

Convergent development of low-relatedness supercolonies in *Myrmica* ants

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Many ant species have independently evolved colony structures with multiple queens and very low relatedness among nestmate workers, but it has remained unclear whether low-relatedness kin structures can repeatedly arise in populations of the same species. Here we report a study of Danish island populations of the red ant *Myrmica sulcinodis* and show that it is likely that such repeated developments occur. Two microsatellite loci were used to estimate genetic differentiation (F_{ST}) among three populations and nestmate relatedness within these populations. The F_{ST} values were highly significant due to very different allele frequencies among the three populations with relatively few common alleles and relatively many rare alleles, possibly caused by single queen foundation and rare subsequent immigration. Given the isolation of the islands and the low investment in reproduction, we infer that each of the populations was most likely established by a single queen, even though all three

extant populations now have within-colony relatedness <0.06 . All populations can be regarded as supercolonies with high nest densities and competitive monopolisation of the habitat patch excluding all other ant species. The investment in new sexuals was extremely male-biased ($>95\%$), and the genetic differentiation of nests showed a significantly positive correlation with the distance between them. Both male-biased sex-ratio and genetic viscosity are expected characteristics of populations where queens have very local dispersal and where new colonies are initiated through nest-budding. Based on a comparison with other *M. sulcinodis* populations we hypothesise a distinct succession of population types and suggest that this may be a possible pathway to unicoloniality, ie, development towards a complete lack of colony kin structure and unrelated nestmate workers. *Heredity* (2002) 89, 83–89. doi:10.1038/sj.hdy.6800098

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Introduction

In eusocial insects, workers increase their inclusive fitness by rearing the brood of related individuals (Hamilton, 1964). The workers thus have an interest in keeping within-colony relatedness high. This is indeed the case when nestmate workers are the offspring of a single, once-mated queen and nestmate relatedness is 75%. There is, however, considerable variation in social structure among extant populations and species. Polygyny (multiple queens per colony) and polyandry (multiple mating of queens) occur frequently (Bourke and Franks, 1995; Crozier and Pamilo, 1996) decreasing the genetic relatedness among nestmates.

An exceptionally derived form of social organisation is found in the so-called unicolonial ants. In these societies nest boundaries disappear and supercolonies are formed in which workers and queens freely move around (Passera, 1994). Well studied examples of species forming unicolonial populations are the Argentine ant *Linepithema humile*, the red imported fire ant *Solenopsis invicta*, and the red ant *Myrmica rubra*. In all these cases colonies are

highly polygynous, aggression among nests is absent, and populations consequently become genetically homogeneous with relatedness among nestmates close to zero (eg, Ross *et al*, 1996; Seppä and Walin, 1996; Holway *et al*, 1998; Krieger and Keller, 2000).

A unicolonial type of social organisation is expected to be evolutionary unstable because workers help rearing unrelated brood (Sturtevant, 1938; Crozier, 1977; Bourke and Franks, 1995; Queller and Strassmann, 1998) but paradoxically, unicolonial species are among the ecologically most successful ants in the world (Hölldobler and Wilson, 1990; Passera, 1994). The evolutionary origin and maintenance of unicoloniality is thus elusive and a matter of some controversy (eg, Tsutsui *et al*, 2000; Giraud *et al*, 2002) as it was recently proposed that several pathways under different genetic and ecological conditions may lead to this social structure (Chapman and Bourke, 2001).

Ants of the genus *Myrmica* are highly suitable to study the evolution of unicoloniality, because they show an extreme flexibility in kin structure across populations. In both *M. sulcinodis* (Pedersen and Boomsma, 1997) and *M. rubra* (Seppä and Walin, 1996) kin structures varying from maximum nestmate relatedness (0.75) and single colony-queens to zero relatedness and highly polygynous nests have been observed. Allozyme studies from five different locations showed that worker nestmate relatedness in *M. sulcinodis* ranges from less than 0.06 in

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a population in Denmark to more than 0.50 in the Pyrenees, Russia and Finland (Seppä *et al*, 1995; Pedersen and Boomsma, 1997). Also, population structure and inbreeding are very variable in this species. In the Pyrenees, nests were very discrete and widely scattered, while on the Danish island of Samsø multiple nests may belong to the same polydomous colony and nest densities are very high, saturating the entire suitable habitat (Pedersen and Boomsma, 1999a,b). Inbreeding was observed in the Pyrenees and in Russia (involving the production of a high frequency of diploid males; Pedersen *et al*, 1999), while Danish and British populations did not show any detectable inbreeding (Pedersen and Boomsma, 1997). An explanation for this intraspecific variation could be that these *Myrmica* populations follow a distinct succession of social stages, and that they were in different successional stages at the moment of sampling. This scenario would include an invasive colonisation stage, followed by expansion, increasing population densities and declining nestmate relatedness. The Pyrenean and Russian populations would thus represent an early stage, where genetic bottlenecks have caused a low number of alleles at the sex determination locus, causing the high number of diploid males. When populations survive this stage the genetic variation will gradually increase due to immigration, primarily of long distance-dispersing males. Consequently inbreeding decreases and expansion through nest-budding starts. Low relatedness populations such as the one on the Danish island of Samsø would then represent a later stage in the succession process, which is achieved only in long term stable habitats that can maintain high population densities. These latter populations tend to completely saturate the habitat (Pedersen and Boomsma, 1999a), but it is as yet unclear whether habitat saturation is a cause or an effect of this final development towards a supercolony structure (Rosengren and Pamilo, 1983; see also Discussion).

The purpose of the present paper is to analyse the population structure of two other isolated populations of *M. sulcinodis* in Denmark, to estimate the genetic differentiation between these populations and the population from the island of Samsø, and to assess whether these populations represent multiple samples of related end points of social development that disperse from island to island, or whether they are independent multiple end points of social succession towards unicoloniality, starting from a single queen colonisation in every population.

Materials and methods

Sample collection

The red ant *M. sulcinodis* is widely distributed from the Apennines to Arctic Scandinavia and from Portugal to East Siberia (Collingwood, 1979). Sampling was carried out in July and August 2000 on Læsø (57°17' N, 10°54' E), a Danish island in the Kattegat of about 20 × 12 km. During a period of 4 weeks all potentially suitable habitat of *M. sulcinodis* (heathland mainly covered with *Empetrum nigrum*, *Calluna vulgaris* and *Dechampsia flexuosa*) was searched. Two discrete populations were found on opposite ends of the island and will be referred to as Læsø1 and Læsø2, respectively. No other *M. sulcinodis* population was found, but very small populations may

have been overlooked (although not within 200 m from each of the populations, see below).

The two populations were situated ca. 16 km apart, about the maximum distance possible on the island. As previously found on the island of Samsø (Pedersen and Boomsma, 1999b), both populations turned out to be very dense, with *M. sulcinodis* nests being dominant within the occupied area and other ant species being common only at the edges of the patch (see also Results). All nests of the two populations were mapped by searching every possible nest site in one-by-one metre quadrates. The area around the two populations was also searched thoroughly, both for isolated *M. sulcinodis* nests and for nests of other ants, in a radius of up to 200 m or until a road, a dune ridge or dense bushes bordered the suitable habitat. Other ant species were included on the map if found within 10 m from a *M. sulcinodis* nest. Directly after mapping, nests (usually clearly visible as elevated small earth mounds in between the vegetation) were opened and 20–40 workers per nest and all visible winged sexuals were collected. Most males and new queens gather in the upper part of the nest (T van der Hammen and JS Pedersen, personal observation), so although some sexuals may have escaped collection the sex ratio and number of individuals obtained are likely to be comparable to studies where complete nests were excavated (eg, Pedersen and Boomsma, 1998; Walin *et al*, 2001). All ants were brought to the laboratory alive and stored at –80°C until genetic analysis.

Nests used for genetic analysis were chosen by drawing gridlines on the maps and selecting the nests closest to crossings of the gridlines. This resulted in a regular sampling scheme including 14 nests for Læsø1 in a 15 m grid and 17 nests for Læsø2 in a 6 m grid (Figure 1). The genetic analysis further included up to 50 workers from 23 randomly chosen nests previously sampled in a single large population on the Danish island of Samsø (situated 138 km south of Læsø, 55°56' N 10°36' E; Pedersen and Boomsma, 1999a).

Molecular genetic analysis

Ten old workers per nest were analysed for both Læsø1 and Læsø2. Old workers can be recognised by their advanced cuticular melanisation and were chosen because at Samsø workers from different age cohorts are genetically differentiated (Pedersen and Boomsma, 1999a), so that mixing of age cohorts could bias relatedness estimates. DNA was extracted from two legs from each of these workers. The samples were incubated in 100 µl 5% Chelex (in 10 mM TRIS buffer, pH 7.5) at 100°C for 15 min. Alternatively a long extraction version (5% Chelex, 1 h incubation at 55°C and 15 min. incubation at 100°C) or a DNeasy-kit were used to extract DNA. The suspension was stored at –20°C and used directly for PCR. Microsatellite primers developed for *M. tahoensis* (Evans, 1993) were tested and revealed genetic variation at the loci *Myrt2* and *Myrt4* in *M. sulcinodis*. PCR conditions were as follows: *Myrt2* (94°C for 3 min (93°C for 30 sec, 45°C for 40 sec, 72°C for 1 min) × 40 cycles followed by 72°C for 5 min), *Myrt4* (94°C for 3 min, 93°C for 30 sec, 56°C for 40 sec, 72°C for 2 min) × 40 cycles (reduced to 30 cycles when analysed on the automated sequencer) followed by 72°C for 1 h). PCR amplification was subsequently tested on 2% agarose gels. Microsatellites were visualised on spreadex S50 and S100

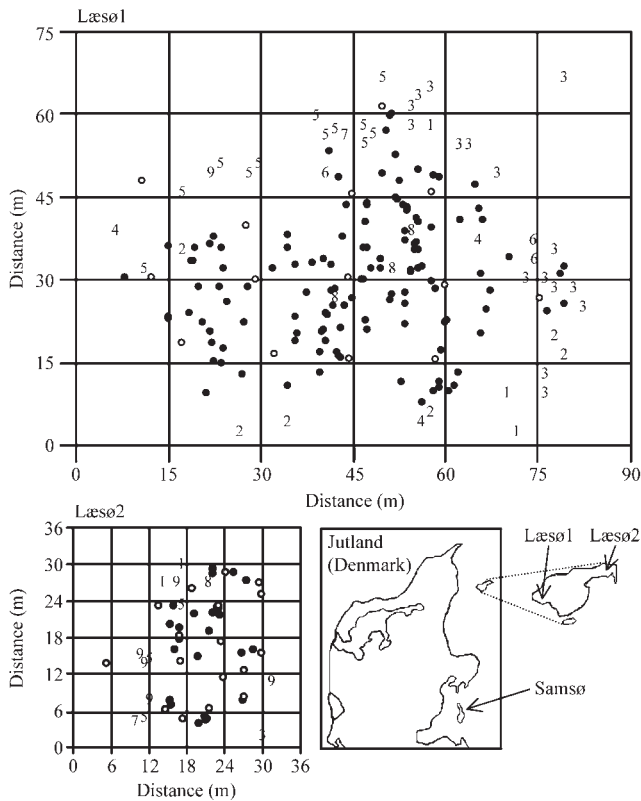


Figure 1 Distribution of *Myrmica sulcinodis* nests (circles) in the two Læsø populations. *M. sulcinodis* nests closest to the crossing of grid-lines were used for genetical analysis and are indicated with open circles. Nests of other ant species inside the *M. sulcinodis* population and within an envelope of ca. 10 m outside the aggregation of *M. sulcinodis* nests are indicated with numbers: (1) *M. ruginodis*, (2) *M. rubra*, (3) *Formica cinerea*, (4) *F. rufa*, (5) *F. fusca*, (6) *Lasius flavus*, (7) *L. psammophilus*, (8) *Leptothorax acervorum*, and (9) *Tetramorium caespitum*.

gels (Kozulic, 1999). Alternatively, when PCR products were ambiguous or spreadex resolution insufficient, analysis was carried out on an ABI 377 automated sequencer.

Statistical analysis

Exact tests for deviation from Hardy-Weinberg equilibrium within populations, genotypic differentiation of nests within populations, and genotypic differentiation of populations were performed by FSTAT 2.9.1 (Goudet, 1995, 2000). This program was further used for calculating matrices of pair-wise F_{ST} estimates for nests within populations. Matrix correlation coefficients and Mantel tests for the association of pair-wise F_{ST} and geographic distance between nests were calculated by the computer programme ARLEQUIN 2.001 (Schneider *et al*, 2001). For data including both Læsø populations these statistics were calculated as partial matrix correlations with a third matrix controlling for the fact that nests were only paired with nested populations).

Estimates of genetic relatedness (r) among nestmates, inbreeding (F_{IS}) and genetic differentiation among populations (F_{ST}) were obtained by the computer program RELATEDNESS 4.2c (Goodnight and Queller, 1996), which uses the algorithms developed by Queller and Goodnight

(1989). Average values were found by weighting nests (for estimates of r and F_{IS}) or populations (for estimates of F_{ST}) equally. Standard errors were obtained by jackknifing over nests (r and F_{IS}) or populations (F_{ST}). Parametric t -tests were used to test whether estimates of r and F showed significant deviations from zero. The effective queen number (n_e) was calculated using equations 1 and 5 in Pedersen and Boomsma (1999a). This represents the average number of equally fertile queens per colony that would lead to the estimated relatedness among nestmate workers (cf. Ross, 1993).

Results

The two populations of *M. sulcinodis* on Læsø (Læsø1 and Læsø2) consisted of 143 and 44 nests, respectively (Figure 1). Within each of the patches *M. sulcinodis* was very abundant and hardly any other ant nests were found, whereas other ant species were common at the edges and outside the *M. sulcinodis* nest-aggregations (Figure 1). The nest density in the larger Læsø1 population was lower than in the Læsø2 population (0.064 vs 0.098 nests/m² of inhabited area) which is reflected in a significantly longer distance between nearest neighbour nests (distance \pm SD = 2.1 \pm 1.7 vs 1.6 \pm 1.8 m; Mann-Whitney U -test; $P = 0.027$).

Both microsatellite loci were highly variable with 11 alleles detected at *Myrt2* and 21 alleles found at *Myrt4* across the two populations (Figure 2). Average expected heterozygosities were 0.64 and 0.85 for *Myrt2* and *Myrt4*, respectively, and 0.70, 0.79 and 0.75 on average for Læsø1, Læsø2 and Samsø, respectively. Estimates of relatedness among nestmate workers were low ($r < 0.06$) but significantly positive for both Læsø populations (Figure 3) with an estimate of $r = 0.051 \pm 0.016$ ($P = 0.002$) for the total sample. The positive nestmate relatedness is further confirmed by the highly significant genotypic differentiation of nests (F_{STAT} , exact tests; $P < 0.0001$ for both populations). None of the Læsø populations showed

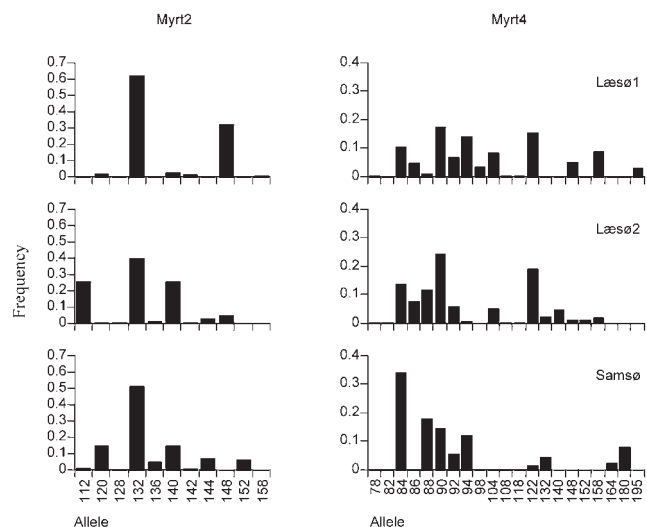


Figure 2 Allele frequencies at the microsatellite loci *Myrt2* and *Myrt4* (Evans, 1993) in workers of the three *Myrmica sulcinodis* populations studied: Læsø1 ($n_{\text{nests}} = 14$, $n_{\text{alleles-Myrt2}} = 249$, $n_{\text{alleles-Myrt4}} = 230$), Læsø2 ($n_{\text{nests}} = 17$, $n_{\text{alleles-Myrt2}} = 310$, $n_{\text{alleles-Myrt4}} = 315$), and Samsø ($n_{\text{nests}} = 23$, $n_{\text{alleles-Myrt2}} = 86$, $n_{\text{alleles-Myrt4}} = 65$).

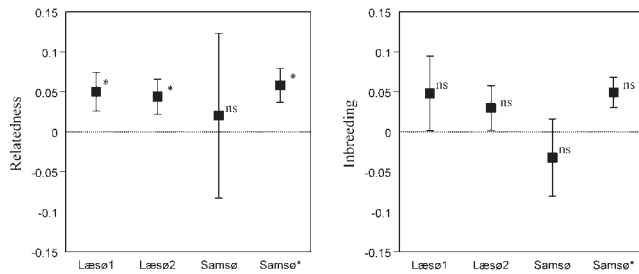


Figure 3 Estimates of worker-worker relatedness (r) and inbreeding (F_{IS}) for *Myrmica sulcinodis* in the three populations. Values are means for nestmate workers \pm SE estimated by jackknifing over nests. Asterisks in the figure indicate significant deviation from zero ($P < 0.05$). Samsø* are previous estimates based on allozyme loci from Pedersen and Boomsma (1999a).

any detectable inbreeding (Figure 3; common $F_{IS} = 0.037 \pm 0.024$; $P > 0.16$) or a significant deviation from Hardy-Weinberg equilibrium at the population level ($P > 0.10$ for both populations). A comparison of these results with estimates of relatedness and inbreeding for Samsø showed that the present markers essentially produce the same results for all three populations. Furthermore, estimates based on microsatellites are equivalent to previous estimates based on allozyme loci (Figure 3), the only difference being larger SEs because of lower sample size for the microsatellite analysis of the Samsø population.

The low estimates of worker relatedness result in a high effective queen number per colony, being 16 on average for the Læsø populations under the assumption of random mating and unrelated, single mated and equally reproducing queens. This estimate increases to $n_e = 19$ when nestmate queens are assumed to be equally related as nestmate workers, which is common in polygynous ant societies (Crozier and Pamilo, 1996).

The relatedness between workers in pairs of nearest nests on the grid analysed was significantly smaller than the average within-nest relatedness ($\Delta r = 0.034 \pm 0.017$; paired t -test; $P = 0.028$) indicating that nests at this scale (6–15 m) are not part of the same polydomous colony. Thus the nests analysed are likely to represent different, independent colonies. However, at the scale of the total population, close nests tend to be more genetically similar than more distant nests as shown by a relatively strong and significantly positive correlation of genetic and spatial distance ($r = 0.17$; Figure 4).

Exact tests revealed that the gene pools of all three populations studied were significantly differentiated (Bonferroni corrected $P < 0.001$ for all three comparisons) with estimates of pairwise F_{ST} being 0.076 for Læsø1–Læsø2 and 0.062 for both Læsø1–Samsø and Læsø2–Samsø. The estimated average F_{ST} among all three populations was 0.067 ± 0.009 . Hence, the two populations at Læsø, which are about an order of magnitude closer to each other than to the Samsø population, are not genetically more similar to each other than to the population on Samsø.

The number of winged sexuals found during sampling at Læsø1 was relatively low ($n = 89$) and all but two were males. This gave a highly male biased numerical sex ratio of 0.978, corresponding to a relative investment in new queens of only 0.042 (calculation based on dry weights in Pedersen and Boomsma, 1998). The sex ratio of Læsø1

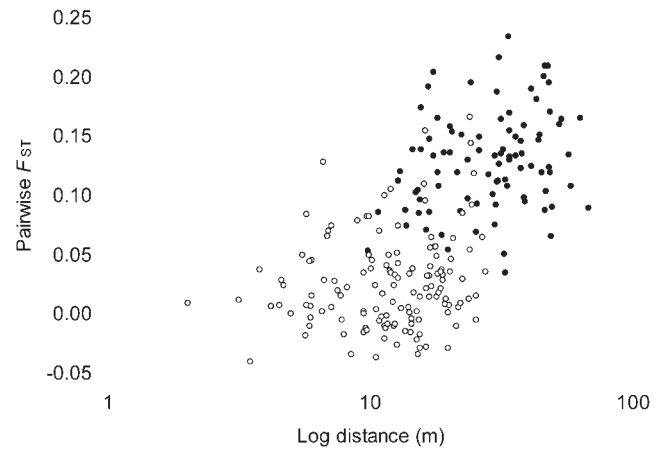


Figure 4 Association of genetic distance measured as pairwise F_{ST} and spatial distance between pairs of nests in Læsø1 (closed circles; matrix correlation $r = 0.22$; Mantel test, $P = 0.070$) and Læsø2 (open circles; $r = 0.21$; $P = 0.046$). The positive correlation was highly significant in the combined test ($r = 0.17$; Mantel test with nested populations, $P < 0.001$).

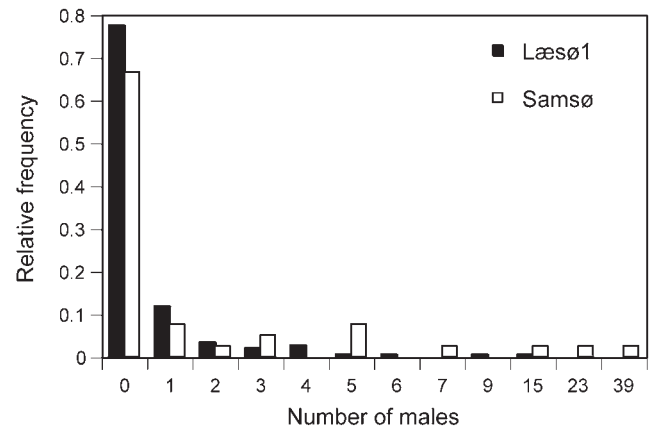


Figure 5 Distribution of males found in nests of the Læsø1 population (closed bars; $n = 87$) and the Samsø population (open bars; $n = 107$). The two distributions are not significantly different (Mann-Whitney U -test; $P = 0.18$).

was very close to a previous sex-ratio estimate for the Samsø population ($M/F = 107/1$; Pedersen and Boomsma, 1998), and also the distribution of males across nests was remarkably similar for the two populations (Figure 5). Sampling of Læsø2 was too late in the season to find sexuals.

Discussion

Similar low-relatedness supercolonies

Both populations of *M. sulcinodis* on Læsø have essentially the same kin structure as the previously studied population on Samsø. Nest densities were high in all cases, saturating the occupied habitat and effectively excluding all other ant species (Pedersen and Boomsma, 1999a). This is remarkable, because the sizes of the populations were very different. The two Læsø populations differed by a factor three in both area covered and nest number, but both were still small compared to the Samsø

population, which covered an area of 15 ha and comprised about 11 000 nests (Pedersen and Boomsma, 1999a), ie, about two orders of magnitude more than the Læsø populations. The nest density is the same at both islands, being 0.070 nests/m² as a weighted mean for the Læsø populations and 0.073 on Samsø. However, the patch structure of the habitat on Samsø results in a more clumped distribution of nests with only 0.35 ± 0.20 m as the mean (\pm SD) distance between nearest neighbour nests within patches (Pedersen and Boomsma, 1999b). This illustrates that the Samsø population is very patchy and completely saturated, because virtually every suitable patch is occupied by densely clustered nests of *M. sulcinodis*. This was not the case at Læsø, where empty nest sites still seemed to be available and where the present nests were relatively discrete. The important conclusions of these across-island comparisons are that (1) *M. sulcinodis* is able to monopolise habitat patches of vastly different sizes, and that (2) such populations have a very similar kin-structure (low, but significantly positive relatedness), breeding system (slightly positive but non-significant inbreeding) and mode of reproduction (extremely male-biased sex ratio).

Genetic differentiation of populations

Genetic differentiation (F_{ST}) among the three *M. sulcinodis* populations studied was highly significant. Moreover, about one-third (12/35) of the alleles were only found in a single population and the high variability of the microsatellite markers used most likely implies that genetic differentiation is underestimated by F -statistics (Hedrick, 1999). Perhaps surprisingly, a comparison of the pair-wise estimates of F_{ST} showed that the two Læsø populations were not genetically more similar to each other than they were to the Samsø population, suggesting that long-distance migration between populations is equally rare at various larger distances and that common alleles are likely to represent the alleles carried by the founding individuals (founder effect).

Where do the new queens come from?

The extremely male biased sex ratio and the relatively strong correlation of genetic and spatial distance at the scale of total populations (Figure 4) firmly support the argument that the dominant mode of reproduction of *M. sulcinodis* on Læsø is nest budding within the habitat patch. Queens joining a group of workers to initiate a bud-nest avoid the mortality associated with long-distance dispersal and the risk of not finding a suitable nest site after short-distance dispersal. The latter may be particularly advantageous in dense, saturated populations where budding is likely to be an optimal strategy for colonies to monopolise resources in the suitable patch (Rosengren and Pamilo, 1983; Pedersen and Boomsma, 1999b). Considerable genetic viscosity, a likely result of nest budding, has earlier been shown to occur in other polygynous ant species of the genera *Myrmica* (Seppä and Pamilo, 1995), *Formica* (Chapuisat *et al*, 1997; Liautard and Keller, 2001), and *Rhytidoponera* (Tay *et al*, 1997).

An intriguing remaining question is how low-relatedness populations of *M. sulcinodis* recruit their replacement queens, as it seems that they hardly produce gynes in most breeding seasons. In an earlier study on Samsø, Pedersen and Boomsma (1998) also found a very low overall sexual productivity of nests (108 sexuals of

both sexes in 39 nests over 2 years), whereas the productivity of British populations with moderate nestmate relatedness was an order of magnitude larger (Elmes, 1987; see also below). Although we cannot exclude the possibility that both island studies accidentally sampled bad years for sexual production, in different years and on different locations, this possibility seems less likely than assuming that saturated habitats in fact select for low investment in sexual reproduction, relative to the production of new workers needed for colony budding (cf. Pamilo, 1991). Another case of apparent underproduction of replacement queens was recently reported for a population of *M. rubra* with comparable social structure (Walin *et al*, 2001).

As a working hypothesis for explaining the paradox of queen recruitment in *M. sulcinodis* (ie, low rates of production of new queens *vs* high abundance of breeding queens), we suggest that colonies either must produce more gynes in some breeding seasons that we have so far failed to sample, or that they rely on immigrating queens from isolated nests or long-distance populations with a more distinct kin-structure and a more even sex ratio. Such populations are known to occur in *M. sulcinodis* (see below) and are probably more easily overlooked than saturated populations like the ones investigated on Samsø and in the present study. However, immigration would tend to remove the observed genetic viscosity of the population (Figure 4) and is thus not likely to be a major source of new queens. A third and not mutually exclusive explanation is that *M. sulcinodis* queens are very long-lived, so that these supercolonies can maintain high effective queen-numbers and thus long-term stability with low recruitment of new queens. This contrasts with the common view that a short lifespan is typically expected in polygynous ants (Keller and Genoud, 1997) including *Myrmica* (Elmes and Petal, 1990; Seppä, 1994; Evans, 1996).

The succession of *M. sulcinodis* populations

In the introduction we have outlined a hypothetical scenario of how low relatedness supercolonies may develop in *M. sulcinodis* and other polygynous ants. We now return to this scenario for a more detailed discussion in light of the results of the present study.

The long distance between the three genetically differentiated *M. sulcinodis* populations suggests that they were founded by a single inseminated queen. We realise that we cannot prove this 'single-queen-foundation' scenario, but the alternative explanation based on short distance multiple colonisation followed by extinction of neighbouring, equally polygynous, populations is much less likely. The alternative explanation would automatically raise the question how these larger original populations were founded, which would ultimately require a scenario where the entire island or the area comprising both islands has once been covered by a network of interconnected supercolonies. Such a starting point, followed by gradual but large-scale extinction and substantial genetic drift, could, in theory, explain the differentiation between the three extant supercolonies on the two islands, but there are strong arguments that make this alternative a rather academic case. First, the topography of the island of Læsø indicates that the heathland habitats of the two extant supercolonies at opposite ends of the island have always been separated by zones of lower altitude without

suitable habitat for *M. sulcinodis*. Second, this scenario would predict that the allele frequencies of the two Læsø populations are more similar to each other than to the Samsø population, and this is clearly not the case as documented by comparable estimates of pairwise F_{ST} .

Assuming single-queen foundation implies that early colony expansion after colonisation would only have been possible if such single founding colonies in Læsø1 and Læsø2 produced winged reproductives of both sexes and if these young budding populations were able to survive a considerable load from diploid male production and perhaps reduced viability due to inbreeding depression. We know that such populations do exist in *M. sulcinodis*. Small and scattered populations in both the Pyrenees and Russia are characterised by a high proportion of mating among relatives leading to F_{IS} values of 0.23–0.33 and with up to 65% of the males being sterile diploids (Pedersen and Boomsma, 1997; Pedersen *et al*, 1999). Many young populations may not survive such a bottleneck, but the few that do would probably start an exponential growth phase of expansion, during which inbreeding depression would gradually diminish owing to occasionally (but still rare) immigrating males or gynes. Intermediate population structures would thus arise, characterised by moderately low relatedness and no detectable inbreeding. Examples of *M. sulcinodis* populations in this stage of development are known from Dorset, UK, where moderate relatedness of worker nest-mates and less male biased sex ratios were found in most years (Stoborough: $r_{workers-old} = 0.366 \pm 0.061$; $F_{IS} = 0.042 \pm 0.067$; $n = 44$ nests. Winfrith: $r_{workers-old} = 0.346 \pm 0.091$; $F_{IS} = -0.046 \pm 0.085$; $n = 27$ nests. Pedersen, 1997). The final stage of development, perhaps only reached in the most long-lived populations in highly stable habitats, would then be represented by the Danish island populations of Læsø and Samsø, where relatedness approaches zero and where habitat saturation is almost complete.

This hypothetical scenario would predict that marker genes of founders remain dominant in the supercolony for quite some time. This implies the expectation that the allele frequency distribution of each highly variable microsatellite locus should be characterised by two or three common alleles (representing the inseminated founding queen and multiplied during the initial population growth phase) and a series of rare alleles (representing later immigrants). The patterns in Figure 2 do indeed seem to give some support for this idea, especially for the *Myr2* locus. A similar scenario may apply to *M. rubra*, where populations are also highly variable in colony kin-structure (Seppä and Walin, 1996) and where low-relatedness supercolonies have repeatedly been found in isolated, long-term stable habitats (Pearson, 1983; Seppä and Pamilo, 1995; Seppä and Walin, 1996). In *M. rubra*, development has gone even further in that a smaller, microgyne phenotype of queen has evolved, possibly as an adaptation to the typical kin structure in these supercolonies (Pearson and Child, 1980; Elmes and Keller, 1993).

We conclude that *M. sulcinodis* and most likely also *M. rubra* are apparently able to predictably develop low-relatedness supercolonies in long-term stable isolated habitats. The mechanism behind this extreme flexibility in the expression of social traits may be convergent natural selection in each patch over a series of queen generations. However, given the limited gene flow between

populations, the likely high longevity (and thus long generation time) of queens and the identical end product of these patch-wise developments in spite of large differences in allele frequencies at neutral marker loci, it seems more likely that phenotypic plasticity with context-dependent expression of social traits is responsible for the repeated emergence of low-relatedness supercolonies in *Myrmica* ants.

A recent paper by Chapman and Bourke (2001) suggested that several pathways may lead to unicolonial structures in social insects depending on the ecological and genetical conditions. The succession of population types known for *Myrmica* ants is at least partly consistent with the 'ecological pathway' hypothesis, because unicoloniality is ultimately mediated by habitat saturation (Chapman and Bourke, 2001). This scenario allows that genetic processes interact with habitat saturation processes. Our study adds considerable detail to this scenario for a single case in point and indicates that these genetic effects are perhaps most prominent during the intermediate stages of social development towards supercolonies in *Myrmica* ants.

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