

# Genetic substructuring as a result of barriers to gene flow in urban *Rana temporaria* (common frog) populations: implications for biodiversity conservation

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The ability to maintain small populations in quasi-natural settings is an issue of considerable importance in biodiversity conservation. The genetic structure of urban common frog (*Rana temporaria*) populations was determined by allozyme electrophoresis and used to evaluate the effects of restricted intersite migration. Despite the lack of any absolute barrier to movement between ponds, substantial genetic differentiation was found between sites separated by an average of only 2.3 km. Genetic distances between these town ponds correlated positively with geographical distances and were almost twice as great as those found between rural sites separated by an average of 41 km. Measures of genetic diversity and fitness were always lowest in the town, where the degree of subpopulation differentiation ( $F_{ST} = 0.388$ ) was high. Population decline was not evident in the town, but molecular and fitness data indicated the presence of genetic drift and inbreeding depression. The long-term survival of artificially restricted populations, particularly of relatively sedentary species, may require molecular monitoring, if genetic diversity is not to be lost by chance when facets of the species niche prove to be poorly understood.

**Keywords:** allozymes, biodiversity, conservation genetics, fitness, genetic drift, *Rana temporaria*.

## Introduction

Continuing anthropogenic disruption of natural population structure has become a major issue among conservationists (Diamond, 1989; Tilman *et al.*, 1994; Kruess & Tscharntke, 1994). Fragmented populations, especially those of small size, are likely to encounter genetic drift as the major mechanism of genetic change, and this is essentially a random process of loss equivalent to sampling error (see, for example, Falconer, 1989). Such effects have been demonstrated in the natural mosaic habitat of the Ozark Mountains of the United States, where genetic isolation in local populations of a variety of ectothermic animals, with concomitant reduction in subpopulation polymorphism, resulted from historical, geographical and climatic influences (Templeton *et al.*, 1990). Local habitat fragmentation can also affect populations adversely if connectivity is disrupted, as was shown in the Glanville fritillary

(*Melitaea cinxia*), in which the probability of extinction was greatest in small, isolated habitat patches where immigration rates were comparatively low (Hanski *et al.*, 1995). Such findings have resulted in a variety of initiatives to reduce the present and predicted attrition of species numbers. In addition to the creation of nature reserves, these include: plans for wildlife corridors to facilitate colonization and link partitioned ranges (Lindenmayer & Nix, 1992; Mann & Plummer, 1995); governmental policies for biodiversity conservation (EEC, 1992); legislation for the protection of endangered species and habitats; and inauguration of reintroduction schemes (Martin, 1994). The majority of these measures, however, involve limitations on population size, immigration and emigration.

To quantify the effects of barriers to gene flow in animal populations on a local scale, we have exploited the quasi-natural environment of an urban town and used the relatively abundant and widespread common frog *Rana temporaria* (Arnold, 1995) as an example. Concerns over declines in this

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amphibian have been expressed for some time (e.g. Cooke, 1972), although not in the context of urban gardens where provision of new habitat (i.e. garden ponds) was associated with increasing numbers (Beebee, 1979). *Rana temporaria* is a semiaquatic species, which breeds communally in shallow ponds in early spring (Cooke, 1975a) and exhibits faithfulness to a summer home range (Smith, 1973; Haapanen, 1974) that is usually within 750 m of the breeding pond (Glandt, 1986). *Rana temporaria* lays an average of 1300 eggs (Cooke, 1975b) in a single spawn clump. This facilitates study, as subsamples of the progeny of each individual can be taken without confusion of identity or diminution of the population. It was hypothesized that urban Brighton (Sussex, UK) would demonstrate the effects on frog populations of isolating factors consisting of inhospitable intersite terrain created by man and imposed within a comparatively short time period. In 1799 Brighton was an isolated fishing village but today is part of an intensively developed, extensive conurbation with many transecting major roads and railways. It was estimated (Beebee, 1979) that Brighton and its adjoining coastal towns contained >7000 *R. temporaria* breeding ponds. By selecting ponds within the town with no recent history of introductions, it was possible to sample the gene pools of a range of subpopulations. Raising spawn and tadpoles in captivity permitted measures of fitness to be obtained, which could in turn be related to genetic diversity and the developmental history of Brighton. In the surrounding county, large rural

populations of the same species existed, which were also sampled and provided a control against which the urban data could be evaluated. The effect on genetic structure of barriers to gene flow is a question likely to be encountered with increasing frequency in communities worldwide.

## Methods

### *Site selection, sampling and rearing*

Thirteen breeding sites were selected to assess genetic diversity and relative fitness in their *R. temporaria* populations (Table 1). The criteria for selection were that ponds were long-established, traditional breeding sites hosting populations of *R. temporaria* and the toad *Bufo bufo* (the subject of a separate study) every year. Eight sites were urban, of which six were owned privately and two were public parks where, over many years, humans are known to have removed and deposited spawn. Five rural Sussex sites with large frog populations were also assayed. With the exception of the parks, no site had any known record of introductions within the past 35 years.

The number of breeding anurans was recorded as twice the direct count of spawn clumps made at all urban sites sampled, supplemented by rural population size estimates (this survey and Banks & Elliott, 1990). Female *R. temporaria* produce one spawn clump per year, and the operational sex ratio is generally 1:1 (Cooke, 1975a). Embryos were collected by gently removing a small portion of every

**Table 1** Details of sampling sites and populations of *Rana temporaria*

Site	Type	Sample number	Population size
Urban			
1	Private crematorium	70	140
2	Private garden	7	40
3	Private garden	7	14
4	Public park	32	64
5	Public park	33	66
6	Private garden	5	10
7	Private garden	80	160
8	Private garden	11	22
Rural			
A	Wooded heathland	100	> 1000
B	River valley marsh	20	100–1000
C	Coastal marsh	84	> 1000
D	Old parkland	100	2000–4000
E	Lake in agricultural setting	60	> 1000

Sample number is the number of spawn clumps from which samples were taken. Population size is twice the spawn clump count.

accessible spawn clump at each site, up to a maximum of 100 such samples. In all but one urban site (2, where 13/20 spawn clumps sank), every spawn clump at every site was sampled. These were stored temporarily in water from the breeding site and returned immediately to the laboratory, where a total of 100 embryos were carefully removed, by gentle separation of adjacent capsules, from the maximum possible number of these spawn subsamples. Thus, each final sample represented the fullest possible range of genotypes at the breeding location. Spawn was examined briefly under a lower power light microscope at 7–40× magnification to confirm that it was undamaged, and the embryos transferred to containers with approximately 8 L of dechlorinated tap water for culture under uniform conditions.

After resorption of gills, free-swimming larvae were maintained at densities of  $<7\text{ L}^{-1}$  and fed rabbit pellets and blanched lettuce *ad libitum*. Developmental abnormalities visible to the naked eye and survival were recorded during growth. At about Gosner stage 37 (Gosner, 1960), 24 individuals were randomly selected from each site sample and stored in liquid nitrogen ( $-196^\circ\text{C}$ ) for subsequent allozyme analysis. All harvested tadpoles and, at the termination of the experiment, all cohort survivors were examined briefly (7–40× magnification) to discern structural defects.

Survival curves were constructed from counts made during the growth period, which enabled inter-population statistical comparisons to be made. The log-rank statistic, LR (Hutchings *et al.*, 1991), which gives an approximation of the conditional log-rank test (Peto & Pike, 1973), was used in comparisons of survival. Survivorship ( $s$ ) is calculated as the number alive in a cohort at the beginning of an interval ( $i$ ), where  $S$  is total  $s$  at the same time point. LR is then calculated between populations by:

$$\text{LR} = \frac{(d_n - E_n)^2}{E_n} + \frac{(d_n - E_n)^2}{E_n} + \dots \text{ for all populations,}$$

where  $d_n$  is the sum of the interval deaths in population  $n$ , and  $E_n$  is the sum of the expected deaths in population  $n$ . Expected deaths are calculated from the product of  $(s_i/S_i)$  and  $D$ , the total of deaths occurring among populations in any  $i$ . LR was calculated similarly within each time period, using  $d_{in}$ , the number of deaths occurring between two census dates and  $e_{in}$ , the expected deaths in that interval. The log-rank statistic follows a chi-squared distribution and degrees of freedom are  $n - 1$ , where  $n$  is the number of populations. One rural site (E)

exhibited atypically high mortality and developmental abnormality of a type associated with nongenetic factors (possibly caused by parasite infection or pesticide pollution) and was therefore excluded from the analysis of survival.

#### Allozyme analysis

Protein extracts were homogenates of entire individual larvae, thawed sufficiently to permit crushing and then liquidized on ice in 150  $\mu\text{L}$  or 200  $\mu\text{L}$  (depending on tadpole size) of ice-cold 50 mM Tris-HCl (pH 8) extraction buffer. Homogenates were frozen at  $-20^\circ\text{C}$  for 15 min to ensure lysis of subcellular organelles, spun for 15 min at 8000  $g$  in a chilled MicroCentaur microcentrifuge to thaw and pellet debris, and supernatants either used immediately or re-stored in liquid nitrogen. An extensive series of trials using standard methodologies (as described, for example, in Richardson *et al.*, 1986) identified 16 enzymes from 19 loci which produced reliable banding patterns on gels during repeated assays. These loci could be resolved using three buffer systems adapted from May (1992). Buffer 1, amine-citrate (morpholine): electrode 40 mM citric acid, *N*-(3-aminopropyl morpholine) to pH 7.1, gel 1:9 dilution of electrode buffer; buffer 2, discontinuous borate: electrode 320 mM  $\text{H}_3\text{BO}_3$ , 60 mM LiOH (pH 8.1), gel 50 mM Tris, 5 mM citric acid, 10 mL  $\text{L}^{-1}$  electrode buffer (pH 8.5); buffer 3, Tris citrate: electrode 240 mM Tris, 94 mM citric acid, 50 mM NaOH (pH 7.2), gel 8 mM Tris, 3 mM citric acid, 3 mM NaOH (pH 7.7). All gels were 10.5–11.0 per cent (depending upon batch) potato starch (Sigma, UK) and included individuals from different populations as indicators of relative migration. Staining protocols were derived mainly from May (1992), except that buffers were either 100 mM sodium phosphate (pH 7) or 50 mM Tris (pH 7 and pH 8) with 100 mM KCl.

The loci used, abbreviations and buffer system(s) were: aspartate aminotransferase cytosolic, *Aat-C*, 2 and mitochondrial, *Aat-M*, 1; aconitase, *Acon*, 3; adenosine deaminase, *Ada*, 2; alcohol dehydrogenase, *Adh*, 2, 3 or 1; creatine kinase, *Ck*, 3; nonspecific esterase, *Est*, 2 loci, 2; fumarate hydratase, *Fum*, 2; glutathione reductase, *Gr*, 3 or 1; isocitrate dehydrogenase, *Idh*, 3; lactate dehydrogenase, *Ldh*, 2 loci, 3; mannose-P-isomerase, *Mpi*, 3 or 1; nucleoside phosphorylase, *Np*, 2; peptidase A, *Pep-A* (substrate glycyl-leucine), 2; peptidase B, *Pep-B* (substrate leucyl-glycyl-glycine), 2; phosphoglucuronate dehydrogenase, *Pgd*, 2; phosphoglucosylmutase, *Pgm*, 2.



### Genetic analyses

Genotype data from biochemical studies were analysed using the computer programs BIOSYS-1 release 1.7 (Swofford & Selander, 1981), FSTAT version 1.2 (Goudet, 1995), GENEPOP version 1.2 (Raymond & Rousset, 1995) and PHYLIP (Felsenstein, 1993). Standard statistical analyses were performed using MINITAB release 8. Conformance of loci to the Hardy–Weinberg equilibrium was assessed by chi-squared tests using Yates's correction for degrees of freedom (d.f.) = 1, and, where d.f. > 1 and the low values typical of genotype frequencies occurred, a method analogous to the Fisher exact test was employed (Swofford & Selander, 1981, BIOSYS-1). Population subdivision was assessed by  $F$ -statistics and  $Nm$ , equivalent to the number of migrants per generation (Wright, 1987). Loci were used in further analysis only when polymorphism was such that a significant difference in allele frequencies existed among populations. Genetic distances were calculated through PHYLIP using Cavalli-Sforza & Edwards (1967) chord distance with a divisor equal to the degrees of freedom (the number of alleles minus 1). This genetic distance assumes that gene frequency change occurs by drift alone without mutation but not that population sizes have remained constant (Felsenstein, 1993), a model appropriate to the short chronological period involved in the development of the urban environment. Nei's standard genetic distance (Nei, 1987) was also calculated to permit comparison with other studies. The PHYLIP CONTML restricted maximum likelihood method, which is based on the models of Cavalli-Sforza & Edwards (1967) and Felsenstein (1993), was used to construct trees of relatedness between populations. Consensus trees were obtained by bootstrapping 500 times through the PHYLIP CONSENSE program. The significance of the association between genetic and geographical distance was assessed using a Mantel test (GENEPOP), as the interdependence of matrix data values invalidates significance testing of the product–moment correlation  $r$ .  $r$  is given, however, as a convenient descriptive statistic of the linear plots of these distances.

To facilitate evaluation of urban Brighton genetic data, an examination of the developmental history of the town was made using Ordnance Survey (OS) maps and local population statistics. Intersite urban distances were calculated by measurement from large-scale OS maps, which allowed precise location of street boundaries, then plotted onto a 1994 edition OS Pathfinder (1307, TQ20-30) 4 cm:1 km

map. 'Major roads' reported were those defined as >4 m wide by this OS map edition.

## Results

### Genetic analysis

Although no formal crosses were carried out, the subunit structures of the enzymes assayed are well characterized as also are the numbers of genetic loci controlling their expression (Noltmann, 1972; Webb, 1984). Without exception, the banding patterns observed in this study were identical to predictions from the above investigations and to those previously carried out on anurans in general (Guttman, 1985) and on *Rana*, including *R. temporaria*, in particular (Nishioka *et al.*, 1992). The mean number of alleles per locus, the mean observed heterozygosity ( $H$ ) per locus and the percentage of these loci that were polymorphic (per cent  $P_m$ ) in each *R. temporaria* population are given in Table 2. The polymorphic loci are given as the percentage actually observed and at the 95 per cent criterion (Wright, 1987).

The urban and rural average number of alleles per locus,  $H$  and  $P_m$ , did not differ significantly, although in all instances genetic diversity was greater in rural than in urban populations. The average conformance of loci to the Hardy–Weinberg equilibrium in urban populations was approximately 58 per cent and in rural populations approximately 69 per cent (after Bonferroni correction for multiple tests of hypothesis; Samuels, 1989, p. 505), and the deviations were always deficiencies of heterozygotes.  $F$ -statistics and  $Nm$  estimates averaged across all loci are given in Table 3. All three  $F$ -statistics in all three population groupings were significantly greater than zero ( $P = 0.001$ ; Goudet, 1995). The relative level of inbreeding within populations ( $F_{IS}$ ) was slightly lower in rural locations than in urban locations. When the individual was considered relative to the expected heterozygosity of pooled sites ( $F_{IT}$ ), that difference was greater.  $F_{IT}$  reflects both drift and nonrandom mating (Hartl & Clark, 1989).  $F_{ST}$ , the measure of population subdivision, was more than twice as high in urban than in rural sites.

Urban and rural  $F_{ST}$  (median: urban 0.35, rural 0.11) and  $Nm$  (median: urban 0.48, rural 2.13) values were significantly different at the 5 per cent level when judged by Mann–Whitney tests.

Geographical and genetic distances between sites are given in Table 4. The Cavalli-Sforza & Edwards (1967) chord genetic distance correlated positively with distance in km between urban sites ( $r = 0.50$ ,  $P$

**Table 2** Genetic summary statistics for the *Rana temporaria* populations

Site	Mean alleles per locus	±SE	Mean <i>H</i> per locus	±SE	<i>P<sub>m</sub></i> (%)	95% <i>P<sub>m</sub></i>
1	1.53	0.19	0.036	0.023	36.8	21.1
2	1.84	0.26	0.077	0.031	47.4	42.1
3	1.37	0.17	0.042	0.027	26.3	26.3
4	1.68	0.19	0.033	0.013	52.6	31.6
5	1.74	0.23	0.056	0.028	42.1	42.1
6	1.74	0.23	0.043	0.017	47.4	36.8
7	1.68	0.28	0.076	0.042	31.6	31.6
8	1.74	0.24	0.075	0.032	47.4	36.8
A	1.63	0.22	0.050	0.028	47.4	31.6
B	2.11	0.36	0.106	0.041	57.9	36.8
C	1.53	0.19	0.063	0.037	36.8	26.3
D	2.11	0.35	0.072	0.040	57.9	36.8
E	1.58	0.19	0.058	0.028	36.8	36.8
All	1.71		0.061		44.3	33.6
Urban	1.65		0.055		41.5	33.6
Rural	1.83		0.073		50.0	33.7

*H*, heterozygosity; *P<sub>m</sub>*, percentage of polymorphic loci, total or at the 95 per cent level (see Methods).

(Mantel) = 0.023) and, with town parks excluded (where genetic mixing was believed to take place), the relationship was more pronounced ( $r = 0.71$ ,  $P = 0.002$ ). In contrast, comparison of rural genetic distance and geographical distance showed no pattern of association ( $r = 0.032$ ) and this changed little when urban populations were averaged and included as a single Brighton (Br) site ( $r = 0.095$ ). The difference between the urban (average = 0.2320) and rural (average = 0.1158) genetic distance means was highly significant ( $t = 5.56$ ,  $P < 0.0001$ ). Nei's genetic distance (data not shown) correlated strongly with the Cavalli-Sforza chord measure ( $r = 0.967$ ,  $P < 0.001$ ) and also showed a similar, albeit weaker, correlation with intersite distances in kilometres between urban sites but not rural ones. Nei's genetic distances between urban sites (average = 0.105) were also significantly higher than the intersite rural distances (average = 0.036;  $t = 5.78$ ,  $P < 0.0001$ ).

**Table 3** Summary *F*-statistics and gene flow estimates for *Rana temporaria* populations

Site type	<i>F<sub>IS</sub></i>	<i>F<sub>IT</sub></i>	<i>F<sub>ST</sub></i>	<i>N<sub>m</sub></i>
All	0.565	0.706	0.324	0.522
Urban	0.582	0.745	0.388	0.394
Rural	0.539	0.606	0.145	1.474

The association between genetic and geographical distances within Brighton was investigated further by examination of urban boundary development in the town over time from OS maps. The number of major roads and railways intersecting the intersite transects was also determined, as roads are known to be significant barriers to migrating amphibians (Cooke, 1972; Fahrig *et al.*, 1995). Weighting the distance in kilometres by  $1 \times$  the proportion of the transect developed post-1932 improved correlation with the Cavalli-Sforza chord genetic distance slightly, to  $r = 0.58$ ,  $P$  (Mantel) = 0.004 with town parks and  $r = 0.74$ ,  $P$  (Mantel) = 0.001 without them. Road numbers showed a slight positive association with genetic distance, but the data points were very scattered and, as the urban study area is almost entirely developed, they correlated linearly with distance in any case ( $r = 0.88$ ,  $P < 0.001$ ).

Figure 1 shows a map of the developmental history of Brighton and the urban study sites, together with a CONTML tree of the genetic relationships of *R. temporaria* breeding ponds with approximately proportional branch lengths. The CONTML tree was strongly supported by bootstrapping (500  $\times$ ), with all clusters except 7–8 occurring in 87 per cent of cases. The greatest area of uncertainty (the node linking sites 7–8, 62 per cent occurrence) resulted from rearrangement of the order of the three terminal sites. The correlation between

**Table 4** Genetic and geographical distances matrix for *Rana temporaria* populations

## (a) Urban

Site	1	2	3	4	5	6	7	8
1		1.8	2.7	3.6	3.3	4.4	3.5	4.1
2	0.1712		0.9	1.8	2.6	2.6	2.3	2.7
3	0.2459	0.1291		1.0	2.7	1.8	2.0	2.2
4	0.1955	0.2561	0.2820		2.7	0.8	1.7	1.6
5	0.2314	0.2585	0.2541	0.0614		2.9	1.1	1.7
6	0.4100	0.3705	0.4078	0.1987	0.1637		1.8	1.4
7	0.3221	0.2960	0.3213	0.1206	0.0817	0.1344		0.7
8	0.3420	0.3089	0.3544	0.1848	0.1292	0.1886	0.0752	

## (b) Rural + Brighton (mean of eight urban sites)

Site	A	B	C	D	E	Br
A		22	37	73	34	29
B	0.1417		27	66	24	11
C	0.0422	0.1460		93	52	36
D	0.1297	0.1550	0.1113		42	57
E	0.0726	0.1563	0.0789	0.1470		16
Br	0.1183	0.1323	0.0939	0.0929	0.1182	

Cavalli-Sforza (1967) chord genetic distances (lower triangles) and geographical distances in km (upper triangles).

geographical and genetic (tree) locations of the study sites was striking.

Although no relationship between genetic distance and distance in kilometres was found for the five rural sites, they were also analysed by CONTML. The resultant tree (not shown) maintained its branching structure after bootstrapping but clusters had low percentage reproducibility ( $\leq 57$  per cent).

*Fitness assessments*

The survivorship curves for urban and rural *R. temporaria* larvae cultured under controlled conditions are given in Fig. 2.

Survivorship among the 12 sites, as tested by the log-rank statistic (LR), was not uniform, either within the time intervals examined or between cohorts (Table 5).

The highest levels of significance were associated with early development, when the greatest mortality occurred. Clustering deaths during all four census intervals as either urban or rural showed significant overall differences between the two groups (Mann-Whitney test,  $P = 0.042$ ) with median urban deaths in any census interval equal to 5 and median rural deaths equal to 2. The difference between the

number of developmental abnormalities observed in urban as opposed to rural tadpoles was also significant ( $P = 0.042$ ) when evaluated by the Mann-Whitney test (urban median = 8.00, rural median = 3.50). A statistically significant negative correlation ( $r = -0.80$ ,  $P = 0.002$ ) existed between survival at day 80 and  $\log_{10}$  abnormalities, although these factors may not be truly independent: abnormal tadpoles may have died, contributing to lowered survival. Similarly, malformations, such as tail-tip kinks, in early larvae may have become invisible as development proceeded.

**Discussion**

Urban populations of common frogs exhibited higher degrees of genetic substructuring and lower levels of average genetic diversity than their rural counterparts, and also experienced higher levels of mortality and developmental abnormality when larval growth was compared in controlled conditions. It is generally believed that breakdown of developmental homeostasis is an indicator of inbreeding (Mitton & Grant, 1984) and the phenomenon of developmental abnormality has a long association with lack of genetic variety. The correlation of



genetic distance with small-scale geographical distance within the town suggests that, despite the lack of absolute barriers to migration, these populations have become differentiated genetically by drift. Genetic distances between urban *R. temporaria* ponds separated by 0.7–4.4 km were, on average, twice as great as those between the much more widely dispersed rural sites (mean separation 41.3 km). Historical and ethological considerations suggest that the causative isolating factor was the increasingly inhospitable terrain of the urban environment, as the association between genetic and geographical distance was enhanced if weighted for recent development. This finding was reflected in the high degree of urban subpopulation differentiation, which resulted in  $Nm$  values averaging 0.394, considerably less than the value of 1, below which allele frequency changes mediated by drift are prob-

able (see, for example, Slatkin, 1985). In comparison, mean rural  $Nm$  was 1.474. The distinctive relationship between the urban genetic tree and the geographical locations emphasized the strength of the isolating factors. The juxtaposition on the CONTML tree of the admixed populations of urban parks, which were distant geographically, provided supporting evidence for the validity of the tree branching structure.

This investigation indicates that the impact of habitat change on the genetic diversity of a species can be rapid and superficially undetectable at the phenotypical level. Amphibians appear to be well adapted to the urban habitat, are locally abundant and will colonize nearby ponds, but there appears to be insufficient progressive migration to maintain panmixia over even relatively short distances. What might be considered a metapopulation (Hanski,

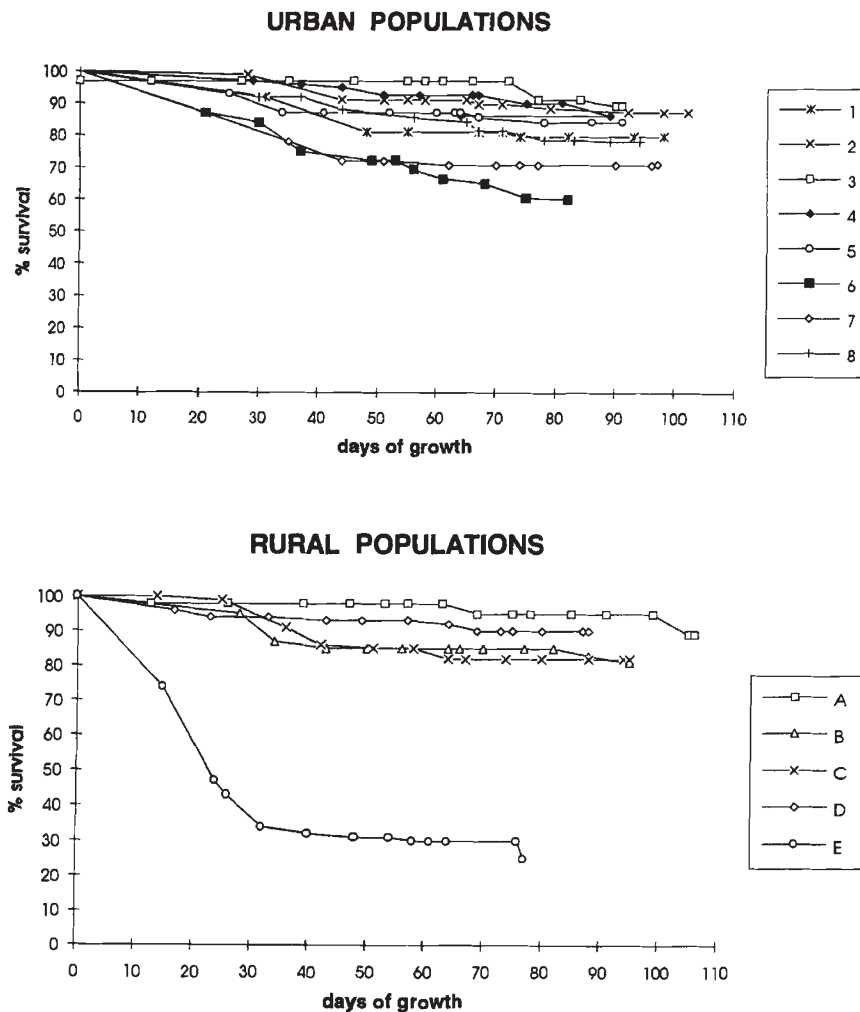


**Fig. 1** Geographical and genetic relatedness of populations of *Rana temporaria*. Top: urban Brighton with site numbers and boundaries of development at the given dates. Major roads are shown by solid lines. TQ28–34 gridline = 6 km. Bottom: CONTML tree of genetic relatedness of populations in the urban sampling sites from allozyme allele frequencies.

1991; Hanski & Gilpin, 1991) because of the very large number of urban ponds with opportunity for random extinction and recolonization, appears instead to be a collection of small, relatively independent populations becoming progressively more isolated in genetic space as barrier strength increases. It therefore seems that the urban environment has prevented frog populations from exchanging sufficient individuals to avoid genetic drift, that this is a continuing process despite the existence of many suitable ponds in the area and that this random loss of alleles has resulted in fitness reduction by inbreeding depression. It is of some interest in this context that disease epidemics in urban frog populations, perhaps related to reduced fitness, have received considerable attention recently (e.g. Cunningham *et al.*, 1993). On the other hand, translocations of frogs from garden environments to start new populations have proved successful (e.g. Cooke

& Oldham, 1995), perhaps because spawn from multiple sites was pooled for the purpose.

Table 6 shows some comparative genetic data for amphibian taxa. Genetic variation of Brighton and Sussex *R. temporaria* was in a range similar to that observed among 12 *Rana* species (Nishioka *et al.*, 1992). In Germany, 20 rural populations of *R. temporaria* (Reh & Seitz, 1990) had higher overall genetic diversity and exhibited significant associations of genetic distance with intersite motorways and railways. *H* was, however, comparatively low in Brighton *R. temporaria* and, in urban populations, *H* and  $F_{ST}$  were similar to those reported in salamanders. Mobility in the Amphibia is inherently restricted by a requirement for a moist environment, and high values of  $F_{ST}$  have been reported regularly, for example in a group of 22 salamanders (Larson *et al.*, 1984) with mean  $F_{ST} = 0.53$  and consequently low estimates of migration, *Nm*. It was considered



**Fig. 2** Survival during growth in controlled conditions of *Rana temporaria* larvae. The keys refer to individual sites.



by these authors that the relatively high levels of  $F_{ST}$  could be explained by a history of glacial isolation and recolonization, coupled with the inherently low mobility of small amphibians.

The imposed isolation of common frogs in Brighton is typical of many other species in the UK. In present-day England, >40 per cent of the land area is converted to intensive human use, with about 10 per cent urban and the remainder arable, and a

further third exists as managed grassland (Barr *et al.*, 1993). Such intensive land management imposes discontinuity on the remaining habitat (Thornton & Kite, 1990) as segments of unsuitable terrain (urban, industrial and agricultural) are interspersed among the undeveloped, seminatural areas, with resultant fragmentation of population connectivity.

The majority of natural population losses result from extensive habitat change caused by the recent rapid increase in the human population of the world, from approximately  $0.7 \times 10^9$  in 1700 to approximately  $5.0 \times 10^9$  at present (Demeny, 1990; Kates *et al.*, 1990). Widespread concern exists over the extinction of species (Ehrlich & Wilson, 1991), with widely varying estimates of rates and numbers of losses, including the suggestion (May *et al.*, 1995) that the average species lifetime may have been reduced from  $10^6$  years to 300 years. Internationally, habitat destruction has been highlighted in tropical rainforests, where recent estimates of annual rates of deforestation in the Brazilian Amazon approached  $2.0 \times 10^6$  ha (Skole *et al.*, 1994).

It is apparent that careful consideration of all facets of a species niche and lifestyle will be necessary if the genetic diversity of populations is to be maintained in natural communities. A wildlife corridor

**Table 5** Survivorship differences between *Rana temporaria* larvae from different populations

Census period (days)	Log-rank statistic	Significance ( <i>P</i> ) d.f. = 11
(a) Between populations at a single census time		
0–24	43.486	0.001
25–39	41.293	0.001
40–59	28.701	0.01
60–79	23.156	0.05
(b) Between total deaths in cohorts		
0–79	70.204	0.001

Data are from 12 urban and rural populations. Probabilities are of departures from expected values.

**Table 6** Comparative amphibian genetic diversity statistics

Key	Subject	Mean alleles per locus	Mean <i>H</i> per locus	Mean $P_m$ (%)	$F_{ST}$
1	<i>Bufo</i> spp.	1.4	0.115	39*	—
2	<i>B. japonicus</i>	1.5	0.107	36*	0.244
3	<i>B. bufo</i>	1.2	0.023	18	0.487
4	<i>Rana</i> spp.	1.4	0.101	33*	0.289
5	<i>R. temporaria</i>	2.1	0.170	66	—
6	<i>R. temporaria</i>	1.7	0.061	44	0.324
7	<i>Salamandra salamandra</i>	1.5	0.078	36	0.403
8	<i>Triturus italicus</i>	1.3	0–0.09	19	0.448

1. *Bufo* spp. (Nishioka *et al.*, 1990). Twenty-three populations of seven bufonid species, including six subspecies of *B. japonicus*.

2. *Bufo japonicus* (Kawamura *et al.*, 1990). Five subspecies in 40 Japanese populations.

3. *Bufo bufo* (Hitchings & Beebee, in preparation). Data from the urban and rural locations of this report.

4. *Rana* spp. (Nishioka *et al.*, 1992). Thirty populations of 12 ranid species.

5. *Rana temporaria* (Reh & Seitz, 1990). Twenty rural German populations of 10–360 individuals.

6. This study, overall averages.

7. *Salamandra salamandra* (Alcobendas *et al.*, 1996). Two conspecific lineages and 13 populations.

8. Italian newt *Triturus italicus* (Ragghianti & Wake, 1986). Eleven populations in southern Italy.

will remain a barrier if it does not provide habitat for species that cannot simply walk straight through. Using amphibians as an example, ponds and suitable non-breeding season territory need to be within an uninterrupted territorial range if they are to function as staging posts in a sequential process of colonization. It will be most important to monitor genetic diversity, at least in a representative group of relatively sedentary taxa, to assess changes that might commence because an essential aspect of the species requirements has been overlooked or underestimated.

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