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ORIGINAL PAPER

Life-history traits and potential invasiveness of introduced pumpkinseed *Lepomis gibbosus* populations in northwestern Europe

Julien Cucherousset · Gordon H. Copp · Michael G. Fox · Erik Sterud · Hein H. van Kleef · Hugo Verreycken · Eva Záhorská

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Abstract To evaluate the potential invasiveness of pumpkinseed *Lepomis gibbosus* introduced to northwestern European inland waters, growth and reproduction traits were examined in ten populations along a trajectory spanning northwestern Europe (Norway, England, Holland, Belgium and France) and evaluated

J. Cucherousset

Salmon & Fisheries Team, Centre for Environment, Fisheries & Aquaculture Science, Lowestoft, Pakefield Road, Lowestoft, Suffolk NR33 OHT, UK

J. Cucherousset

EcoLab Laboratoire d'Écologie Fonctionnelle, UMR 5245 (CNRS-UPS-INPT), Université Paul Sabatier, Bât. 4R3, 118, route de Narbonne, 31062 Toulouse, Cedex 9, France

Present Address: J. Cucherousset School of Conservation Sciences, Bournemouth University, Poole, Dorset, UK

G. H. Copp (🖂)

Salmon & Freshwater Team, Centre for Environment, Fisheries & Aquaculture Science, Lowestoft, Pakefield Road, Lowestoft, Suffolk NR33 OHT, UK e-mail: gordon.copp@cefas.co.uk

G. H. Copp School of Conservation Sciences, Bournemouth University, Poole, Dorset, UK

M. G. Fox

Environmental and Resource Studies Program and Department of Biology, Trent University, Peterborough, ON K9J 7B8, Canada in light of published dataset from Europe. In the 848 pumpkinseed captured, maximum age was 3-4 years, with a sex ratio near unity in all but one population. Significant variations with increasing latitude were observed in adult growth (age 2-3 increment in total length, TL) and mean age at maturity (A_M), with

E. Sterud

National Veterinary Institute, P.O. Box 8156 Dep., 0033 Oslo, Norway

Present Address:

E. Sterud National Institute of Technology, St. Hanshaugen, Akersveien 24 C, P.O. Box 2608, 0131 Oslo, Norway

H. H. van Kleef

Bargerveen Foundation, Department of Animal Ecology and Ecophysiology, Department of Environmental Science, Radboud University, Heyendaalseweg 135, 6525 AJ Nijmegen, The Netherlands

H. VerreyckenResearch Institute for Nature and Forest, Duboislaan 14,B-1560 Groenendaal–Hoeilaart, Belgium

E. Záhorská Faculty of Natural Sciences, Department of Ecology, Comenius University, Mlynská dolina B2, 842 15 Bratislava, Slovakia non-significant variations observed in juvenile growth (TL at age 2), sex ratio and gonado-somatic index. As observed elsewhere in Europe, mean $A_{\rm M}$ decreased significantly with increasing TL at age 2. Using this relationship, which has been proposed elsewhere as a potential predictive model of pumpkinseed invasiveness, eight of the ten populations could be provisionally categorized as 'non-invasive' (five populations), 'transitional' (one population) and 'potentially invasive' (two populations), with two populations not categorized due to insufficient data. Based on the available knowledge on each population, the relationship between juvenile growth and age at maturity appeared to predict reasonably the status of pumpkinseed in northwestern Europe and its applicability to other species should be tested.

Keywords Reproduction · Body condition · Life-history strategy · Latitudinal clines · Non-native species · Centrarchidae

Introduction

An invasive fish species has been defined as an "indigenous or nonindigenous species that spreads, with or without the aid of humans, in natural or seminatural habitats, producing a significant change in composition, structure, or ecosystem processes, or cause severe economic losses to human activities" (Copp et al. 2005). Invasive fishes are of increasing concern to the biodiversity and ecosystem integrity of fresh waters (e.g. Vitousek et al. 1997; García-Berthou et al. 2005), and tools to predict species invasiveness are needed (Kolar and Lodge 2001). Life-history traits have been shown to be particularly good predictors of non-native fish establishment success in many cases (e.g. Fausch et al. 2001; Marchetti et al. 2004; Vila-Gispert et al. 2005; Olden et al. 2006), and life-history perspectives are useful for a mechanistic-based understanding of the patterns of fish invasions and species invasiveness (Olden et al. 2006; Ribeiro et al. 2008). Intra-specific variation in life-history traits at large spatial scales (i.e. latitudinal clines) have also been demonstrated, due to the relationship between latitude and temperature. For instance, the decrease in growth and the increase in age at maturity with latitude has been observed in many fish species (Blanck and Lamouroux 2007 and references therein). Because these traits influence species potential invasiveness, latitudinal changes in the invasiveness between populations of a non-native species can be expected across its introduction range. Consequently, more integrative approaches of life-history traits of non-native species along latitudinal clines are needed to assess their potential invasiveness (García-Berthou 2007). Indeed, studies on life-history traits of non-native species usually consider only a limited latitudinal gradient, without the most northerly or southerly populations (e.g. Copp and Fox 2007; Fox et al. 2007; Ribeiro et al. 2008).

Recent applications in the biology of invasive fishes are now addressing the adaptations in lifehistory traits of non-native fishes in their new environments, in particular the introduction of North American fish species (Fausch et al. 2001; Copp et al. 2004; Vila-Gispert et al. 2005; Copp and Fox 2007; Ribeiro et al. 2008). One of the most successful fish introductions to Europe has been that of the North American centrarchid, pumpkinseed Lepomis gibbosus (L.). Introduced to European inland waters during the late nineteenth century, the pumpkinseed is now established in at least 28 countries of Europe and western Asia Minor (Copp and Fox 2007). Apart from recent studies of pumpkinseed populations in England (Copp et al. 2002; Villeneuve et al. 2005), all previous studies of pumpkinseed growth and lifehistory under natural thermal conditions have been undertaken in southern and central Europe (Copp et al. 2004; Fox et al. 2007). However, the increased attention to northwestern populations has revealed pumpkinseed to be more widely distributed within countries of northwestern Europe than was previously believed (e.g. Verreycken et al. 2007; Van Kleef et al. 2008), despite the fact that the pumpkinseed is considered a warmwater species. The study of northwestern European populations is especially relevant for understanding how pumpkinseed lifehistory varies according to local climate across a latitudinal cline, resulting in a different invasive potential. Moreover, using the relationship between age at maturity and juvenile growth, the potential invasiveness of pumpkinseed populations can be evaluated (Copp and Fox 2007).

The aim of the present study was to examine the variability of life-history traits of pumpkinseed populations across a latitudinal cline spanning northwestern Europe (southern Norway, northern and southern Netherlands, England, Belgium, northern France) in order to assess the potential invasiveness of the species in northwestern European and compare it with all available European populations (data compiled by Copp and Fox 2007). The specific objectives were to: (1) determine the general biological parameters and test for latitudinal clines in lifehistory traits of pumpkinseed within northwestern Europe, (2) test for the consistency of latitudinal clines in life history traits of pumpkinseed throughout Europe, and (3) examine the data in light of the model proposed by Copp and Fox (2007) to assess potential invasiveness of northwestern European pumpkinseed populations within a wider European context. Water temperature strongly influences the life-history traits of ectotherms in general (e.g. Atkinson 1994) and of pumpkinseed in particular (Fox and Crivelli 2001; Copp and Fox 2007), and is also negatively correlated with latitude. Consequently, we predicted that northwestern pumpkinseed populations would exhibit decreasing growth rates, increased age at maturity and reduced reproductive investment with increasing latitude and, thus, exhibit a lower invasiveness potential than southern Europe populations where potentially invasive populations have been reported (Fox et al. 2007).

Methods

Pumpkinseed were collected from 10 locations spanning northwestern Europe (Fig. 1; Table 1). Einedammen Pond (NEP), a 0.92 ha pond (mean depth = 1.5 m) located outside of Oslo (Norway), contains the most northerly pumpkinseed population of Europe (Sterud and Jørgensen 2006). In the Netherlands, pumpkinseed were sampled in the Schoapedobbe Pond (HSP; surface area = 0.5 ha; mean depth = 1.2 m) and in the Meeuwven Pond (HMP; surface area = 3 ha; mean depth = 1.0 m). The English sites were composed of two ponds and one stream section. Batts Bridge Stream (EBS) is a small tributary of the River Ouse (Sussex); it flows southward through a number of on-line ponds (Klaar et al. 2004; Villeneuve et al. 2005). Cottesmore School Pond (ECS) is a small man-made pond of



Fig. 1 Map of the locations in northwestern Europe where pumpkinseed (*Lepomis gibbosus*) populations were sampled in the summers of 2005 and 2006 (site codes in Table 1)

Country	Code	Site name	Sampling dates	Latitude	Longitude
Norway	NEP	Einedammen Pond	1 July 2006	59:49:00:N	10:27:00:E
Holland	HSP	Schoapedobbe Pond	21 June 2006	52:57:07:N	06:15:30:E
	HMP	Meeuwven Pond	22 June 2006	51:23:28:N	05:29:59:E
England	ECS	Cottesmore School Pond	8 June 2006	51:05:14:N	00:13:15:W
	EBS	Batts Bridge Stream	8 June 2006	51:00:47:N	00:04:27:W
	EAP	Isle of Wight	15 June 2005	50:39:11:N	01:11:03:W
Belgium	BHP	Webbekomsbroek Pond	23 June 2006	50:58:06:N	05:04:36:E
-	BHS	Slangebeek Stream	23 June 2006	50:57:22:N	05:18:10:E
France	FBM	Brière Marsh	14-17 May 2006	47:21:58:N	02:18:45:W
	FGL	Grand-Lieu Lake	19 May 2006	47:05:00:N	01:39:00:W

 Table 1
 Country, site code, site name, sampling dates, latitude and longitude of pumpkinseed Lepomis gibbosus populations sampling in northwestern Europe

about 0.7 ha area and constitutes the upper-most water body of a small stream system (Copp et al. 2002). Airport Pond (EAP) is a small pond of about 0.72 ha area with a mean depth of 1.5 m located in the Isle of Wight (English Channel). Although Cottesmore and Airport ponds were originally artificial water bodies, they have undergone ecological succession over numerous decades to achieve a pseudo-natural character. In Belgium, the two sites were located in the northeast of the country. Slangebeek Stream (BHS) is a small tributary of the River Demer and the study stretch was approximately 2 m in width with a depth of 0.15–0.30 m. Webbekomsbrock Pond (BHP) is an artificial water body of ≈ 2 ha surface area (depth varying between 0.5 and 1 m), with characteristics of near-natural floodplain ponds (Van Thuyne and Breine 2005). In France, the two sites belong to the River Loire drainage, in the northwest part of the country. Brière Marsh (FBM) is freshwater system (9,000 ha) composed of a complex network of permanently-flooded canals (Cucherousset et al. 2006). Grand-Lieu Lake (FGL) is a large, shallow, naturally turbid and eutrophic lake of 4,000 ha area (Paillisson and Marion 2006).

Pumpkinseed were collected in 2005 and 2006 during their spawning period (mainly from mid-May to early July) by various means (electrofishing: EBS, EAP, BHP, BHS, FBM, FGL; netting (because electrofishing was not possible): HSP, HMP; funnel trapping: ECS; angling (due to low water conductivity and dense ligneous debris): NEP; see Table 1 for site codes). Sample size at some sites was restricted due to low pumpkinseed density or reduced sampling efficiency (i.e. NEP and HSP). Captured fish were immediately killed by an overdose of anaesthetic and placed in a slurry of ice water and chilled to freezing.

After defrosting, each specimen was processed according to Fox (1994), which involved measurement for total length (TL, nearest mm) and wet weight (nearest 0.1 g), removal of the gonads by dissection for determination of weight (nearest 0.01 g), sex and determination of maturity status. Female pumpkinseed with ovaries containing nonvolked or indistinguishable eggs were classed as immature, and those with ovaries containing yolked eggs were classed as mature. Scale samples were collected from between the lateral line and dorsal fin of each specimen. Importantly, when sample sizes for a given size-class were large, only a subset of the specimens were aged. Age was determined as per Steinmetz and Müller (1991) using scale impressions on acetate strips, read on a micro-projector (magnification: $47 \times$), with age determination cross-checked using independent readings by a second reader on a sub-sample of specimens. Back-calculation of TL at age was undertaken using the linear relationship between scale radius and TL (Creaser 1926). Backcalculated TL at age estimated from this formula employed a fixed body-scale intercept value of 24 mm so as to reduce bias due to differences in the size distribution used to generate the intercept (Fox and Crivelli 2001). To provide an estimator of juvenile growth rate, we calculated the mean TL at age 2, the earliest age at which female pumpkinseed mature in cool-water populations (Fox 1994). As an estimator of adult growth rate, annual length increment was calculated as the difference in TL between ages 2 and 3, as TL at age 3 was the maximum age found in many of the sample sites.

Mean age at maturity (A_M) was calculated from the percentage of mature females in each age-class using the formula of DeMaster (1978) as adapted by Fox (1994):

$$\alpha = \sum_{x=0}^{w} (x) [f(x) - f(x-1)]$$

where α is the mean age at maturity, *x* is the age in years, *f*(*x*) is the proportion of fish mature at age *x*, and w is the maximum age in the sample. A modified version of this formula, using 10 mm TL intervals in place of age-classes (Trippel and Harvey 1987), was used to calculate mean total length at maturity (TL_M) according to Fox and Crivelli (2001). Gonadosomatic index (GSI: 100 × ovary weight \div [total body weight – ovary weight]) was calculated for mature females only, as GSI is not a good indicator of reproductive allocation in male pumpkinseed (Danylchuk and Fox 1994). The sex-ratio was expressed as the number of males divided by the number of females.

As described by Ricker (1975, 1979), the linear relationship for TL versus weight was determined using all sampled individuals. Mean body condition factor was calculated according to Copp (2003) using the formula of Le Cren (1951), which requires populations to be sampled during the same period of the year: $K_{\rm LC} = w/w'$, where w is the observed weight of each individual and w' is the expected weight using the length-weight relationship $(\log_{10} W = -a + b)$ $\log_{10} L$, where in this case a = -5.382 and b = 3.341; $r^2 = 0.9845, df = 847, P = 0.0001). K_{LC}$ values > 1 or < 1 indicate that the individual is in better or worse condition, respectively, than the average individual of the same TL range. The slope parameter 'b' from the regression equation $(\log_{10} W = b (\log_{10} TL) \pm a)$ was used as an estimator of 'generalised' condition (sensu Pitcher and Hart 1982).

Relationships between latitude and life-history traits (i.e., juvenile growth, adult growth, age at maturity, reproductive investment; proposed by Copp et al. 2002) and between age at maturity and juvenile growth were examined using linear regression, initially on the northwestern European data set and subsequently on the entire dataset available for Europe (as compiled by Copp and Fox 2007). The relationship between age at maturity and juvenile growth is used as a means of assessing the potential invasiveness of pumpkinseed populations in Europe, whereby the transition phase between non-invasive and potentially invasive pumpkinseed populations is defined at its lower extent by the minimum age at maturity (age 1) and at its upper extent by the end of juvenile growth (which for many pumpkinseed populations is age 2, Fox 1994; Fox and Crivelli 2001; Copp et al. 2004). In this relationship, noninvasive populations of pumpkinseed, such as observed in England (Villeneuve et al. 2005), are characterised by slow juvenile growth (small at age 2) and late maturity. Whereas, potentially invasive pumpkinseed populations, such as reported for southern Europe (Fox et al. 2007), are characterised by fast juvenile growth (large at age 2) and early maturity. In the present study, we identified the position of each pumpkinseed population in northwestern European on the age at maturity versus juvenile growth rate axis, and evaluated whether each population was likely to be 'non-invasive', in the 'transitional' phase or 'potentially invasive' based on its position on the axis and the available knowledge on each population.

Results

Of 848 pumpkinseed examined for condition and reproduction traits, 653 were aged. Total length (TL) ranged from 18 to 158 mm (n = 848), and maximum observed age was 4 years in six populations and 3 years in four populations (Table 2). Males outnumbered females in all populations, with sex ratio ranging from 1.1 to 3.2 (Table 2). TL at age 2 varied from 55 to 93 mm, with mean annual growth increments for ages 1 to 2 varying from 19 to 55 mm, and for ages 2 to 3 varying from 11 to 28 mm. Mean body condition factor ($K_{\rm LC}$) varied from 0.88 to 1.19, and generalized condition (*b*) varied from 2.47 to 3.64 (Table 2).

Age at maturity (A_M) and total length at maturity (TL_M) could be estimated in eight of the ten populations of northwestern Europe (only mature fish were captured at NEP and only one immature fish (age 1, 51 mm TL) was captured at HSP): A_M varied

Table 2 For pumpkinseed populations in northwestern Europe, the site (see Table 1 for codes), total number of specimens $(n_{\rm T})$, sex ratio (number of males \div number females); number of specimens subjected to age analysis $(n_{\rm A})$, the back-calculated TL at age (mm) and annual length increments for

ages 1–2 (L1–L2) and 2–3 (L2–L3), mean body condition factor ($K_{\rm LC}$, Le Cren 1951), generalised condition (b, slope of total length (TL) vs. weight relationship), mean age ($A_{\rm M}$, in years) and total length (TL_M, in mm) at maturity, and mean gonado-somatic index (GSI, in %)

Country	Site	n_{T}	Ratio	Mean TL at age				Increments		Condition		Reproduction			
				n _A	1	2	3	4	L1-L2	L2–L3	K _{LC}	b	$A_{\rm M}$	TL_M	GSI
Norway	NEP	38	3.2	38	47	85	97	_	38	12	0.91	2.47	_§	_§	7.22
Holland	HSP	54	1.3	53	38	93	113	124	55	20	1.00	3.05	_§	_§	10.04
	HMP	100	1.5	100	36	55	66	70	19	11	0.91	2.80	2.00	60.0	8.78
England	ECS	66	2.8	60	35	56	76	92	21	20	0.88	3.64	2.80	75.0	9.80
	EBS	100	1.1	66	36	61	83	-	25	22	0.97	3.40	2.50	78.3	4.13
	EAP	29	1.5	29	36	72	97	121	36	25	1.07	3.26	2.43	95.0	6.71
Belgium	BHP	150	1.2	86	46	89	109	120	43	20	1.04	3.50	1.63	80.3	6.10
	BHS	106	1.4	45	41	60	84	-	19	24	1.05	3.20	2.45	80.7	3.14
France	FBM	105	1.7	102	46	83	111	131	37	28	1.04	3.33	1.00	65.0	11.20
	FGL	100	2.0	74	40	73	95	-	33	22	1.19	3.31	1.71	63.0	11.36

[§] Value not available due to missing age and size classes critical to calculation

Table 3 Statistics of the linear regressions (y = bx + a) between latitude (lat) and juvenile growth rate (total length at age 2; TL_{L2}), adult growth increment (TL_{L2-3}), age at maturity

 $(A_{\rm M})$ and gonado-somatic index (GSI) and between juvenile growth rate and age at maturity of pumpkinseed in northwestern Europe and Europe

Populations	x	у	b	а	r^2	df	P-values
Northwestern Europe	Lat	TL _{L2}	0.928	25.530	0.051	8	0.529
	Lat	TL _{L2-3}	-1.044	73.897	0.461	8	0.031
	Lat	$A_{\mathbf{M}}$	0.237	-9.781	0.495	6	0.051
	Lat	GSI	-0.238	20.055	0.084	8	0.418
	TL _{L2}	$A_{\mathbf{M}}$	-0.035	4.466	0.561	6	0.033
All Europe [†]	Lat	TL _{L2}	-1.232	129.758	0.205	41	0.001
	Lat	TL _{L2-3}	0.026	17.447	< 0.001	41	0.905
	Lat	$A_{\mathbf{M}}$	0.091	-2.052	0.299	31	0.002
	Lat	GSI	0.008	7.164	< 0.001	34	0.927
	TL_{L2}	$A_{\mathbf{M}}$	-0.036	4.798	0.571	31	0.001

[†] The entire dataset available for Europe, i.e. those from Copp and Fox (2007) plus the present study

from 1.0 to 2.8 years and TL_M varied from 60 to 95 mm. Mean gonado-somatic index (GSI) was highly variable, ranging from 3.14 to 11.36% (Table 2). As expected, $A_{\rm M}$ decreased significantly with increasing juvenile growth rate (Table 3). Adult growth (TL increment between ages 2 and 3) decreased significantly with increasing latitude, and the increase in age at maturity with increasing latitude was very close to significant (Table 3). No significant variation with latitude was observed in the sex ratio ($r^2 = 0.25$, P = 0.14) nor in the juvenile

growth and GSI of the pumpkinseed populations of northwestern Europe (Table 3).

When the northwestern European data were combined with those from elsewhere in Europe, no significant latitudinal cline was observed in adult growth increment or GSI, but A_M increased significantly with latitude and juvenile growth decreased with increasing latitude (Table 3). Using the relationship between age at maturity and juvenile growth derived for European pumpkinseed populations (Fig. 2; Table 3) as a predictor of potential



Fig. 2 Mean age at maturity (A_M , in years) as a function of mean juvenile growth (TL at age 2) for European pumpkinseed populations—redrawn from Copp and Fox (2007) with superimposed new data (from Table 2) indicated with filled squares and circles as well as the site code given in Table 1. The proposed physiological transition phase between non-invasive and potentially invasive pumpkinseed populations is hypothesized as extending from the minimum age at maturity (the 45° *line* that traces from the intercept, at 'i') and the end of juvenile growth (which for many pumpkinseed populations is age 2; the 45° *line* that traces through the age 2 intercept with the regression slope, at 'ii')

invasiveness, five of the study sites would be categorized as 'non-invasive' (ECS, EBS, BHS, HMP, EAP), one as 'transitional' (FGL) and two as 'potentially invasive' (BHP, FBM). The two sites for which age at maturity are missing (NEP, HSP) remain unclassified.

Discussion

Life-history traits, and in particular age at maturity and juvenile growth rate, have been advocated as a means of predicting the potential invasiveness of pumpkinseed in Europe (Copp and Fox 2007). As predicted, pumpkinseed populations in northwestern Europe exhibited slower growth with increasing latitude, though this was restricted to the age 2–3 growth interval. This is consistent with the pattern of faster centrarchid growth in warmer waters reported elsewhere (McCauley and Kilgour 1990; Fox and Crivelli 2001). However, juvenile growth was not related to latitude when only the northwestern European populations were considered, whereas a significant relationship was established when the entire Europe level dataset was considered. This discrepancy might support the notion that environmental factors other than temperature become more important when considering a smaller latitudinal range.

The present study extends considerably the latitudinal range of current knowledge on the growth variability of introduced pumpkinseed, providing also information on sex ratio (not reported in previous studies). Sex ratio has been found to be a distinguishing, environmentally-linked character in another species invading northern Europe, the gibel carp *Carassius gibelio* (Vetemaa et al. 2005). However, in pumpkinseed, a species in which gynogenetic reproduction has not been reported, sex ratio did not vary with latitude in northwestern populations. As well, the present study provides information on pumpkinseed body condition, which has been reported elsewhere in Europe to vary greatly in pumpkinseed according to local conditions (Villeneuve et al. 2005), and this also appears to be the case in the northwestern parts of Europe (Table 2).

Assuming that latitude is an reasonable surrogate measure of general climatic conditions (water temperature and air temperature during the growing season), the results of the present and recent studies (Copp and Fox 2007) suggest that metabolism and growth performance changes with higher water temperature and longer growing seasons (e.g. higher food intake, higher food assimilation efficiency and/or lower energy expenditure; see also Dembski et al. 2006). Also as predicted, pumpkinseed populations in northwestern Europe exhibited a trend of later age at maturity than populations at lower latitude (Table 3), and this relationship was significant when all available data for Europe were included. The latitudinal clines in juvenile growth and age at maturity proposed previously (Copp et al. 2002; Villeneuve et al. 2005) were only partially corroborated in the present study, as juvenile growth was not related to latitude when only the northwestern European populations were considered, nor to gonado-somatic index with either population database. This suggests that age at maturity is a more useful (and reliable) means of assessing, even in smaller data sets such as this (Table 2), the reproductive response of pumpkinseed to environmental variability than other reproductive indices,

such as GSI or length at maturity (Copp and Fox 2007). As such, age at maturity should be a good parameter for estimating the potential invasiveness of pumpkinseed by virtue of its rapid response to growth potential. Whereas, the lack of relationship between GSI and latitude is in contrast to the initial assessment for pumpkinseed in Europe (Copp et al. 2002). Also, one-off assessments of GSI can misrepresent the true allocation to reproduction (Copp and Fox 2007), since the species is known to spawn over shorter periods, and with fewer batches, in colder water (Fox and Crivelli 2001). Therefore, further investigation of reproductive investment, undertaken throughout the spawning season, is needed to better understand the influence of environmental factors on this life-history trait.

Despite its relatively specialized reproductive strategy (polyphilous nest-guarder, sensu Balon 1975), the pumpkinseed exhibits great plasticity, even 'opportunism' in its range of life-histories (Fox et al. 2007), tolerating a wide range of environmental conditions (Vila-Gispert et al. 2002). The ability to switch to a more opportunistic life-history strategy may facilitate the expansion of an invading species, even though the primary life-history strategy of the species is equilibrium (Fox et al. 2007). The parental care that is exhibited by the pumpkinseed and which is typical of many equilibrium strategists has been associated with successful invaders (e.g. Olden et al. 2006; Statzner et al. 2008), even though at least some of these invaders appear to adapt to the novel environment by putting more effort into reproduction (Fox et al. 2007).

Strong juvenile growth and precocious maturity, and thus shorter life-span, appear to be adaptive responses to elevated water temperatures, as is predicted for most ectotherms (Atkinson 1994). The distribution, age structure and invasiveness potential of pumpkinseed populations in Europe are expected to change under conditions of climate warming (Klaar et al. 2004; Villeneuve et al. 2005; Dembski et al. 2006). Indeed, fish life-history traits have been used as correlates of introduction success and extirpation events (review in Olden et al. 2006). Growth and maturation characteristics of a population are influenced by ambient environmental conditions (e.g. climate, food and habitat availability, absence/presence of predators and of other fish species-see Belk and Hales 1993; Fox 1994; Fox and Crivelli 2001; Dembski et al. 2006). The model proposed by Copp and Fox (2007) is based on the relationship between juvenile growth and maturation, which is predicted for animals in general (Stearns 1992) and fish in particular (Fox 1994). It provides a useful means for assessing the potential invasiveness of pumpkinseed populations in Europe (Fig. 2). Indeed, almost all of the populations from southern Europe, where the species is considered invasive, exhibit fast juvenile growth and early maturity, i.e. the highest potential for population expansion by virtue of their short generation time (Copp and Fox 2007).

Within this context, two pumpkinseed populations located in the lower latitudes of the northwest Europe dataset (BHP, FBM) expressed $A_{\rm M}$ and juvenile growth characteristic of 'potentially invasive' populations (Fig. 2), and at both locations pumpkinseed were present in high densities. In contrast, the populations of two nearby sites (BHS, FGL) had A_M and juvenile growth values characteristic of noninvasive and transitional populations, respectively; and the densities of pumpkinseed at these sites were lower than the other nearby locations. In Belgium, this result may be due to the ecosystem characteristics (i.e. stream versus pond). Consistent with the original model (Copp and Fox 2007), the three studied English populations (ECS, EBS, EAP) fell into the non-invasive category, though the latter population borders the 'transition' zone (Fig. 2).

At present, neither of the most northerly populations (NEP, HSP) can be classified due to missing values and small sample sizes (see Table 2). Both of these populations were in very low abundance and shared their respective ponds with few or no other fish species at the time of sampling. The site near Oslo (NEP) is a recently introduced population (Sterud and Jørgensen 2006) and therefore may be at the start of an expansive establishment phase. And the site in northern Netherlands (HSP) had been subjected to an attempted eradication the year before sampling and therefore had been reduced by this human intervention to a status similar to that of the Norwegian population.

In conclusion, the use of the model proposed by Copp and Fox (2007), based on the relationship between juvenile growth and age at maturity, appeared to predict reasonably the status of pumpkinseed in Europe. However, other life-history characteristics (e.g. mortality rate, plasticity, reproductive strategy) are also likely to affect invasiveness (Olden et al. 2006; Statzner et al. 2008) and the model presented here undoubtedly does not capture all of the important biotic factors relating to invasive potential. As well, the alteration of the environment by human activities might increase species invasiveness (Vila-Gispert et al. 2005; Olden et al. 2006). Integrative tools are needed to predict species invasiveness (Kolar and Lodge 2001) and the applicability of this model to other species should be investigated. However, reliable growth and reproductive data for other species across their introduced range should be collected and the relationships between life-history traits and population invasiveness should be determined at the specific level.

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