

Importance of dispersal for the expansion of a Eurasian lynx *Lynx lynx* population in a fragmented landscape

Fridolin Zimmermann, Christine Breitenmoser-Würsten and Urs Breitenmoser

Abstract Dispersal allows recolonization of previous areas of habitat following severe depression of a population but the significance of this is not clear in felids. There is little evidence to support the general belief that subadult felids will colonize new areas, although this is a crucial assumption in reintroduction or recovery projects. Eurasian lynx *Lynx lynx* were reintroduced into the Swiss Alps and have subsequently spread over part of their potential range but the expansion halted in the mid 1980s. We postulated that high lynx densities would lead to an expansion of the population, and to assess the potential of this population to expand we compared the dispersal characteristics of 22 subadults from the north-west Swiss Alps, where an increase in lynx abundance occurred from 1995 onwards, to 17 individuals from the Jura Mountains, an area with a lower lynx density.

Dispersal data came mainly from radio-telemetry. Dispersal rates and distances for subadults that completed dispersal were lower in the north-west Swiss Alps than in the Jura Mountains. In general, subadults exhibited little ability to cross major barriers such as highways. The hypothesis that high density alone will foster the expansion of the population was therefore not confirmed. This has consequences for the reintroduction and recovery of carnivores in fragmented landscapes. To establish only one strong source population may not be an optimal strategy, and population nuclei should therefore be founded in several neighbouring patches.

Keywords Density dependence, dispersal, fragmented landscape, *Lynx lynx*, population expansion, Switzerland.

Introduction

Natal dispersal, the movement of subadult animals from their natal range to the place where they will breed (Howard, 1960), is an important mechanism in the colonization of new areas. Individuals compete for local resources, and dispersers need to obtain access to unexploited resources. In a saturated population such resources are at the edge of or outside the colonized area. Hence, dispersal regulates abundance by expanding distribution (Dieckmann *et al.*, 1999). Dispersal also allows recolonization of former habitats after a population decrease (Lubina & Levin, 1988), and this is occurring in large carnivores in Europe and North America. Brown bears *Ursus arctos*, nearly extinct in the early 20th century, have recolonized large parts of

Scandinavia (Swenson *et al.*, 1995). The range outside the core areas was occupied by young bears, predominantly by males 2–4 years old, the age of most active dispersal (Swenson *et al.*, 1998). Wolves *Canis lupus* from the Italian Apennine population recolonized the Alps, where the species had been eradicated in the 19th century (Breitenmoser, 1998). Recolonization of former range by dispersing wolves has also been observed in Minnesota (male-biased; Mech, 1987) and in the central Rocky Mountains (both sexes; Boyd & Pletscher, 1999).

The significance of dispersal for the spread of a population is less clear in felids. Long-range dispersal has been described in Eurasian lynx *Lynx lynx* (J. Linnell, pers. comm.), in Canadian lynx *Lynx canadensis* (reviewed in Mowat *et al.*, 2000) and in puma *Puma concolor* (Sweaner *et al.*, 2000) but these were animals dispersing between or within extant populations. There is little evidence that dispersing subadult felids will colonize new territory, although this is a crucial assumption in reintroduction or recovery projects.

Eurasian lynx were reintroduced into the Swiss Alps in the early 1970s and have subsequently spread over the north-west of the Alpine arc (Breitenmoser *et al.*, 1998). The expansion slowed down in the mid 1980s, despite the availability of suitable unoccupied habitat, and in the 1990s little or no evidence for lynx presence

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was found in some regions that had been previously colonized (Molinari-Jobin *et al.*, 2001). After 1995 an increase in lynx abundance in the north-west Swiss Alps resulted in a controversy about the return of this species. Hunters, sheep breeders and the regional authorities demanded that the lynx population in this area be reduced (Breitenmoser *et al.*, 1999). Such permission was, however, not given at that time, as the policy of the Federal Office for the Environment was to have lynx distributed throughout the Alps, and there was a general belief that the population pressure in the north-west Swiss Alps would further the expansion of the population.

The hypothesis that population pressure will result in an expansion of the occupied area (Hell, 1961) implies two assumptions: the population expands through natal dispersal, and dispersal is positively density dependent. Therefore both the rate and the mean distance of dispersal are greater when lynx are abundant. These assumptions have consequences for the colonization of new or former areas, and hence for the design of reintroduction projects. Furthermore, the nature of any dispersal is determined by resource availability, landscape (habitat and topography), and the social structure of the species. To assess dispersal in the north-west Swiss Alps during a period of peak lynx density (1.4–1.5 resident lynx per 100 km²; Breitenmoser-Würsten *et al.*, 2001) we compared it with data from an earlier, similar study (Zimmermann, 1998) in the Jura Mountains where lynx density was low to average (0.7–0.8 resident lynx per 100 km²; Breitenmoser-Würsten *et al.*, in press). To assess the potential of the population of the north-west Swiss Alps to expand we examined the dispersal rates, distances, directions, habitat and linear barriers crossed by dispersers compared to the Jura Mountains and postulated that (1) rate and distance of dispersal should be greater in the higher density population, and (2) most subadults would leave the north-west Swiss Alps and settle in neighbouring areas. Hence we assumed that the monitoring system for lynx in Switzerland (Capt *et al.*, 1998) would reveal an increase of lynx presence, after a time lag, in the areas adjacent to the north-west Swiss Alps.

Materials and Methods

Study Areas

The north-west Swiss Alps are a 2,800 km² area more or less isolated from the rest of the Alps and fragmented both by natural and artificial barriers (for details see Zimmermann, 2004; Fig. 1). Roe deer *Capreolus capreolus*, lynx's main prey (Breitenmoser & Haller, 1987), were locally depressed during the peak of lynx density at the

end of the 1990s. The number of roe deer harvested per 1 km² forest, assumed to reflect roe deer abundance, has continuously decreased in the north-west Swiss Alps since 1994 (Fig. 2). The overall resident lynx density in the north-west Swiss Alps, based on the distribution of radio-collared animals and information about additional, untagged individuals from camera-trapping, was estimated at 1.4–1.5 per 100 km² (Breitenmoser-Würsten *et al.*, 2001). Areas with lynx adjacent to the north-west Swiss Alps are Valais to the south, and the west central Swiss Alps to the east.

The Jura Mountains are a secondary limestone mountain chain forming the north-west border of Switzerland with France (for details see Breitenmoser *et al.*, 1993a; Fig. 1). They are less fragmented than the Alps, forming a block of contiguous suitable habitat of 6,670 km² (Zimmermann & Breitenmoser, in press). In contrast to the north-west Swiss Alps, prey base was not a limiting factor during the study period. The number of roe deer killed per 1 km² of forest increased continuously since 1990 in the southern part of the Jura Mountains (Fig. 2), as well as in France (Stahl *et al.*, 2001). The overall lynx density in the main study area remained fairly constant over the whole study period and was estimated at 0.7–0.8 residents per 100 km² (Breitenmoser-Würsten *et al.*, in press).

The study area in each region was defined as the area formed by the 1 km buffered 100% minimum convex polygon (MCP) of all adult resident females from which juveniles were caught and observed during dispersal (Fig. 1).

Field study

From 1988 to 2001 dispersal characteristics (Table 1) were obtained for 22 and 17 lynx in the north-west Swiss Alps and Jura Mountains, respectively. Dispersal data came mainly from radio-telemetry; additional information was available from kittens tagged at the den and later live-trapped or photographed by a camera trap. Details of capture, handling, radio-telemetry and number of locations are in Zimmermann *et al.* (2005). Only radio fixes with an accuracy of 1 km² or better were considered. We located dispersing lynx almost every day when they moved through new terrain, and at least every week once home ranges were established for >1 month. After independence subadults were considered dispersers when they established a home range that overlapped by ≤5% of their natal (maternal) home range (based on a 90% MCP) or were last located outside their natal area for those that either died or lost contact before forming a home range (Sweaner *et al.*, 2000). Percentage overlap of total ranges and home ranges between animals A and B was calculated as

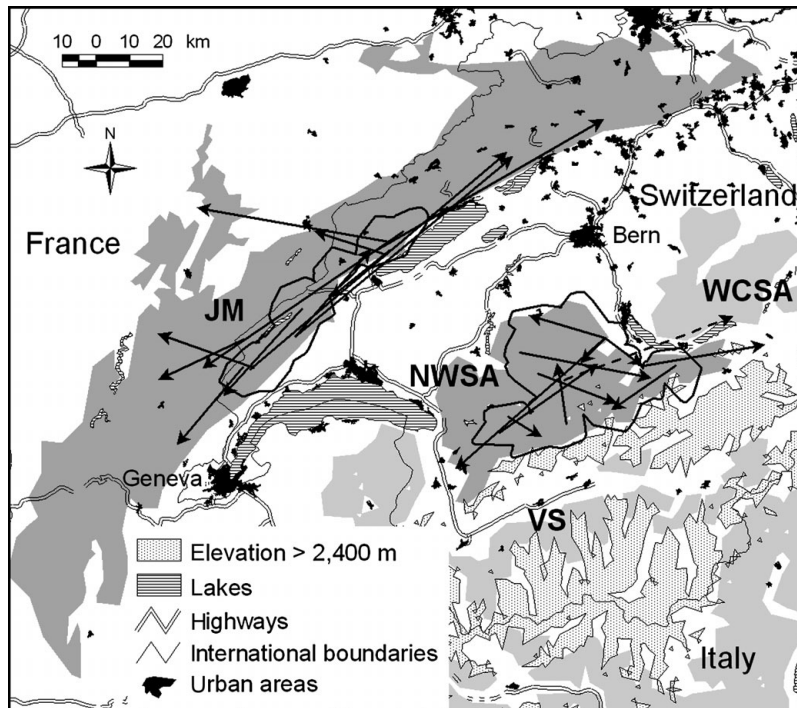


Fig. 1 Location of the study sites in Switzerland. Suitable lynx habitat patches in the north-west Swiss Alps (NWSA) and the Jura Mountains (JM) are dark grey; the adjoining patches of Valais (VS) and west central Swiss Alps (WCSA) are light grey. Thick black lines show the main study areas (see text for further details). Arrows indicate dispersal directions, distances and endpoints for 12 lynx in NWSA and 12 lynx in JM. Movements are shown as straight lines from a lynx's natal home range centre or its capture site to its independent home range centre, mortality site or last location. All subadults settled in their natal habitat patch, except one male that left NWSA (dashed line).

$$\sqrt{(\text{overlapAB}/\text{rangeA}) * (\text{overlapAB}/\text{rangeB})}$$

All other independent progeny (those establishing home ranges with >5% overlap with their natal range) were considered philopatric (Ph). Dispersal began when a subadult made its first move outside its natal home range without returning. Dispersers were classified as: (1) dispersers (Di) that most likely completed their natal dispersal and exhibited 6 months of site fidelity suggestive of home range establishment and/or reached sexual maturity, or (2) failed dispersers (fD) that exhibited <6 months site fidelity and/or died before they established a home range. Lynx normally reach sexual maturity at 2.75 years for males and 1.75 years for females. In one study (Kvam, 1991) some males and almost 50% of females reached sexual maturity one year earlier but we never observed such early maturity. A limit of 6 months was chosen for site fidelity because all dispersers, with the exception of one male that stayed for 200 days, left their transient home range after 72–114 days (median 113 days, $n = 5$; Zimmermann *et al.*, 2005).

Data analysis

The dispersal direction was defined as the vector from the centre (arithmetic mean of x and y coordinates of all radio fixes) of the natal home range to the centre of

the independent home range. Directional data were transformed to unimodal data and subjected to Rayleigh's test (Zar, 1984) to examine whether dispersal directions were distributed uniformly or not. Centroid distance (CD; Fig. 3) was the distance from the centre of a progeny's natal home range to the centre of its independent home range. When complete dispersal information was not available, dispersal distances and directions were calculated based on one of the following combinations: centre of natal

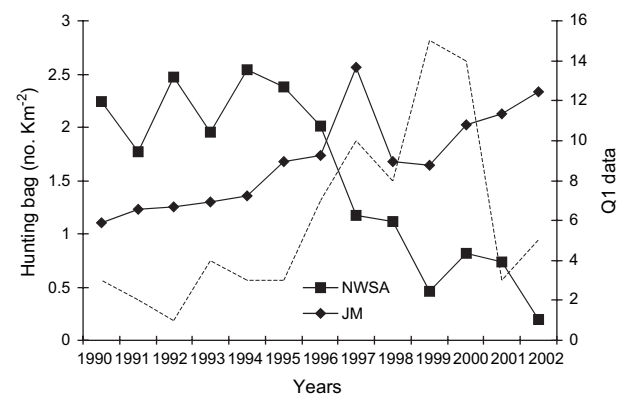


Fig. 2 Roe deer hunting bag (number killed per km² forest) from 1990 to 2002 in the north-west Swiss Alps (NWSA) and the Jura Mountains (JM). The hunting quotas are fixed each year according to estimates of population size and trend. The broken line shows the evolution of the Q1 (see text for details) signs of lynx presence in the north-west Swiss Alps.

Table 1 Characteristics and fate of juvenile lynx (M, male; F, female; FB, females ear-tagged as juveniles) followed in the north-west Swiss Alps and the Jura Mountains (Zimmermann *et al.*, 2005), with the maternal lines, date of first observation, method used to track each individual, and the fate of each individual.

Lynx	Kitten of female	Date of first observation	Method ¹	Date of last observation	Fate ²
North-west Swiss Alps					
M18	F37	22/01/1997	rt, ct	16/05/1999	Alive, [reproduced]
M20	F30	25/03/1997	rt, ct	02/12/1998	Alive
M24	F47	24/02/1998	rt, ct	27/01/2004	Alive, reproduced
M25	F33	28/06/1997	den, rt	14/09/2000	Alive, [reproduced]
M28		30/01/1999	rt	12/02/1999	Died, starvation
M29	F34	25/06/1998	den, rt, ct	07/05/2003	Alive, [reproduced]
M30		05/03/1999	rt	21/09/1999	Died, disease
M31	F52	08/12/1998	ct, rt	10/08/2000	Alive
M35	F35	31/10/1998	ct, rt, ct	12/02/2004	Alive, [reproduced]
F31	F32	06/01/1997	rt, ct	26/09/1998	Alive, [reproduced]
F33 ³		14/01/1997	rt	23/07/1998	Alive, reproduced
F40		13/03/1997	rt	20/03/1998	Unknown
F42 ³		05/04/1997	rt,ct	11/01/2004	Alive, reproduced
F46 ³		15/11/1997	rt	20/01/1998	Died, car accident
F48	F38	05/07/1997	den, rt	18/05/1998	Unknown
F49	F34	23/06/1997	den, rt	27/08/1999	Alive, [reproduced]
F50	F37	13/03/1998	rt	19/05/1998	Unknown
F54 ³		14/08/1999	rt	23/06/2000	Alive, [reproduced]
F56	F32	08/02/2000	rt	25/02/2000	Unknown
F57 ³		08/05/2000	rt	10/03/2001	Alive
FB22	F34	25/06/1998	den, ct, cr	18/10/2001	Alive, [reproduced]
FB44	F34	11/12/2001	ct	01/08/2003	Alive, [reproduced]
Jura Mountains					
M11	F21	19/06/1993	den, rt, cr	08/02/2002	Alive, reproduced
M13	F18	18/06/1993	den, rt	27/06/1995	Alive, [reproduced]
M14	F21	31/12/1995	rt	26/02/1998	Alive, [reproduced]
M15	F18	23/06/1995	den, rt	21/09/1996	Died, illegal killing
M16	F11	08/03/1990	rt	19/06/1990	Died, disease
F12	F11	20/04/1988	rt	20/12/1988	Died, illegal killing
F13	F11	23/03/1989	rt	17/08/1989	Died, disease
F17	F14	25/03/1990	rt	23/05/1990	Died, car accident
F19	F18	04/03/1991	rt	05/05/1991	Unknown
F20	F11	16/03/1991	rt	??/11/1995	Alive, reproduced
F22	F18	04/03/1992	rt	25/11/1994	Alive, reproduced
F23	F15	14/03/1992	rt	15/02/1995	Alive, reproduced
F25	F30	20/12/1995	rt	02/07/1996	Unknown
F26	F18	23/06/1995	den, rt, ct	08/01/2003	Alive, reproduced
F27	F24	30/06/1995	den, rt	27/04/1996	Unknown
F28	F24	27/06/1995	den, rt	04/03/1996	Unknown
F36	F29	04/03/1997	rt	24/06/1998	Alive, [reproduced]

¹Den, individual ear-tagged as kitten; rt, radio-telemetry; ct, camera-trap; cr, carcass retrieved

²Alive, survived the full year after separation from mother; reproduced, strong evidence from genetic analyses or field observations that individuals have reproduced; [reproduced], lynx reached sexual maturity (according to definition of Kvam, 1991) but there was no proof of reproduction

³First observed as subadults after separation from mother

home range to mortality site or last location, or capture site to centre of independent home range, mortality site or last location. The dispersal distance was also expressed relatively as the number of sex-specific home ranges crossed during dispersal. The size of male and female home ranges, respectively, was calculated as the diameter (d) of a circle with an area equal to the average home range size (HR) for adult male and female lynx in

the study areas (north-west Swiss Alps: Breitenmoser-Würsten *et al.*, 2001; Jura Mountains: Breitenmoser-Würsten *et al.*, in press):

$$d = 2\sqrt{HR/\pi}$$

The total dispersal distance (TD, Fig. 3) was the sum of distances between consecutive locations (only one radio-telemetry fix per day considered) of subadults

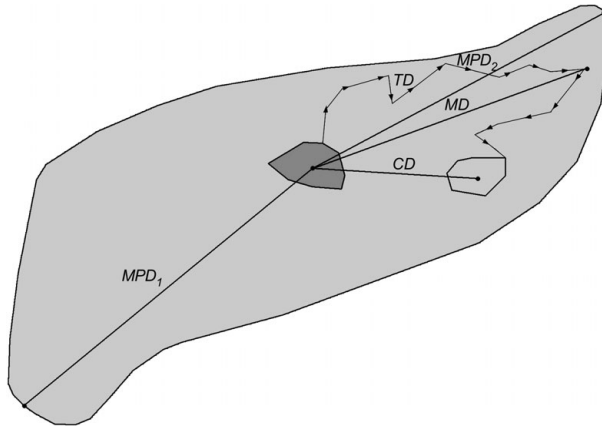


Fig. 3 Definitions of the measured dispersal distances of subadult lynx. Centroid distance (CD) is distance from arithmetic centre of natal home range (solid grey minimum convex polygon, MCP) to arithmetic centre of independent home range (light grey MCP). Total distance (TD) is sum of distances between consecutive locations during dispersal (measured from the point the subadult left the natal home range to the point when it entered its independent home range); maximum distance (MD) is longest distance a dispersing lynx was ever located from the centroid of its natal range; maximum possible distance 1 (MPD_1) is distance from the centroid of the natal range to the farthest edge of good lynx habitat (light grey area); maximum possible distance 2 (MPD_2) is distance from the centre of the natal home range to the most distant good lynx habitat edge in the initial dispersal direction.

during their dispersal. The measures were taken from the point the subadult left the natal home range to the point when entering its subsequent home range. When subadults established more than one home range, the total distances between the ranges were summed. The maximum dispersal distance (MD; Fig. 3) was the largest distance a dispersing lynx was ever located from the centre of its natal home range. We compared MD with two maximum possible dispersal distances: from the centre of the natal range to the farthest edge of good lynx habitat that could be accessed within the same area (MPD_1 ; Fig. 3) and to the most distant good lynx habitat edge in the initial dispersal direction chosen by the respective subadult (MPD_2 ; Fig. 3). The areas of good lynx habitat were derived from lynx habitat models developed for the Alps (Zimmermann, 2004) and the Jura Mountains (Zimmermann & Breitenmoser, in press). We used the Mann-Whitney U-test (Zar, 1984) to compare dispersal distances between sexes and study areas, and the Wilcoxon paired-sample test (Zar, 1984) to compare MD with MPD_1 , MPD_2 and CD. Comparisons were made between (i) all subadults (Ph, fD and Di), (ii) only individuals that dispersed (fD, Di) and (iii) only those that completed dispersal (Di). Centroid distance of Ph, fD and Di is equivalent to the recovery distance,

with the centroid distance of Di describing the effective dispersal (Trehwella *et al.*, 1988).

To assess the effect of increased lynx abundance in, and dispersal from, the north-west Swiss Alps on lynx presence in neighbouring areas, we used information from a standardized monitoring system (see Capt *et al.*, 1998, for details). All data were calibrated according to the criteria defined for the pan-Alpine monitoring (Molinari-Jobin *et al.*, 2001), which defines three levels of reliability (Q1, Q2 and Q3). We considered only confirmed data (Q1 and Q2) to compare lynx presence and population trends in neighbouring areas but excluded depredation reports, as livestock is not consistently available in all areas.

Results

Only two females out of 14 subadults in the Jura Mountains and one male out of 13 subadults in the north-west Swiss Alps (F23, F36 and M30; Table 2) remained philopatric, and M30 died after a short dispersal (Fig. 3). Seven out of 9 subadults from the Jura Mountains that completed dispersal left the main study area whereas three out of 12 did so in the north-west Swiss Alps. None of the subadults left the Jura Mountains whereas one (M18) left the north-west Swiss Alps.

In both areas no difference between sexes was observed for centroid, total or maximum dispersal distances whichever dispersal category was considered (all $P > 0.05$). Males and females were therefore pooled for all subsequent analyses. Centroid distances were 4.5–56.0 km in the north-west Swiss Alps and 2.1–97.3 km in the Jura Mountains (Table 2). Subadult lynx that completed and/or had a fatal dispersal dispersed further in the Jura Mountains than in the north-west Swiss Alps: centroid distance was higher in the Jura Mountains than in the north-west Swiss Alps when either both fatal (fD) and completed dispersal (Di) or when only completed dispersal (Di) was considered but did not differ when all individuals (Ph+fD+Di) were included (Table 3).

Median recovery distance (CD of Ph, fD and Di) in the north-west Swiss Alps was 2.0 (range 0.4–5.6) times the mean circular resident female's home range diameter and 1.4 (range 0.3–3.8) times the mean circular resident male's home range diameter. In the Jura Mountains, it was 2.0 (range 0.1–6.6) and 1.5 (range 0.1–5.1) times the respective means. Median effective dispersal distance (CD of Di) was 2.1 (range 0.4–5.6) times female home range diameter and 1.5 (range 0.3–3.8) times the mean resident male's home range diameter in the north-west Swiss Alps. In the Jura Mountains it was 4.7 (range 0.1–6.6) and 3.6 (range 0.1–5.1) times the respective means.

Table 2 Dispersal type of 13 subadult lynx in the north-west Swiss Alps and 14 in the Jura Mountains with centroid, total, maximum dispersal distances, and maximum possible distances (MPD₁ and MPD₂), and % overlap with maternal home range. Centroid distance is also expressed as number of mean circular resident female and male home range diameters, respectively. See text and Fig. 3 for further details of the dispersal distances.

		Centroid distance			Total distance (km)	Maximum distance (km)	MPD ₁ (km)	MPD ₂ (km)	Overlap with maternal home range (%)
Lynx	Dispersal type ¹	km	No. ♀ HR diameter	No. ♂ HR diameter					
North-west Swiss Alps									
M18	Di	50.7	5	3.5			50	50	0
M20	Di	15.2	1.5	1			53	53	0
M24	Di	4.5	0.4	0.3	157	43.2	68	68	0.8
M25	Di	33.1	3.3	2.3	36	28.8	73	73	0
M29	Di	56	5.6	3.8	81	68.2	65	65	0
M30	fD	5.1	0.5	0.3	62	17.2	65	4	+ ²
M31	Di	36.7	3.6	2.5	164	51.2	63	63	0
M35	Di	20.1	2	1.4			76		0
F31	Di	22.1	2.2	1.5	55	22	58	58	0
F40	Di	33.3	3.3	2.3	69	35.9	68	25	0
F49	Di	7.4	0.7	0.5	2	17.4	65	65	0
FB22	Di	10.7	1.1	0.7			65		0
FB44	Di	20.6	2	1.4			65		0
Median		20.6	2	1.4	65.5	32.4	65	60.5	
Jura Mountains									
M11	Di	31.4	2.1	1.6			156		0
M13	Di	97.3	6.6	5.1			173		0
M14	Di	81.1	5.5	4.3	86	75.7	156	156	0
M15	fD	27	1.8	1.4	14	21	175	104	0
M16	fD	19.1	1.3	1	52	28	157	157	0
F12	Di	80.7	5.5	4.2	70	68.9	150	128	0
F13	fD	18.4	1.3	1	19	18.4	155	155	0
F17	fD	31.9	2.2	1.7	68	51	151	127	0
F20	Di	54.9	3.7	2.9	99	45.7	159	159	0
F22	Di	27.6	1.9	1.4	72	35.5	172	105	0
F23	Ph	11	0.7	0.6	51	34	143	143	12.2
F25	fD	24.9	1.7	1.3	91	46.8	170	112	0
F26	Ph	2.1	0.1	0.1	52	30	176	176	47.2
F36	Di	68.8	4.7	3.6	110	53.3	164	164	0
Median		29.5	2	1.5	69	40.6	158	149	

¹Ph, remained philopatric; Di, individuals that most likely completed their dispersal (6 months of site fidelity suggestive of home range establishment and/or surveyed until sexual maturity); fD, failed dispersal (<6 months site fidelity and/or died before establishing a home range)

²Overlap, but maternal home range not known exactly

Dispersal directions of individuals that either failed or completed dispersal (Table 2) were randomly distributed in the north-west Swiss Alps, but not in the Jura Mountains (Rayleigh test; north-west Swiss Alps: $Z = 2.1$, $n = 12$, $P > 0.1$; Jura Mountains: $Z = 5.5$, $n = 12$, $0.002 < P < 0.005$). The main dispersal direction in the Jura Mountains was to the south-west and north-east, corresponding approximately to the orientation of the predominant ridgelines of this mountain range (Fig. 1). The total and maximum dispersal distances were 2.0–164.0 km and 17.2–68.2 km, respectively, in the north-west Swiss Alps and 14.0–110.0 km and 21.0–75.7 km, respectively, in the Jura Mountains (Table 2). Total and maximum dispersal distances did not differ significantly

between the two areas whichever dispersal category was considered (Table 3).

In the north-west Swiss Alps CD was smaller than MD, indicating a circular dispersal, when all individuals, and only individuals that exhibited fatal or completed dispersal, were compared (Table 4), but did not differ for individuals that completed dispersal. No difference between CD and MD was observed in the Jura Mountains (Table 4).

In both areas the maximum distance was smaller than MPD₁ whichever dispersal category was considered but was significantly smaller than MPD₂ for all comparisons in the Jura Mountains but did not differ in the north-west Swiss Alps, indicating that in the latter most of the

Table 3 Statistical comparison between the centroid, total and maximum dispersal distances (medians with sample size in parentheses) of subadult lynx in the north-west Swiss Alps (NWSA) and the Jura Mountains (JM), using the Mann-Whitney *U* statistic. See text and Fig. 3 for further details of the three distances.

Dispersal type*	Centroid distance (km)				Total distance (km)				Maximum distance (km)			
	NWSA	JM	<i>U</i>	<i>P</i>	NWSA	JM	<i>U</i>	<i>P</i>	NWSA	JM	<i>U</i>	<i>P</i>
Ph+fD+Di	20.6 (13)	29.5 (14)	64	0.19	65.5 (8)	69 (12)	46	0.88	32.4 (8)	40.6 (12)	36	0.35
fD+Di	20.6 (13)	31.7 (12)	42	0.05	65.5 (8)	71 (10)	38	0.86	32.4 (8)	46.3 (10)	28	0.28
Di	21.4 (12)	68.8 (7)	11	0.01	69 (7)	86 (5)	12	0.37	35.9 (7)	53.3 (5)	7	0.09

*See footnote to Table 2

subadults reached the edge of suitable habitat during their dispersal (Table 4). The suitable habitat patch in the Jura Mountains is larger than in the north-west Swiss Alps as MPD₁ and MPD₂ were significantly higher in the former (all *P* < 0.05).

In the north-west Swiss Alps four out of nine subadults, all males, went beyond the edge of good lynx habitat while dispersing. Only M18, however, reached the neighbouring area of the west central Swiss Alps (Fig. 1). The three other males (M24, M29 and M30) returned after having spent a few days in the vicinity of a highway. Two of them (M24 and M30) turned back close to the place where they initially started their dispersal. M30 in the north-west Swiss Alps (Fig. 4) and M16 in the Jura Mountains, covered a considerable distance through sparsely wooded areas of the Swiss Plateau before returning to their respective mountain ranges.

The monitoring data showed a significant increase in lynx abundance in the north-west Swiss Alps from the

middle to the end of the 1990s followed by a decrease (Fig. 5b). No significant increase in lynx abundance could, however, be observed in the neighbouring areas of Valais and the west central Swiss Alps, even after a time lag of 3 years (Fig. 5a,c).

Discussion

Subadults from the north-west Swiss Alps and the Jura Mountains appeared to have the same dispersal potential as there were no observed differences between the two areas in the total and maximum distances dispersed. Contrary to our expectation, centroid dispersal distances did not differ between the two areas when all individuals were considered, and were only significantly smaller in the north-west Swiss Alps than in the Jura Mountains when individuals that most likely completed dispersal were considered. However, a larger proportion of individuals in the north-west Swiss Alps, all males, moved through unfavourable habitat but all stopped at fenced highways and turned back, except M18 which left the area. Two of the individuals settled in, or in the vicinity of, their maternal home range. Subadult lynx appear, therefore, to have a low capability to move through unfavourable habitat and to cross linear barriers such as fenced highways. Observations from adult radio-tagged lynx, which sometimes roam far outside their home ranges show, however, that lynx can cross such obstacles (Breitenmoser-Würsten *et al.*, 2001; Ryser *et al.*, 2004).

The apparent reduced ability of subadults to cross barriers led to circular dispersal in the case of two males in the north-west Swiss Alps. Similarly, severe habitat restriction led to philopatry in male cougar *Puma concolor coryi* in Florida, where they returned to the vicinity of their natal areas after unfruitful dispersal attempts (Maehr *et al.*, 2002). A similar process was described for the Iberian lynx *Lynx pardinus* (Ferrerías *et al.*, 2004). Habitat quality and barriers also shaped dispersal directions. In the north-west Swiss Alps, with no parallel ridgelines, dispersal directions were oriented randomly

Table 4 Statistical comparison between the centroid distance (CD) and maximum distance (MD) and between the maximum distance (MD) and maximum possible distances (MPD₁ and MPD₂) of dispersing subadult lynx in the north-west Swiss Alps and the Jura Mountains using a Wilcoxon paired-sample test (*Z*). See text and Fig. 3 for further details of CD, MD, MPD₁ and MPD₂.

Comparison*	NW Swiss Alps			Jura Mountains		
	<i>n</i>	<i>Z</i>	<i>P</i>	<i>n</i>	<i>Z</i>	<i>P</i>
CD/MD						
Ph+fD+Di	8	1.96	<0.05	12	1.07	0.29
fD+Di	8	1.96	<0.05	10	0.18	0.86
Di	7	1.69	0.09	5	1.48	0.13
MD/MPD₁						
Ph+fD+Di	8	2.38	0.02	12	3.06	<0.01
fD+Di	8	3.06	<0.01	10	2.80	<0.01
Di	7	2.20	0.03	5	2.02	0.04
MD/MPD₂						
Ph+fD+Di	8	1.54	0.12	12	3.06	<0.01
fD+Di	8	1.54	0.12	10	2.80	<0.01
Di	7	1.86	0.06	5	2.02	0.04

*See footnote to Table 2

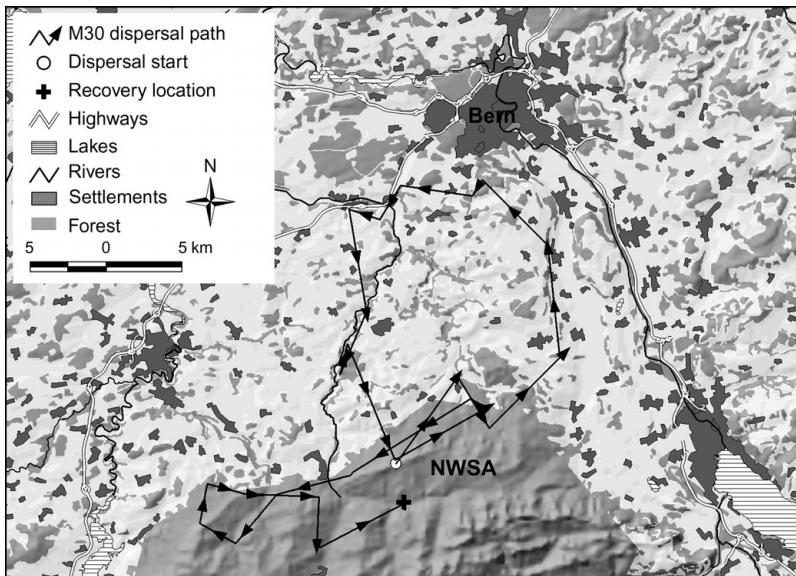


Fig. 4 Dispersal route of lynx M30. He traversed a sparsely wooded part of the Swiss Plateau and moved north until he reached the surroundings of Bern where he turned west. After moving into an area 8 km west of Bern he followed a fenced highway for >4.5 km and spent a week in the vicinity of both the highway and a railway. He returned to the north-west Swiss Alps (NWSA) by moving along a river course.

but the ridges of the Jura Mountains run south-west to north-east and this appeared to shape the movements of dispersing subadult lynx. The lack of south-east dispersal in the Jura Mountains may indicate hesitance to leave the continuous forest and travel over open agricultural areas. Similar behaviour has been reported in American black bears *Ursus americanus* in west Virginia that used the predominant ridgelines of the Appalachian Mountains as corridors (Lee & Vaughan, 2003), and in red fox *Vulpes vulpes* in North Dakota where dispersal directions were altered by a 4-lane interstate highway (Allen & Sargeant, 1993).

The most important outcome of our analysis, however, was that we failed to detect any positive density-dependent effects in lynx dispersal and hence cannot confirm the hypothesis that high population density will encourage the expansion of the population. Contrary to expectation a high proportion of subadults also dispersed in the Jura Mountains, an area with a lower lynx density and relatively high prey availability compared to the north-west Swiss Alps. Only one out of 12 individual lynx that completed dispersal actually left the north-west Swiss Alps, and the proportion of individuals leaving was smaller than in the Jura Mountains. As a consequence of the low rate of successful dispersals, the population pressure in the north-west Swiss Alps did not trigger an increase in lynx abundance in the two neighbouring areas (Fig. 5).

High lynx abundance can depress locally the numbers of roe deer and chamois *Rupicapra rupicapra*, lead to an increase in depredation on livestock, and ultimately diminish the acceptance of lynx by local people (Breitenmoser *et al.*, 1999). In 2000 at least eight individuals were known to have been killed illegally

(Breitenmoser-Würsten *et al.*, 2001) and four animals were removed as stock raiders during 1997–2001. In addition, six lynx were taken from the north-west Swiss Alps in 2001 for a translocation programme into the eastern Swiss Alps (Molinari-Jobin *et al.*, 2001). All of these removals lead to a considerable reduction in abundance (Fig. 5).

Generally, there is good evidence in natural populations that dispersal rate increases with increasing competition for limited resources (Lambin *et al.*, 2001) but dispersal rates have also been reported to be negatively density dependent, with a smaller fraction of individuals dispersing at higher densities (Wolff, 1997; Lambin *et al.*, 2001). Furthermore, those that disperse may only move relatively short distances (McCarty, 1997). This pattern has been reported for Townsend's voles *Microtus townsendii* (Lambin, 1994), red foxes (Trehwella *et al.*, 1988), and Canadian lynx (Breitenmoser *et al.*, 1993b). The little data available (Schmidt *et al.*, 1997; Sunde *et al.*, 2000) indicate that median recovery distance may be negatively correlated with lynx density and positively with mean male and female home range diameter. With the exception of the hunted Norwegian population, lynx dispersed roughly 1.5 male or 2.0 female home range diameters when recovery distances were considered. If dispersal is negatively correlated with density, the longest dispersal distances should be observed in areas where the lynx density is low and the home ranges large. The data (Table 5) seem to confirm this. However, central and northern Europe (where densities are generally much lower) are not necessarily comparable.

In contrast to wolves and bears, which can disperse over long distances (Swenson *et al.*, 1998; Merrill & Mech, 2000; Stratman *et al.*, 2001), subadult lynx are

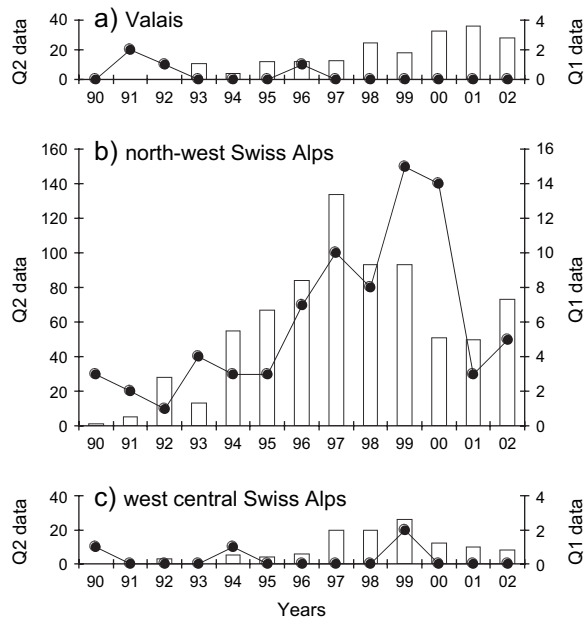


Fig. 5 Reported signs of lynx presence over 1990–2002 in (a) the Valais, (b) the north-west Swiss Alps and (c) the west central Swiss Alps. Reports of lynx killed or found dead, or young orphaned lynx caught and taken into captivity (Quality 1, continuous line, right y-axis), records of wild prey remains, tracks, scats, sightings, and vocalizations confirmed by trained personnel (Quality 2, columns, left y-axis).

conservative dispersers because of their life history and land tenure system and therefore maintain close contact to conspecifics (Zimmermann *et al.*, 2005). Their low ability to cross major barriers such as fenced highways hampers the colonization of a fragmented landscape such as the Alps. In the long-term the population may yet disperse but because of prey depression and conflicts

with local people high population densities are not maintained over long periods.

Natural spread of the species could potentially be increased by the use of wildlife crossings but the type of crossings that lynx will use have yet to be determined. Riparian vegetation may serve as natural corridors, as shown by the dispersal path of M30 (Fig. 4). Connectivity may be enhanced if wildlife crossings are established in the direction of predominant ridgelines or other landscape level features such as forests and riparian vegetation that may aid or direct dispersing lynx.

Our findings may have consequences for the design of carnivore reintroductions (Breitenmoser *et al.*, 2001) or recovery programmes in fragmented landscapes. Rather than creating one population, population nuclei could be founded in several neighbouring patches through reintroductions or artificial transfer of individuals to neighbouring areas. Considering that dispersal may be inversely density dependent, such an approach may be a better release strategy for reintroduction, as dispersing animals may encounter conspecifics when moving away from the release site. In a reintroduction project in Austria lynx dispersed in all directions from the site of release (Gossow & Honsig-Erlenburg, 1986) without founding a population.

To compensate for the lack of expansion of the Swiss populations six lynx from the north-west Swiss Alps and three from the Jura Mountains were translocated to the eastern part of Switzerland during the winters of 2000/2001 and 2002/2003. This allowed reduction of locally high abundance in the north-west Swiss Alps and accelerated the desired spread of the species. Paradoxically, it is possible that the intervention in the north-west Swiss Alps and the resulting reduction of the population will now lead to an increased dispersal rate.

Table 5 Data on dispersal distances of Eurasian lynx in four European areas, sorted from low to high density, with mean resident male and female home range diameters (HR ϕ) and population densities (number of resident lynx 100 km⁻²), and median recovery distances and effective dispersal distances (km; see text for further details), with median number of mean circular resident female and male home range diameters crossed.

Location	Mean HR ϕ (km)		Density	Recovery distance			Effective dispersal		
				Median, km (n)	Median no. of HR ϕ crossed		Median, km (n)	Median no. of HR ϕ crossed	
	♂	♀			♂	♀		♂	♀
Norway ¹	48.4	26.2	0.3 ⁵	40.6 (6)	0.8	1.6	41.2 (5)	0.9	1.6
Jura Mountains ²	19.1	14.7	0.8	29.5 (14)	1.5	2.0	68.8 (7)	3.6	4.7
Poland ³	16.9	13.0		25.0 (6)	1.5	1.9	50.5 (4)	3.0	3.9
Alps ⁴	14.6	10.1	1.5	20.6 (13)	1.4	2.0	21.4 (12)	1.5	2.1

¹Sunde *et al.* (2000)

²Breitenmoser-Würsten *et al.* (in press)

³Schmidt *et al.* (1997)

⁴Breitenmoser-Würsten *et al.* (2001)

⁵Managed population

Even though there are more vacant territories for youngsters close to their birth place in controlled populations there is no evidence that dispersal is less common (Macdonald & Johnson, 2001). If anything, control seems only to increase dispersal (Frank & Woodroffe, 2001). Exchange between established neighbouring populations will almost certainly take place later on, as resident adult lynx, especially males, show a higher propensity to cross barriers than subadults, particularly during the mating season. This could help to improve genetic exchange between established subpopulations.

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