Aspi et al., 2009; Jansson et al., 2012). No data appear to exist on migration to and from Kola. In addition to empirical support for \(m_{Fi-Ka} \approx 1\), the OMPG referred to in EU management guidelines for large carnivores (Linnell et al., 2008) stresses the need for evaluating the effect on \(N_e\) under this migration rate.

RESULTS
General aspects of metapopulation-\(N_e\) in conservation
First, some general aspects of effective population size in the context of metapopulations should be highlighted for clarity. \(N_e\) is only defined

\[ N_{eK} = N_{e} + \sum_{i > j} f_{ij} N_{eij} \]

\[ N_{eij} = N_{e} \frac{f_{ij}}{f_{ii} + f_{jj} - 2f_{ij}} \]

\[ f_{ij} = f_{ij}^{(0)} \frac{N_{eij}^{(0)}}{N_{eij}} \]

\[ N_{eij}^{(0)} = N_{e} \frac{f_{ij}^{(0)}}{f_{ii}^{(0)} + f_{jj}^{(0)} - 2f_{ij}^{(0)}} \]

\[ N_{eK} = N_{eK}^{(0)} \frac{f_{ij}^{(0)}}{f_{ii}^{(0)} + f_{jj}^{(0)} - 2f_{ij}^{(0)}} \]

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for populations where (i) there exists genetic variation and (ii) inbreeding increases or remains constant (when \( N_e = \infty \)). For populations with no genetic variation or with a decreasing level of inbreeding \( N_e \) is undefined. In a metapopulation, inbreeding level of a separate subpopulation can decrease as an effect of immigration, and in such situations the realized effective size of the receiving subpopulation \( (N_{eR}) \) becomes undefined. For instance, if subpopulation \( x \) with \( f_x > 0 \) starts receiving immigrants from an unrelated donor population \( y (f_y = 0) \) and with an immigration rate large enough for \( f_x \) to decrease, then the realized \( N_e \) of subpopulation \( x (N_{eR}) \) becomes undefined. In many situations \( f_x \) will eventually start to increase again, and at that ‘turning point’, when inbreeding goes from decreasing to increasing, \( N_{eR} \) will become definable again; initially it may be very large because the rate of increase of inbreeding is typically low at the turning point.

Similarly, for a metapopulation where subpopulations are completely isolated from each other (that is, \( m = 0 \)) we have \( N_{eE} = \infty \) because the subpopulations will eventually become fixed for different alleles; identity by descent of alleles of different subpopulations will not increase but remain the same generation after generation in the system as a whole. Thus, metapopulation-\( N_e \) is irrelevant for systems without migration between subpopulations, and as \( m \) approaches 0 and the system is close to migration–drift equilibrium, meta-\( N_e \) will approach its undefined state.

Conservation genetic goals for metapopulation-\( N_e \)

To our knowledge, conservation genetic goals reflecting long-term genetic viability have not been dealt with specifically for metapopulations. Rather, an implicit suggestion has been that for a subdivided population the same rule of thumb should apply as for a single, isolated one, that is, \( N_{eMeta} \geq 500 \) should reflect long-term viability for the metapopulation as a whole (Hansen et al., 2011; Laikre et al., 2013). We suggest, however, that \( N_{eMeta} \geq 500 \) cannot be the only focus for long-term genetic viability of a metapopulation. Rather, the inbreeding rates within the separate subpopulations must also be considered. We base this suggestion on the dynamics of a purely hypothetical subdivided population analyzed as an example here. We consider a subdivided population of constant size that consists of eight subpopulations, two of effective size \( N_e = 50 \) (subpopulations 1 and 2) and six of \( N_e = 25 \) (subpopulations 3–8) (Figure 3a). In this example, migration follows an island model such that in every generation each subpopulation receives \( m \) immigrants drawn at random from an infinitely large migrant pool to which the other subpopulations have contributed equally.

In the absence of subdivision, that is, under panmixia, the effective size of this population is \( N_e = 250 \), corresponding to the sum of the separate subpopulation effective sizes. Subdivision increases the total effective size (\( N_{eMeta} \)), but this increase is rather modest at high migration rates, and \( N_{eMeta} \) as well as the realized effective sizes (\( N_{eR} \)) of the separate subpopulations will approach the eigenvalue effective size (\( N_{eE} \)) within a few generations. For \( m = 5 \), for example, we have \( N_{eE} = 260 \), and all the realized effective sizes are within 90% of this value in <15 generations (not shown). At lower migration rates, however, \( N_{eE} \) grows progressively larger and the approach to equilibrium is much slower (Figure 3a). At \( m = 1 \) we have \( N_{eE} = 299 \) and it takes about 100 generations for the system to approach equilibrium; the corresponding value for \( m = 0.2 \) (one migrant every fifth generation) is \( N_{eE} = 504 \) and \( N_{eMeta} \) is clearly below this value even after 200 generations (Figure 3a).

As indicated in Figure 3, however, \( N_{eE} \) and \( N_{eMeta} \) are poor indicators of the rate of inbreeding in the separate subpopulations before equilibrium has been attained. At \( m = 0.2 \) the eigenvalue effective size is \( N_{eE} = 504 \) for this system, thus reflecting an overall rate of inbreeding of \( \Delta f \approx 0.001 \) per generation, so eventually the long-term genetic target will be reached for the metapopulation. Before this has happened, however, the realized effective size of

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**Figure 3** Realized (\( N_{eR} \)) and eigenvalue (\( N_{eE} \)) effective size (a) and inbreeding (b) over 200 generations in a hypothetical subdivided population of constant size that consists of eight subpopulations, two of effective size \( N_e = 50 \) (subpopulations 1 and 2) and six of \( N_e = 25 \) (subpopulations 3–8). Note that this is a hypothetical example that does not relate to the Fennoscandian wolves. Migration follows an island model such that in every generation each subpopulation receives on average \( m \) immigrants drawn at random from an infinitely large migrant pool to which the other subpopulations have contributed equally. Initial inbreeding and kinship is zero (0) within and between all subpopulations. The two curves in b marked \( N_e = 50 \) and 500 show the accumulation of inbreeding in isolated populations of that size (short- and long-term targets). Black curves, \( m = 1 \); gray curves, \( m = 0.2 \). See text for details.
subpopulations has been low for extended periods of time, resulting in an initial accumulation of inbreeding that for the smaller subpopulations (3–8) and for the system as a whole even exceeds that of the short-term conservation goal (Figure 3b). Thus, although meta- \( N_e \) eventually will reach the long-term conservation genetic goal, the inbreeding rates of separate subpopulations will initially (over 50–75 generations) exceed acceptable levels for even short-term conservation goals.

We suggest that the conservation genetic target for metapopulations to reflect long-term genetic viability should imply that the rate of inbreeding in the system as a whole, as well as in the separate subpopulations, should not exceed \( \Delta f = 0.001 \) (as for an \( N_e \) of 500), that is,

1. metapopulation effective size is \( N_{eMeta} \geq 500 \)
2. realized effective size of each subpopulation equals or exceeds 500 (\( N_{eRx} \geq 500 \)).

Further we suggest that goals 1–2 should be reached within, say, 5–20 generations, that is, within the time frames typically considered in the context of short-term conservation (for example, Allendorf and Ryman, 2002). We now apply this proposed conservation genetic target for long-term viability to the Fennoscandian wolf metapopulation.

**Meta-\( N_e \) of a closed Fennoscandian population assuming no initial inbreeding**

First we modeled a ‘closed’ Fennoscandian population using current estimates of local effective sizes, no inbreeding or kinship, and migration patterns implying that all subpopulations both receive and contribute migrants (that is, two-way migration; Figure 2a; Supplementary Appendix 1; Supplementary Table 1). The results indicate that it is not possible to reach a metapopulation \( N_{eMeta} \geq 500 \) for this system without a very restricted gene flow that results in excessive rates of inbreeding within the separate subpopulations as well as for the system as a whole (Figure 4). With the very restricted migration rate of 1 migrant every 20th generation (\( m = 0.05 \)), the equilibrium metapopulation \( N_{e} \) is \( N_{e} = 871 \) (Figure 4a). However, the system is not even close to this equilibrium within the first 200 generations (corresponding to c. 1000 years assuming a generation length of 5 years as estimated for the Scandinavian population; Forslund, 2009).

Further, for \( m = 0.05 \) metapopulation \( N_{e} \) (\( N_{eMeta} \)) increases from 57 to 161 over the initial period of 200 generations and stays below 115 for the first 100 generations, as do all the subpopulations. Thus, inbreeding rates are much above the long-term target rate over this period, and over the first c. 50 generations inbreeding accumulates almost as fast as for the short-term conservation goal of \( N_e = 50 \) in the Karelian and Finnish subpopulations, as well as in the metapopulation as a whole (Figure 4b). In Kola, the effect of immigration does not reduce the accumulation of inbreeding below the short-term goal until after more than 100 generations, and the rate of inbreeding in the Scandinavian population is, of course, also much higher than the target of \( \Delta f = 0.001 \) for the initial 200 generation period (Figure 4b).

With the higher migration rates of \( m = 1 \) and 3, the rate of inbreeding drops below the short-term target in all subpopulations, as well as for the system as a whole within the first few generations (Figures 4d and f). However, the long-term target of metapopulation \( N_e \geq 500 \) is never reached, as \( N_e = 250 \) and 233 for \( m = 1 \) and 3, respectively (Figures 4c and e). Attaining equilibrium is much quicker than at low migration rates, however; the system as a whole is close to equilibrium within 50–150 generations, and at that point realized effective size of all the subpopulations are close to the eigenvalue effective size (\( N_{eRx} \approx N_{eMeta} \approx N_{e} \)).

Clearly, meta-\( N_e \) increases above the sum of the local effective sizes at low migration rates and decreases with increasing migration. When migration rates are high enough to make the system as a whole approach panmixia, meta-\( N_e \) will approach the sum of local population sizes. In the present case the system is close to \( N_{eMeta} = N_{e} = 225 \) (100 + 50 + 50 + 25) at \( m = 3 \). Thus, low migration results in a high \( N_e \) at the cost of high rates of inbreeding in the subpopulations for many generations, and thereby of the system as a whole.

In a next step, we doubled the subpopulation effective sizes (that is, \( N_{eFsc} = 200 \), \( N_{eFi} = 100 \), \( N_{eKol} = 100 \) and \( N_{eKol} = 50 \)) and found that \( N_{e} \) becomes close to or above 500 for all three migration rates (not shown). Again, \( N_{e} \) is large (\( \approx 1742 \)) with the lowest migration rate of \( m = 0.05 \), but it takes a very long time to approach equilibrium; \( N_{eMeta} \) is far below 500 and only increases from 113 to 210 over the first 200 generations. Also, with the higher migration rates of \( m = 1 \) and 3 the equilibrium values of \( N_{e} = 499 \) (\( m = 1 \)) and \( N_{e} = 466 \) (\( m = 3 \)) are not reached until after c. 100 generations or more.

In another model we increased subpopulations of Scandinavia and Finland proportionally to a total size of 400 (\( N_{eFsc} = 267 \) and \( N_{eFi} = 133 \)), while assuming that Kola and Karelia remain at \( N_{eKol} = 50 \) and \( N_{eKol} = 25 \), and using \( m = 3 \). The rationale for increasing only Scandinavia and Finland is that both Sweden and Finland are members of the EU and have agreed to the EU target of maintaining long-term viable populations of naturally occurring species, whereas Russia is not a member of EU. The sum of subpopulation effective sizes is now close to 500 (475) and \( N_{e} \) becomes 489, which is reached by \( N_{eMeta} \) at \( t = 100 \).

Clearly, the conservation genetic target that we suggest for metapopulations is not possible to reach for a closed Fennoscandian wolf population unless subpopulation effective sizes (in isolation) increase so that their sum approaches 500. Further, substantial gene flow among subpopulations is necessary (\( m > 3 \)) to reach the target within reasonable time and to avoid excessive inbreeding during the approach to equilibrium.

**Effects of one-way vs two-way migration between Scandinavia and Finland**

Empirical observations suggest that gene flow may have been unidirectional from Finland to Scandinavia since the current Scandinavian population was founded in the early 1980s (Supplementary Appendix 1). Thus, we investigated the potential effects of one-way vs two-way migration between Finland and Scandinavia on the Fennoscandian metapopulation, still assuming no initial inbreeding in the system. We used current estimates of subpopulation effective sizes (Figure 2a) and \( m = 1 \) implying that Finland, Kola and Karelia receive one migrant each from the other two, whereas the migration between Finland and Scandinavia is either two-way or one-way from Finland to Scandinavia (Figure 2a).

Eigenvalue effective size (\( N_{e} \)) is almost doubled under two-way migration as compared with one-way migration from Finland to Scandinavia (\( N_{e} = 250 \) and 132, respectively; Figure 5). On the way to equilibrium, however, the Scandinavian population benefits from one-way migration over the first c. 100 generations. The realized \( N_e \) for Scandinavia increases more rapidly with one-way migration, and at generation \( t = 75 \), for example, we have \( N_{eFsc} = 244 \) and 214 for one-way and two-way migration, respectively. This benefit for the Scandinavian population occurs at a cost for the other subpopulations and for the metapopulation as a whole, however; at \( t = 75 \) we have \( N_{eMeta} = 169 \) vs 221 for one-way vs two-way migration (Figure 5). Similarly, realized \( N_e \) for Finland,
Karelia and Kola are all at \( N_e = 132 \) under one-way migration, as compared with \( N_e = 241, N_e = 222 \) and \( N_e = 214 \) with two-way (not shown). In fact, neither of Finland, Karelia or Kola reaches a realized \( N_e \) above 132 with one-way migration from Finland to Sweden, while they are above this value already in generation 22 under two-way migration (not shown).

At first glance these results may appear surprising, but they can be explained as follows. Under one-way migration (from Finland
to Scandinavia) the Scandinavian population behaves as a ‘genetic sink’ only receiving migrants from a metapopulation comprising Finland, Kola and Karelia. The effective size of this smaller metapopulation (without Scandinavia) is \(N_{\text{meta},\text{Kol},\text{Ka}} = 132\), and with the present population sizes and migration rates the ultimate fate of the Scandinavian subpopulation is completely determined by the Finland–Kola–Karelia population system. Thus, under one-way migration each of the four populations will eventually approach a realized effective size of \(N_e = 132\). In contrast, under two-way migration all the four subpopulations are connected through two-way exchange of migrants with at least one of the other subpopulations. This results in a larger metapopulation with an effective size of \(N_{\text{meta}} = 250\), and the realized effective size of each of the subpopulations will eventually approach this number (\(N_e = 250\)). Further, the reason why \(N_{\text{meta}}\) is larger under one-way migration than under two-way is that Scandinavia then receives immigrants that are unrelated during a longer time than with two-way migration when Scandinavian exports migrants to Finland and can receive descendants of those migrants back (cf. Hössjer, 2015).

**Meta-\(N_e\) of a closed Fennoscandian population assuming initial inbreeding**

Next, we explored meta-\(N_e\) for the Fennoscandian wolf when there is initial inbreeding in Scandinavia (\(f_{\text{fi}} = 0.25\)) and Finland (\(f_{\text{fi}} = 0.1\)) as observed in these populations (Supplementary Table 1), including a kinship between Scandinavia and Finland of \(f_{\text{fi}} = 0.05\). Other starting parameters were as before (Figure 2a). We examined both one-way and two-way migration between Finland and Scandinavia, using \(m = 1\) in all cases of migration.

Now, \(N_e\) becomes undefined for the Scandinavian and Finnish subpopulations during an initial period when inbreeding decreases from the starting values of \(f_{\text{fi}} = 0.25\) and \(f_{\text{fi}} = 0.1\) (Figure 6a). With one-way migration the decrease of inbreeding continues over 39 generations for the Scandinavian population when \(f_{\text{fi}}\) levels off around 0.24 and then begins to increase again (not shown). In the Finnish subpopulation the decrease only occurs for two generations; \(f_{\text{fi}} = 0.092\) in \(t = 2\) and then starts to increase slowly again.

With two-way migration the decrease of inbreeding continues until \(t = 7\) for Scandinavia when \(f_{\text{fi}}\) reaches its lowest value at \(f_{\text{fi}} > 0.24\), and until \(t = 4\) for Finland when \(f_{\text{fi}} = 0.09\). At the turning points where the decrease of inbreeding levels off and inbreeding starts to increase again, the increase is initially very small and \(N_{\text{Efi}}\) and \(N_{\text{ERfi}}\) are therefore very large. For instance, at generation \(t = 8\) with two-way migration \(N_{\text{Efi}} \approx 18,000\), and \(N_{\text{ERfi}} = 493\) (as compared with \(N_{\text{Efi}} \approx 116\), and \(N_{\text{ERfi}} = 113\) in the case of no initial inbreeding; cf. Figure 6a vs Figure 4c). The final equilibrium values are, however, the same as in the case of no initial inbreeding, that is, \(N_e = 132\) and 250 for one-way and two-way migration, respectively (Figure 6). Thus, the initial reduction of inbreeding in Scandinavia and Finland results in realized effective population sizes of those subpopulations that to begin with are larger than when starting with no inbreeding, but with \(m = 1\) this early effect on \(N_{\text{ER}}\) has leveled out around \(t = 150–200\) both in the case of one-way and two-way migration (Figure 6a).

As observed for the case with no initial inbreeding (Figure 5), one-way migration from Finland to Scandinavia also results in a ‘favorable’ situation for Scandinavia (with respect to the rate of inbreeding). The
reason is that the immigrants are less related to the Scandinavian wolves than in the situation with two-way migration when Scandinavia exports genes, which can later migrate back again. For example, at $t = 75$ and initial inbreeding we have $N_{eRu} \geq 570$ and 355 for one-way and two-way migration, respectively (Figure 6a).

When starting with inbreeding in the Finnish and Scandinavian populations the average inbreeding coefficient of the metapopulation as a whole increases from $f_{Meta} \geq 0.13$ to 0.49 and 0.43 in generation $t = 200$ for one- and two-way migration between Finland and Scandinavia, respectively. Clearly, after 200 generations the effect of initial inbreeding is minor relative to the case of no initial inbreeding (cf. Figure 4d).

With two-way migration and initial inbreeding the level in Finland decreases from the starting value of $f_{E} = 0.1$ to 0.088 in generations $t = 5–8$ and then increases again, passes $f_{E} = 0.1$ in $t = 16$ and reaches $f_{E} = 0.16$ in $t = 50$. In Kola and Karelia $f_{Col}$ and $f_{Ka}$ pass 0.1 in generation $t = 16$ and 21, respectively. At $t = 50$, $f_{Col} \approx 0.20$ and $f_{Ka} \approx 0.19$.

**Meta-$N_e$ with gene flow from Russia east of Fennoscandia**

Clearly, a Fennoscandian wolf metapopulation cannot reach the conservation genetic target we propose unless the sum of subpopulation effective sizes approaches 500 and migration between subpopulations is extensive. Further, the results above indicate that inbreeding levels will remain high if the Fennoscandian population is a closed population. We modeled the effects of inflow of genes from the Russian population east of the Russian provinces of Kola and Karelia (Figure 1). Census estimates indicate around 3000 wolves in this area (Supplementary Appendix 2), and assuming $N_e/N = 0.25$ this corresponds to an effective size of $N_{eRu} = 750$ (Figure 2b).

As expected, currently estimated inbreeding levels in Fennoscandia can be reduced with inflow from Russia. However, for a positive effect of immigration from Russia the migration within the Fennoscandian metapopulation must be extensive. For example, with a bidirectional gene flow within Fennoscandia of $m = 1$, a unidirectional inflow from Russia of 10 wolves per generation (Figure 2b) will never result in a reduction of average $f$ of the metapopulation ($f_{Meta}$) below the initial value of 0.13. In contrast, with $m = 5$ within Fennoscandia, $f_{Meta}$ drops to 0.1 within the first few generations, and it is reduced to this level also in Scandinavia around generation 30, with $f_{E}$ below 0.15 in generation 12 (not shown).

Metapopulation effective size of a Fennoscandian wolf population with currently estimated population sizes can increase to $N_{eMeta} \geq 500$ with immigration from Russia. Again, gene flow within Fennoscandia is critical for how long this will take. For example, even if 10 immigrants from Russia enter Fennoscandia via Karelia each generation $N_{eMeta}$ will not reach 500 until generation $t = 90$ if migration within Fennoscandia is $m = 1$. Assuming a generation interval of 5 years (Forslund, 2009) this translates into 430 years. With $m = 2$, 3 or 5 within Fennoscandia the time to $N_{eMeta} \geq 500$ is 52, 37 and 26 generations, respectively (that is, in 130–260 years). With only one immigrant from Russia every generation $N_{eMeta} = 500$ is reached at generation $t = 99$ (that is, in ~500 years) with a genetic exchange within Scandinavia of $m = 5$. Clearly, if applying a shorter generation length of, for example, 3.4 years as estimated for the Finnish population (Supplementary Appendix 1) these time frames in years are shorter.

**DISCUSSION**

We used recently developed theory to address issues of long-term conservation genetic targets of metapopulations, with particular focus on the Fennoscandian wolf population. We suggest that the well-established rule of thumb for long-term genetic conservation of $N_e \geq 500$, corresponding to an increase of inbreeding of $\Delta f \leq 0.001$, should apply to the metapopulation as a whole as well as to each of the subpopulations. Thus, long-term genetic viability of metapopulations includes meeting the criteria of (i) metapopulation effective size $N_{eMeta} \geq 500$ and (ii) realized effective size of each subpopulation is $N_{eSub} \geq 500$. We also suggest as a conservation target that these two goals should be reached within the time frames of short-term conservation, that is, typically 5–20 generations. We argue that less rigorous criteria than those of long-term conservation would be acceptable during short-term conservation periods, but long-term criteria should be met when short-term conservation periods have passed.

When applying these criteria to the empirical case of the Fennoscandian wolf our findings can be summarized as follows:

1. The current Fennoscandian wolf population cannot reach the above long-term conservation target. Rather, assuming generally recommended rates of gene flow ($m \approx 1$; the OMPG rule) and current local effective population sizes, the meta-$N_e$ that can be reached is 132 or 250 (see next point).

2. Two-way migration between all subpopulations is essential for the genetic health of the metapopulation. Unidirectional gene flow from Finland to Scandinavia, rather than bidirectional, greatly reduces metapopulation effective size. With one-way migration between those subpopulations $N_{eE} = 132$, while $N_{eS} = 250$ with two-way migration (assuming one migrant per generation).

3. To reach the above conservation criteria subpopulation effective sizes must increase such that their sum approaches 500. Further, the bidirectional migration rates among subpopulations must increase substantially above one migrant per generation. Thus, the OMPG rule is not sufficient in this particular case.

4. If subpopulation effective sizes are not allowed to increase, the genetic viability of the Fennoscandian wolf population must be dependent on gene flow from the northwestern Russian population. Several conditions must be met for such gene flow to have the wanted effect, however. First, migration from northwestern Russia must be substantial, c. 5–10 migrants per generation. Second, gene flow among the Fennoscandian subpopulations must be high (c. 3–5 migrants per generation). Further, the Russian population supporting migrants to the Fennoscandian metapopulation must be large and comprise an effective size of at least $N_e = 500$ (data not shown).

Several general aspects of the concept of effective population size and its application to metapopulations have become apparent in this study. First, asymmetrical gene flow can have a great impact on metapopulation effective size and realized $N_e$ of separate subpopulations. In our case study of Fennoscandian wolves this is exemplified by the markedly reduced $N_{eE}$ resulting from one-way migration from Finland to Scandinavia. Second, the parameter $N_e$ can be difficult to interpret for metapopulations where inbreeding decreases in some subpopulations, because it then becomes undefined and can take extremely large values at the turning points when inbreeding starts to increase regardless of actual inbreeding levels (cf. Figure 6), thus poorly reflecting the actual conservation genetic situation. Third, the OMPG rule is not generally applicable in the context of metapopulation genetic conservation. Rather, necessary migration rates to avoid excessive inbreeding depend on the number of subpopulations, their effective sizes, as well as the pattern of gene flow among them.

We have suggested two criteria for long-term genetic viability of metapopulations: (i) $N_{eMeta} \geq 500$; and (ii) $N_{eSub} \geq 500$. In fact, it is sufficient to apply the second criterion because when this criterion is met the first one will also be satisfied for the diplodiploid model of
inbreeding effective size (cf. Hössjer et al., 2015). We chose to keep both criteria for clarity.

International conservation policies agreed by Finland, Norway and Sweden stipulate that viable populations of wolves should be maintained in the area. Such agreements are particularly strong for Finland and Sweden as members of the EU, but also for Norway, which has signed and ratified the Bern Convention (as have Finland, Sweden and other EU members) that lists the wolf in its Appendix II of strictly protected fauna (www.coe.int/bernnovation). In contrast, Russia is not part of either EU or the Bern Convention.

Our results indicate that criteria for long-term genetic viability are not met for the current Fennoscandian wolf population. In order to meet such criteria, local population sizes as well as migration rates between subpopulations need to increase. Alternatively, genetic support from Russia east of the province of Karelia is needed. Even with such gene flow, however, considerably improved connectivity among the Fennoscandian populations is necessary.

Knowledge is scarce on population genetic characteristics, dynamics and connectivity of wolf populations in the Russian provinces of Kola and Karelia of Fennoscandia, as well as of those in neighboring Russian provinces (Supplementary Appendix 1; Supplementary Table 1), and research to generate such information is urgently needed. Similarly, management cooperation is not well developed (Dr Jouni Aspi, University of Oulu, Finland pers.com.). Thus, it may be dangerous for Scandinavia and Finland to rely on gene flow from northwestern Russia in order to meet conservation goals for wolves in Fennoscandia. Further, due to the lack of common conservation policy, management collaboration and information it can be argued that Scandinavia and Finland alone should take responsibility for maintaining a long-term viable wolf population (Nilsson, 2015). This requires that the sum of local subpopulation-Ne is ≈ 500, with a high degree of connectivity. For instance, if Ne = 300 and Nm = 200 the time for NeMeta and NEdxp to reach 500 varies in the range 15–50 generations for two-way migration in the interval 5 < m < 20. Assuming an Nm/N ratio of 0.25 (Forslund, 2009; Bruford, 2015) this would translate into 1200 wolves in Sweden and 800 in Finland, with an exchange of 20–80 individuals per generation.

The present study represents an application of a new mathematical framework to a practical scenario; in fact, it was the case of the Fennoscandian wolves that initiated the mathematical developments by Hössjer et al. (2014, 2015). Clearly, however, this framework can be applied to many other situations where exploration of metapopulation effective size is of interest.

Also, there are several general aspects of metapopulation effective size that we have not been able to address here. For instance, we have noted that it is unclear from the original presentation by Franklin (1980) which N is his Ni ≥ 500 rule applies to. Some authors have suggested that it is the variance effective size (Harmon and Braude, 2010), but since loss of additive genetic variation is associated with loss of heterozygosity (Franklin 1980) it is actually the inbreeding effective size that is applicable here. The haploid inbreeding effective size might be most relevant because it quantifies loss of gene diversity (cf. Hössjer et al., 2014). We elaborate on this in forthcoming work (Hössjer et al., unpublished). NeMeta is larger for the haploid inbreeding model than for the diploid one because rate of inbreeding is quantified based on two alleles drawn at random from the whole population rather than from individuals. However, the difference for Nedxp between the haploid and diploid models is negligible.

Further, in the present model we have ignored the forces of mutation and selection. Adding mutation to the model is not expected to change our basic conclusions because we consider relatively short time frames from an evolutionary perspective— even our long-term perspectives in conservation is typically to be regarded as short-term from an evolutionary standpoint. Selection is difficult to include because a wide range of potential selective regimes are possible. With directional selective pressures, however, rates of inbreeding are expected to increase at a higher rate than in a situation without selection. Similarly, growing immigration into inbred populations such as Scandinavia and Finland may initially result in a genetic rescue effect where less inbred individuals contribute more progeny than others, thus increasing the variance of reproductive success. Such a higher variance is expected to increase the rate of inbreeding and thus reduce realized effective population sizes during the initial phases of the approach to equilibrium. This kind of genetic rescue effect with extensive spread of genes from an immigrating individual followed by increased inbreeding and associated inbreeding depression was documented empirically in the isolated and highly inbred Isle Royale wolf population (Hedrick et al., 2014).

More theoretical work is needed with respect to several aspects of metapopulation effective size such as exploring the effects of selection and the behavior of other types of effective sizes (such as haploid vs diploid Ne, variance, linkage disequilibrium, coalescence effective sizes and so on) in metapopulations. We are in the process of continuing research along the lines of both to explore metapopulation effective size dynamics, including applying our current framework to other practical conservation cases of concern, and to further address the mathematical theory of metapopulation effective sizes.

Concluding recommendations on Fennoscandian wolves

On the basis of the results from this study we find the following management recommendations warranted for the Fennoscandian wolves.

1. Two-way migration between Finland and Scandinavia needs to be secured.
2. Monitoring of migration rates among all subpopulations of Fennoscandia is required.
3. The degree of connectivity needed among subpopulations is in the order of c. 3–5 genetically effective migrants per generation.
4. Local effective population sizes need to increase such that the sum of the subpopulation Ne that comprise the metapopulation approaches 500.
5. Owing to the lack of a common conservation policy between EU member states (Sweden and Finland) and Russia, we propose that the conservation target of meta-Ne ≥ 500 should be reached for wolf populations of Scandinavia and Finland.

DATA ARCHIVING

There were no data to deposit.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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**Supplementary Information** accompanies this paper on Heredity website (http://www.nature.com/hdy)