



Climate changes and post-nuptial migration strategy by two reedbed passerines

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ABSTRACT: Phenology of biological systems is the expression of selective pressures forcing organisms to match their energy requirements to seasonal variations in resource availability. For long-distance migrant organisms, migration strategy is shaped by the availability and quality of stopovers during a precise time period. Ongoing climate changes alter the usual spatial and temporal distribution of resources, and, as a result, migrant species return earlier to their breeding grounds. Less is known on the evolution and determinants of timing of departure to wintering grounds, i.e. post-breeding migration. We analysed timing of post-nuptial migration and stopover strategy with capture–mark–recapture (CMR) data in 2 reedbed insectivorous passerines: the sedge warbler *Acrocephalus schoenobaenus*, specialised on a thermo-dependant prey, and the generalist reed warbler *A. scirpaceus*. Spring temperatures increased through years ($0.16 \pm 0.047^\circ\text{C yr}^{-1}$), and studied species migrated earlier in recent years ($-0.73 \pm 0.086 \text{ d yr}^{-1}$). Autumnal migration phenology was negatively related to spring (March) temperature ($-2.54 \pm 0.19 \text{ d }^\circ\text{C}^{-1}$), whereas it was not related to autumnal temperatures. We suggest that this results from a major time shift (i.e. advance) in biomass production caused by global warming. Stopover duration (estimated by CMR analysis) was positively related to body mass gain, with a trend for a stronger effect in the specialised species. However, neither stopover duration nor body mass gain were related to spring temperature, suggesting that the advance in migration timing efficiently adapted bird phenology to temporal variations in resources.

KEY WORDS: Climate change · Stopover duration · Optimal migration · *Acrocephalus scirpaceus* · *Acrocephalus schoenobaenus* · Capture–Mark–Recapture

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1. INTRODUCTION

Over the past 2 decades spring temperature has increased in the Northern Hemisphere temperate regions (Trenberth et al. 2007). These ongoing global climate changes have disturbed usual phenological patterns of distribution and abundance of resources. Phenology of biological systems is the expression of selective pressures forcing organisms to match their

energy requirements to seasonal variations in resource availability. There is already compelling evidence that plants and animals are affected by climate changes, particularly by modifying their phenological patterns (e.g. Myneni et al. 1997, Roy & Sparks 2000, Root et al. 2003, Cleland et al. 2006, Grosbois et al. 2006, Rivalan et al. 2007). For example, many bird species lay eggs earlier in spring to adapt to changes in prey phenologies (Crick et al. 1997, Crick & Sparks 1999, Both &

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Visser 2001, Both et al. 2005). As migration dates are most likely plastically adjusted to experienced environmental conditions, rapid adjustments of migration phenology to changing environmental conditions are expected. Rapid changes of migration timing were quantified from spring phenology of long-distance migrants (Lehikoinen et al. 2004, Jonzén et al. 2006, Rubolini et al. 2007, this issue, and references therein). Furthermore, since migration date also has a genetic basis (Møller 2001), climate change could induce a heritable, evolutionary response of migration timing.

There is less information available on the impact of climate change on the timing and ecology of autumnal, post-nuptial migration (Gatter 1992, Sokolov et al. 1999, Sparks & Braslavská 2001, Cotton 2003, Jenni & Kéry 2003, Lehikoinen et al. 2004, Gordo & Sanz 2006, Sokolov 2006). Every possible pattern of change of post-breeding migration timing (no change, later departure date, earlier departure date) has been reported (e.g. Lehikoinen et al. 2004). Inter-annual variation in post-breeding behaviour seems to depend on ecology and life-history traits of bird species. For instance, short-distance migrants and species with variable reproductive effort left later in recent warmer years (Jenni & Kéry 2003), suggesting that these species possibly benefit from climate warming through increased reproductive effort. Some species adjust the choice of their wintering ground to new temperature conditions (Austin & Rehfish 2005, Rivalan et al. 2007), or increase the time spent on their breeding grounds (Gordo & Sanz 2006). On the opposite hand, among long-distance migrants, the date of departure from breeding grounds decreased throughout the 20th century, but this relationship is controversial: although several studies report earlier departure of some long-distance migrants in recent warmer years, most found that migrants did not change or even delayed their departure date (review in Lehikoinen et al. 2004, Sokolov 2006). Response of migrant birds to warmer, longer autumn is hence contrasted. Few studies analysed the effect of temperature, and, among those that did, most attempted to correlate autumnal migration timing to temperatures in autumn or during the breeding season (e.g. Gordo & Sanz 2006). However, spring temperatures are likely a better determinant of autumnal migration timing than summer or autumnal temperatures (Sokolov et al. 1999, Sokolov 2006) under 2 main hypotheses that we will develop in the present study: (1) fitness benefits from early arrival at wintering grounds and (2) a temporal constraint results from the phenology of food availability at stopover sites.

The first objective of this study was to test for an advance in post-breeding migration date and, if found, to determine which temperatures from spring or autumn were the best predictors. The second objective

was to test the 2 hypotheses that explain why birds do not take advantage of climate change to spend more time on the breeding grounds and eventually lay more clutches. The third objective was to address which patterns of phenotypic plasticity and/or natural selection are most likely to explain the observed temporal changes in migration phenology, under the assumption that if plasticity is mainly involved, the observed changes in stopover ecology should be better explained by a physical explanatory variable, such as temperature, than by a linear year effect (Jonzén et al. 2006).

These predictions were explored by correlative analysis of the effects of monthly mean temperatures on post-breeding migration timing in 2 trans-Saharan migrant birds: the sedge warbler *Acrocephalus schoenobaenus* (SW) and the reed warbler *A. scirpaceus* (RW). SW diet during post-breeding migration is mainly composed of reed aphids *Hyalopterus pruni* (Bibby & Green 1981), whereas RW is a generalist insectivore (Bibby & Green 1981, Idrissi et al. 2004). These inter-specific diet differences are used hereafter to discriminate the 2 hypotheses about which mechanisms select for advanced migration in response to warmer years. We acknowledge that no causal relationship between diet and behaviour can be formally tested in the present study. Experimental manipulations of food availability and/or comparative analysis involving more species would be required to validate the interpretations of correlative results proposed hereafter.

In the first hypothesis, early arrival at wintering grounds increases fitness, possibly through acquisition of a better wintering territory (Marra et al. 1998, Salewski et al. 2002, Studds & Marra 2007, this issue). As long-distance migrants arrive earlier at their breeding grounds, and breed earlier, in warmer springs (likely for RW, Bergmann 1999, Crick & Sparks 1999, Schaefer et al. 2005, and, for SW, Crick & Sparks 1999; demonstrated for numerous other Palearctic insectivorous passerine migrants: Jonzén et al. 2006), migrants should also depart earlier in autumn, once breeding is achieved, in order to reach their wintering grounds as soon as possible. In this hypothesis, timing of autumnal migration should be at least partly, if not mainly, determined by spring temperatures, and both studied species should migrate earlier in autumns following warm springs. This is analogue to a complete time shift in the annual life cycle of long-distance migrants (Cotton 2003, and see Fig. 2 in Gordo 2007, this issue).

In the second hypothesis, timing of autumn migration is determined by availability of food resources en route. In this hypothesis, the birds are constrained to track the changes in the phenology of their insect prey. Following Bibby & Green (1981), we suppose that

warmer springs induce earlier appearance of the aphid abundance peak in autumn, because all primary production (Cleland et al. 2006) and hence secondary production is advanced by warmer springs. The constraint is then stronger for SW (the aphid specialist) because aphid populations are characterised by a short period of availability. Hence, under this hypothesis, SW should depart breeding grounds earlier in warmer years to track resource availability at stopover sites, and onset of its migration should be correlated to spring temperatures. Because of its generalist diet, RW should be less affected by among-year variations in stopover condition, and its migration timing should not correlate to spring temperatures. In terms of predictions, the second hypothesis predicts that spring temperature will correlate to migration timing in SW only, or at least more strongly in SW, whereas the first hypothesis predicts no interaction between species and temperature. In both hypotheses, we furthermore compared the effects of temperature and year to determine the extent to which plasticity is involved (Jonzén et al. 2006), and evaluated the extent to which a warm autumn can release the constraint by comparing models in which spring and autumn temperatures interact.

To further investigate response of these 2 migrants to a changing stopover environment, we estimated 2 other variables of the stopover ecology: stopover duration (SOD) and daily mass gain (DMG). These 2 variables are linked in the frame of optimal migration theory (OMT; Alerstam & Lindström 1990). Intuitively, the higher the food availability, the quicker birds fatten, and the quicker they leave the stopover site. However, OMT predicts the opposite pattern: individuals should continue to store fat as long as the DMG at the site is higher than the average expected DMG at forthcoming sites, further south in the case of post-breeding migration of RW and SW (Bibby & Green 1981, Alerstam & Lindström 1990, Hedenström et al. 2007, this issue). This strategy should be stronger in SW: if the populations of aphids peak during a specific time frame that is linked with spring temperature, SW that encounter high food availability should not move further south, where the resource is likely to already be scarce; whereas, for RW, the constraint is weaker because they rely on a broader spectrum of food resources likely to be more uniformly distributed along their migration route (Bibby & Green 1981, Schaub & Jenni 2001, Idrissi et al. 2004). The prediction is then that variation in site quality (as experienced by the birds at the time they stop at the site), quantified by a proxy (daily DMG), will correlate more strongly to stopover duration for SW than for RW. This prediction is likely to be impacted by climate change if migrants do not efficiently track the changes in their en route food resource phenology: if migrants arrive at stopover

sites after the abundance peak of their prey, we predict a decrease in DMG and in SOD, particularly for SW (the aphid specialist); however, if the advance in migration timing is sufficient, there should be no such decrease because the experienced food availability does not change.

2. METHODS

2.1. Study species

Acrocephalus schoenobaenus (SW) and *A. scirpaceus* (RW) are small (10 to 14 g in the breeding season), insectivorous passerines breeding in wetlands of the Palearctic. The populations concerned in the present study breed in north-western Europe. They migrate through coastal western France from late July to late September and stopover essentially in reedbeds. Their winter quarters are located in sub-Saharan Africa, where they occupy grasslands and wetlands (Cramp 1992).

2.2. Data collection

Reedbed passerines post-nuptial migration was monitored by capture–mark–recapture (CMR), using mist-nets, at 3 sites that represent the most important reedbeds in coastal western France: Marais du Hode (Site S76: Seine river estuary, 49° 27' N, 0° 26' E; 19 912 SW and 31 864 RW between 1983 and 2005; stopover analysis was performed from 1994 onward because recaptures were too few in earlier years), Etang de Trunvel (Site S29: Audierne's Bay, 47° 55' N, 04° 19' W; 33 271 SW and 13 416 RW between 1994 and 2005; Bargain et al. 2002) and Île du Massereau (Site S44: Loire river estuary, 47° 14' N, 01° 55' W; 37 208 SW and 21 409 RW between 1994 and 2006; Caillat et al. 2005).

At S76, capture sessions lasted 16 d on average (SD = 7.0) in the period from 1 August to 5 September. Capture dates (i.e. dates at which nets were opened) advanced with years (-0.58 ± 0.077 [SE] d yr⁻¹, $r^2 = 0.74$, $p < 0.01$), with the mean capture date on 22 August (SD = 4.0). At S44, capture sessions lasted 46 d on average (SD = 15) between 15 July and 15 September. Capture sessions were earlier in recent years (-1.13 ± 0.30 [SE] d yr⁻¹, $r^2 = 0.55$, $p < 0.01$), with the mean capture date on 18 August (SD = 5.9). At S29, capture sessions lasted 50 d on average (SD = 9.6) between 15 July and 15 September. Capture session dates were similar through years ($p > 0.5$), with the mean capture date on 16 August (SD = 2.1). The catching effort per day, i.e. number of mist-nets and duration of CMR sessions in days, varied within and among years. The dependent variable used to quantify the

migration flux per species was therefore a capture index, noted N_p and computed as the number of individuals captured per meter of net per day.

Temperature data were obtained from the Meteo France stations of Cap de la Hève (49° 30' 36" N, 0° 04' 12" E) for S76, Pointe de Pen Marc'h (47° 47' 48" N, 4° 22' 24" W) for S29, and Saint-Nazaire Montoir (47° 19' 00" N, 2° 10' 00" W) for S44. Monthly means (computed from daily means) are hereafter referred to as T_i standing for temperature in the i th month. Temporal trends on monthly mean temperatures were tested with linear regressions.

2.3. Data analysis

2.3.1. Migration phenology, peak dates and correlations with temperature

Timing of migration was characterised each year for Site S76 by the migration peak dates, i.e. the estimated dates at which N_p was maximum. The trapped birds were most likely heterogeneous in origin, age and quality; hence multimodality in the abundance pattern could be expected. However, visual examination of the data reveals that 1 mode of N_p dominated. A modelling method assuming unimodality thus seemed reasonable to extract the main information on timing of migration. The variation in session durations and dates made necessary the use of a method able to estimate peak date even if the peak occurred outside of one given capture session. For this purpose, 3 alternative non-linear models were fitted to daily N_p data, with the least-squares method. We have used the following parametric functions (where t is the time in days and a , b and c are the parameters to be estimated):

$$f(x) = ax^2e^{-x^2}, \text{ where } x = (t - b)/c \quad (1)$$

$$f(x) = (ae^{-x})/(1 + e^{-x})^2, \text{ where } x = (t - b)/c \quad (2)$$

$$f(t) = at^2 + bt + c \quad (3)$$

(t varying between 1 [1 August] and 36 [5 September]). These functions were considered as possible forms of the abundance curve of the species: Function (1) is asymmetrical with post-peak abundance decreasing more slowly than it increased before peak abundance; Function (2) models a narrow window of high abundance, with few early and late-migrating individuals; and Function (3) models a parabolic curve of abundance. Separately for each year and for each species, we used, among the 3 models, the one with the lowest deviance to compute the date of maximum N_p (and associated SE). If the computation process failed for all 3 models for a given species-year dataset, indicating inappropriate fit of all the models to the data, then no estimate of peak date was available for subsequent analyses.

Theoretically, the above-described method correctly estimated peak dates, even outside of the capture session. To check for the robustness of peak date estimates, we generated 3 simulated datasets, with the migration peak dateset on 5, 17 and 28 August, and a random, normally distributed error with standard deviation taken as the maximum $N_p/10$. These simulated datasets were analysed through the same 3 models and with the same dates of capture as the real dataset. Each of the 3 simulations was repeated 10 times.

Estimated migration peak dates per species and per year were then used as dependent variables in a linear regression analysis to identify the best predictors of observed among-year variations in migration timing. The accurateness of peak date estimates was accounted for by including $1/SE^2$ as a weight of the dependent variable in the regression. Explanatory variables were: species (Sp , 2 modalities), year (Y , taken as a continuous variable), monthly mean temperatures at the given site (T_i for temperature in the i th month, i varying between 02 for February to 09 for September). Preliminary to formal analysis, we tested for a linear temporal change in migration peak date throughout the study period for each species, which could be taken as indicative of a possible effect of climate change. Afterward, the analysis relied on an information theoretic approach (Burnham & Anderson 2002), where the influence of effects on migration peak date was evaluated by multimodel inference. To restrict the number of candidate models to its minimum, models that were *a priori* biologically meaningless and models with interactions involving 3 terms or more were not considered. Step 1 of the selection procedure was the model with a linear effect of year only. We then compared this model with the model including temperature as an explanatory variable. Step 2 was then the exploratory analysis of the effect of temperature in different months of the year on migration peak date. Because of the lack of strong prior knowledge on which temperature should be most influential, we had to consider a relatively high number of models: 8 models of the form $Sp \times T_i$. T_j hereafter refers to the best temperature as selected by this procedure. We compared the model $Sp \times T_j$ and the model with a temporal trend to evaluate if temperature was a better predictor of temporal changes than a simple directional change throughout the study period. Our model selection procedure then followed 4 more steps (1 model per step). Step 3: the interaction $Sp \times T_j$ was removed to determine if the effect of temperature differed among species. Step 4: interaction terms among spring and autumn temperature effects were added to assess the possible interplay of spring and autumnal temperature. If a spring temperature was selected as the best

predictor, the interaction with the temperature at the time of capture (August) was added to the model; if August temperature was selected, we added the interaction with all spring temperatures. Step 5: a linear effect of year was added to the selected temperature effect to evaluate if plastic adaptation alone (no effect of year) to temperature is the main mechanism explaining the observed change in migration peak date, or if directional natural selection was also likely to have a role (effect of year; cf. prediction from Jonzén et al. 2007 Section 1, this issue). Finally, in Step 6, we compared our final model to the model with the mean capture date as a unique explanatory variable. This allowed us to check whether biological determinants of the temporal change in migration peak date were not confounded by the possible methodological bias due to among-year changes in capture session dates. At each step, the best model was identified as the one with the lowest AIC (Akaike's information criterion), with a 2 point AIC difference when compared to other models (Burnham & Anderson 1998). When models differed by <2 points of AIC, their respective probability was assessed by Akaike's weight w_i ; w_i gives the probability that a model is the real model, given the data and given that the real model is among those included in the model selection (Burnham & Anderson 1998, p. 124). At the end of this procedure, estimates and standard deviations of the slopes were presented. The influence of effects of interest is evaluated with Akaike's weights. Goodness of fit (GOF) was assessed by testing the normality of residuals of the best model with a Shapiro-Wilk test and through the dispersion coefficient (residual deviance/residual degree of freedom). All analyses were performed with R (R Core Development Team 2005).

2.3.2. Stopover duration, body mass gain and correlates

Average stopover durations per species per year per site were estimated with the software SODA 2.1.2 (Schaub et al. 2001). This CMR modelling of stopover data is advantageous because it gives access to the 'true' stopover duration, i.e. estimating the total time spent at the stopover site by summing estimates of the time spent at the site *before* (since arrival) and *after* (up to departure) the first capture, while it adjusts SOD estimates for potentially confounding variations in capture probability (hereafter noted P). Thus, it provides robust, daily, local survival (hereafter noted Φ ; Lebreton et al. 1992) and SOD estimates. Note that the SODA method assumes that the stopover duration is constant among birds (within seasons). Efford (2005) suggested that stopover duration may indeed vary

among individuals, following a Poisson distribution. In this case, SODA estimates are shown to be twice the real SOD, and the average time of survival after first capture is a better estimate of the average SOD than SODA estimates. In the present study, we chose to rely on SODA and not on 'local survival' analysis for 3 reasons. (1) According to what is known about behavioural and meteorological determinants of migration waves in small passerines (in Pradel et al. 2005), SOD is likely to be closer to constancy among birds rather than to following a Poisson distribution (Efford 2005); thus SODA seems likely to perform better than Efford's method in a wider range of situations. (2) No statistical method allows us to identify the true distribution of SOD (Pradel et al. 2005); thus, in our case, nothing supports that the 'life expectancy' parameter (Efford 2005) would be a better approximation of SOD than SODA estimates. (3) Our parameter of interest is the SOD, i.e. the resultant of both immigration *and* emigration probabilities (as modelled by SODA), and not only the emigration part of the SOD strategy (Efford 2005). Note that no CMR method can account for the likely dependency of the emigration probability on the time spent at a stopover site. This is a major caveat of SOD approaches with CMR data, indicating that all studies, including ours, are to be considered exploratory (Pradel et al. 2005).

GOF was tested with the software U-Care 2.2.3 (Choquet et al. 2005) for the model [$\Phi(\text{species} \times \text{site} \times \text{year} \times t)$, $P(\text{species} \times \text{site} \times \text{year} \times t)$] to detect transients (Test 3.SR) and trap dependence (Test 3.CT). Overdispersion was tested using the overall test computed from the 4 components. Afterward, for the sake of simplicity, daily local survival was assumed to be constant within years, but varied across years, site and species, and recapture probability was assumed to vary among days and among years {model [$\Phi(\text{species} \times \text{site} \times \text{year})$, $P(\text{species} \times \text{site} \times \text{year} \times t)$]}.

DMG was calculated for individuals that were captured more than once during a single migration period. DMG is the difference in body mass divided by the number of days between 2 consecutive capture events (in g d^{-1} ; averaged per individual if >1 recapture event was available).

Estimates of SOD per species, per site and per year were then used as dependent variables in a linear regression analysis based on model selection (same procedure and criteria as for the migration peak date analysis). The accurateness of SOD estimates was accounted for by including $1/\text{SD}^2$ as a weight of the dependent variable in the regression. In a first step, we computed separate univariate linear regressions for each species to obtain comparable estimates of the slope of the effect of DMG on SOD (potentially useful for future analysis). We also tested the relationship

between DMG and T_i with separate linear models. We then compared a set of models including the effect of DMG in addition to the effects of temperature, site and species. Step 1: we departed from the model $S + Sp \times DMG$, i.e. the model with no temperature or year effect. The exploratory analysis of the effect of temperature was conducted in Step 2, where we compared 8 models of the form $S + Sp \times T_i + Sp \times DMG$ (S for site, other effects noted as in the previous section), where only the effect monthly temperature was allowed to change. T_k refers to the temperature selected during this first phase. The model $S + Sp \times T_k + Sp \times DMG$ was compared to the model of Step 1 to evaluate the importance of temperature as a determinant of SOD. Then, inference was based on 7 more steps of model selection (1 model per step). Step 3: a model without the DMG effect was compared to the model selected in Step 2 to confirm that DMG was an important predictor of the SOD. Step 4: a model with a linear effect of year alone was compared to the model selected in Step 2 to confirm that the predicted plastic adaptation of SOD to DMG and T_k was a better determinant of the estimated SOD than a directional change throughout the study period. Step 5: the interaction $Sp \times T_k$ was removed from model $S + Sp \times T_k + Sp \times DMG$ to determine if species responded differently to temperature. Step 6: the interaction $Sp \times DMG$ was removed to evaluate if species adjusted their SOD to DMG differently. Step 7: a linear effect of year was added to a model $S + Sp \times T_k + Sp \times DMG$, where the interactions not selected in Steps 3 and 4 had been removed. If plastic adaptation to the experienced conditions, as approximated by DMG and T_k , is the only component of the observed variations, the temporal trend should not be selected. Step 8: the interaction $S \times DMG$ was added to check whether the effect of DMG differed among sites. Finally, Step 9: the interaction $T_k \times DMG$ was added to determine if DMG mediated the effect of temperature on SOD. At the end of this procedure, estimates and SD were presented. GOF was assessed by testing the normality of residuals of the best model with a Shapiro-Wilks test and through the dispersion coefficient (residual deviance/residual degree of freedom).

3. RESULTS

3.1. Evolution of mean monthly temperatures

Mean temperatures in February, March, April and September were positively correlated to year (all $r^2 > 0.3$ and $p < 0.01$). Slopes were, respectively, 0.25 ± 0.072 , 0.16 ± 0.047 , 0.09 ± 0.031 and $0.11 \pm 0.033^\circ\text{C yr}^{-1}$. Mean temperature in August was not related to year ($p > 0.5$).

3.2. Phenology, peak date and temperature

The peak dates computed from the simulated datasets with fixed migration timing exhibited no advance with year (all 3 $p > 0.5$) and were correctly estimated (average estimated peak date for the 3 simulated datasets: 5 ± 1.0 , 18 ± 1.5 , 28 ± 1.2 , for simulated peak date of 5, 17 and 28 August, respectively). Thus, the method to compute migration peak dates produced estimates robust to among-year variations in dates of capture sessions in our simulated cases.

In the real dataset, peak dates were obtained for all but 3 species-years (Fig. 1). The second functional form was selected as the best descriptor of the N_p phenological pattern in 29 species-year datasets out of 46.

Peak date was negatively correlated to the year for SW (linear model: $r^2 = 0.51$, slope = $-0.72 \pm 0.16 \text{ d yr}^{-1}$; Fig. 1) and RW (linear model: $r^2 = 0.87$, slope = $-0.78 \pm 0.067 \text{ d yr}^{-1}$; Fig. 1). Pooled together, both species shifted peak date by $-0.73 \pm 0.086 \text{ d yr}^{-1}$. As average temperatures have increased, there was a potential effect of climate change on migration timing to investigate. Temperatures were far better in determining the timing of migration than a linear temporal trend (Table 1, Step 2.2 vs. Step 1). The selection procedure gave temperature in March as the most important factor (Table 1, Steps 2.1 to 2.8: AIC differences between Step 2.2 and the others are >30 points). The interspecific difference in migration timing could be neglected (Table 1, Step 3 vs. 2.2: weight ratio indicates that the model with no interspecific difference is 6.3 more likely than the model with species differences). Both species migrated earlier in years of warmer March temperatures (estimate of the slope in Step 3: $-2.54 \pm 0.19 \text{ d }^\circ\text{C}^{-1}$). The estimated migration timing, which

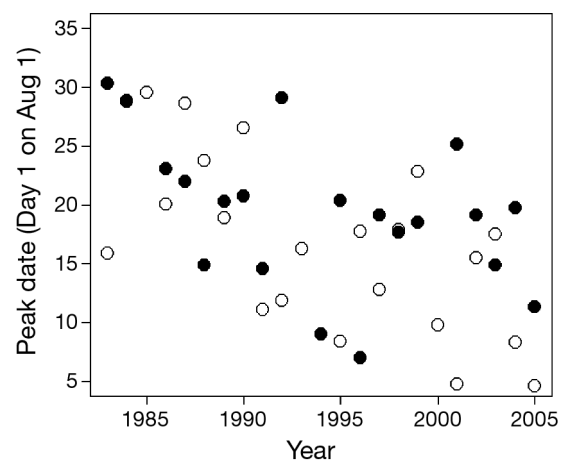


Fig. 1. *Acrocephalus schoenobaenus* and *A. scirpaceus*. Peak date of the post-breeding migration of sedge warblers (SW, ●) and reed warblers (RW, ○)

Table 1. Model selection for peak date, second phase. *Sp*: effect of species; *Y*: linear effect of year; T_i : linear effect of mean temperature in the *i*th month; *C*: linear effect of mean date of capture sessions; *w*: Akaike's weight. Best models are indicated in bold. Shapiro-Wilks normality test on residuals of model $\sim T_{03}$: $w = 0.9774$, $p = 0.5465$; dispersion coefficient: 21.2. AIC: Akaike's information criterion

Step	Model	AIC	w
Linear temporal change			
1	$\sim Y$	334.1	1.87×10^{-7}
Influence of monthly temperatures per species			
2.1	$\sim Sp \times T_{02}$	361.18	2.47×10^{-13}
2.2	$\sim Sp \times T_{03}$	308.85	0.06
2.3	$\sim Sp \times T_{04}$	359.65	5.31×10^{-13}
2.4	$\sim Sp \times T_{05}$	376.33	1.27×10^{-16}
2.5	$\sim Sp \times T_{06}$	368.66	5.87×10^{-15}
2.6	$\sim Sp \times T_{07}$	372.95	6.87×10^{-16}
2.7	$\sim Sp \times T_{08}$	368.82	5.42×10^{-15}
2.8	$\sim Sp \times T_{09}$	339.82	1.07×10^{-8}
Most influential monthly temperature without species differences			
3	$\sim T_{03}$	305.04	0.38
Interaction between most influential monthly temperature and August temperature			
4	$\sim T_{03} \times T_{08}$	305.36	0.33
Temporal linear trend in addition to March temperature			
5	$\sim T_{03} + Y$	306.03	0.23
Effect of capture dates			
6	$\sim C$	326.79	7.25×10^{-6}

mainly takes place in August, may thus be influenced by the interaction of March and August temperatures. However, comparing Step 3 vs. Step 4 provides weak support for such an interaction effect of August temperatures: the model with interaction is about as likely as the model without interaction. The model including the interactions suggests that warm August temperature tended to moderate the effect of warm March temperatures (Table 2: interaction term in the model $\sim T_{03} \times T_{08}$), with no apparent additive effect of August temperatures. The additive temporal trend did not markedly improve the fit (Step 5 vs. Step 3): the model without temporal trend was slightly (1.6 times) more likely than the model with the trend. It suggests that most among-year variation was explained by temperature.

Table 2. Parameter estimates from the model $T_{03} \times T_{08}$ for peak date. T_i : linear effect of mean temperature in the *i*th month

Parameter	Estimate	SE
Intercept	61.3327	27.4964
T_{03}	-6.7763	3.2502
T_{08}	-1.1313	1.5488
$T_{03} \times T_{08}$	0.2219	0.1791

The effect of capture dates was not selected (weight of Step 6 < 0.01 in Table 1), further emphasising that the true temporal variations in migration peak date were not strongly biased by changing capture dates among years. Finally, a linear temporal trend alone was a worse predictor of among-year variations in migration peak date when compared to the predictive power of temperature models, with a probability < 0.01 (weight of Step 1 in Table 1). All models including a year effect (Steps 1 and 5) had a summed weight of 0.23.

3.3. Stopover duration, body mass gain and temperature

Some lack of fit of the general model used for SOD estimation [$\Phi(\text{species} \times \text{site} \times \text{year} \times t)$, $P(\text{species} \times \text{site} \times \text{year} \times t)$] due to the excess of individuals captured only once (transience, Test 3.SR) was detected in 18 species by site by year subsets out of 62 ($p < 0.05$ in Table 3). If we corrected these GOF tests for multiple testing, the detected lack of fit would be even more restricted. Moreover, transience in itself is a biologically meaningful phenomenon in the context of migration stopover, since individuals migrating through the area without remaining for a stopover are de facto transients. Thus, models were not adjusted to discard transients from SOD estimates (Schaub et al. 2001). No other source of lack of fit of the general model to the data was detected (trap-dependence, Test 3.CT: all $p > 0.05$; overall fit, global tests: all $p > 0.05$, indicating absence of overdispersion).

Table 3. *Acrocephalus schoenobaenus* and *A. scirpaceus*. Goodness of fit: p-values of Test 3.SR corresponding to each species-site-year subset of data for sedge warblers (SW) and reed warblers (RW). A value < 0.05 (bold font) indicates a significant transience effect. -: data were too sparse to compute the test

Year	SW			RW		
	S29	S44	S76	S29	S44	S76
1994	0.86	0.20	-	0.84	0.87	-
1995	0.00	1.00	-	0.83	0.24	1.00
1996	0.01	0.29	-	0.31	0.42	-
1997	0.39	0.98	-	0.82	0.09	-
1998	0.01	0.80	-	0.54	0.23	-
1999	0.02	0.48	1.00	0.44	0.00	-
2000	0.96	0.88	0.50	0.27	0.01	0.08
2001	0.96	0.00	0.18	0.79	0.00	0.33
2002	0.61	0.00	1.00	0.40	0.00	0.15
2003	0.12	0.04	0.00	0.08	0.00	0.01
2004	0.03	0.05	1.00	0.09	0.75	0.51
2005	0.04	0.77	0.91	0.01	0.04	0.11

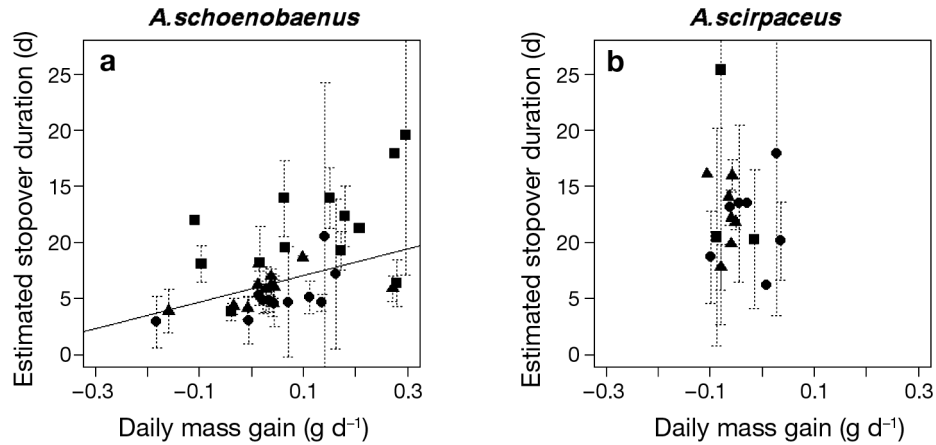


Fig. 2. *Acrocephalus schoenobaenus* and *A. scirpaceus*. Variation of stopover duration estimates with daily mass gain estimates for (a) sedge warblers (SW) and (b) reed warblers (RW). Error bars: SD of the stopover duration estimates. Unbroken line in (a): significant regression (weighted using the variance of the estimations). Sites: S76 (■), S44 (●), and S29 (▲)

Separate linear models built for each species suggested that DMG had a positive effect on SOD in SW ($r^2 = 0.31$, slope = $9.37 \pm 3.79 \text{ d}^2 \text{ g}^{-1}$; Fig. 2), but not in RW ($r^2 = 0.018$; Fig. 2). The effect of DMG was selected by the AIC-based selection procedure (Table 4, Step 3 vs. Step 2.7: >40 AIC points).

The temporal linear trend alone was strongly unlikely (>90 AIC points more than the best model). The AIC-based procedure retained August temperature as the best predictor of SOD estimates (Table 4, Steps 2.1 to 2.8: AIC differences between Step 2.7 and the others is always >18), and this effect of temperature was very likely (Step 2.7 vs. Step 1: weight ratio indicates that the model with the temperature effect is $>10^6$ times more likely than the model without). The interspecific difference in the response to temperature was retained (Table 4, Step 2.7 vs. Step 5: the model with interaction is 4.8 times more likely), but adjustment of SOD according to experienced DMG did not seem to vary among the 2 species (Table 4, Step 6 vs. Step 2.7: the model without interspecific difference is 2.2 times more likely). Adding a linear temporal trend markedly improved the fit (Step 7 vs. Step 6). The interaction between DMG and August temperatures affected SOD estimates (Step 9 vs. Step 7: model with interaction is 2.8 times more likely), and this was a stronger effect than the potential differential effect of DMG among sites (Step 9 vs. Step 8: model including DMG interaction with site was 9 times less likely than model including interaction with temperature, suggesting that among-site differences in SOD response to DMG were low).

Estimates (Table 5) indicated that the warmer the autumn and the higher the experienced DMG, the longer the stopover (Table 5: effects T_{08} and DMG). RW

Table 4. Model selection for stopover duration. *Sp*: effect of species; *S*: effect of site; *Y*: linear effect of year; DMG: linear effect of daily mass gain; T_i : linear effect of mean temperature in the *i*th month; *w*: Akaike's weight. Models are presented in the order followed in the procedure. Best models are indicated in bold. Shapiro-Wilks normality test on residuals of model $\sim S + Sp \times T_{08} + T_{08} \times DMG + Y$: $w = 0.815$, $p < 0.01$; dispersion coefficient: 1.05. AIC: Akaike's information criterion

Step	Model	AIC	w
Model without temperature effect			
1	$\sim S + Sp \times DMG$	202.5	6.85×10^{-10}
Influence of monthly temperatures per species			
2.1	$\sim S + Sp \times T_{02} + Sp \times DMG$	195.32	2.48×10^{-8}
2.2	$\sim S + Sp \times T_{03} + Sp \times DMG$	191.03	2.12×10^{-7}
2.3	$\sim S + Sp \times T_{04} + Sp \times DMG$	198.181	5.93×10^{-9}
2.4	$\sim S + Sp \times T_{05} + Sp \times DMG$	196.85	1.15×10^{-8}
2.5	$\sim S + Sp \times T_{06} + Sp \times DMG$	198.10	6.18×10^{-9}
2.6	$\sim S + Sp \times T_{07} + Sp \times DMG$	202.27	7.68×10^{-10}
2.7	$\sim S + Sp \times T_{08} + Sp \times DMG$	172.08	2.76×10^{-3}
2.8	$\sim S + Sp \times T_{09} + Sp \times DMG$	197.68	7.62×10^{-9}
Model without DMG effect			
3	$\sim S + Sp \times T_{08}$	212.44	4.75×10^{-12}
Linear temporal change			
4	$\sim Y$	255.61	2.01×10^{-21}
Most influential monthly temperature without species differences			
5	$\sim S + T_{08} + Sp \times DMG$	175.20	5.80×10^{-4}
No interspecific difference in response to DMG			
6	$\sim S + Sp \times T_{08} + DMG$	170.49	0.01
Temporal linear trend in addition to former model			
7	$\sim S + Sp \times T_{08} + DMG + Y$	163.12	0.24
Among-site differences in the effect of DMG			
8	$\sim S + Sp \times T_{08} + S \times DMG + Y$	165.49	0.07
Effect of DMG depending on temperature			
8	$\sim S + Sp \times T_{08} + T_{08} \times DMG + Y$	161.09	0.67

Table 5. Parameter estimates from the model $SOD \sim S + Sp \times T_{08} + T_{08} \times DMG + Y$. Intercept is sedge warbler *Acrocephalus schoenobaenus* (SW) and Site S29. SpRW: effect of the species reed warbler *A. scirpaceus*; S44 and S76: effect of those sites; Y: linear effect of year; DMG: linear effect of daily mass gain; T_{08} : linear effect of mean temperature in August

Parameter	Estimates	SE
Intercept	405.74742	122.96557
S44	-2.08310	0.53738
S76	-1.02143	0.67529
SpRW	-31.81739	23.72507
T_{08}	0.99588	0.18343
DMG	61.24120	30.10817
Y	-0.20867	0.06171
$SpRW \times T_{08}$	1.96604	1.26845
$T_{08} \times DMG$	-3.16255	1.71606

tended to stay longer than SW in warm autumns (interaction $SpRW \times T_{08}$ in Table 5). Values of the estimates suggested that temperature was more important when DMG was negative, and reciprocally the effect of DMG was stronger at low temperatures (interaction $T_{08} \times DMG$ in Table 5). When tested with separate linear models, the temperatures (August) had no effect on DMG (regression $DMG \sim T_i$, $p > 0.5$ in both species and at all 3 sites), showing that DMG did not mediate the response to temperature. In addition, there was a significant linear trend for SOD to decrease through years, both species staying for shorter time periods in recent years (effect Y in Table 5).

4. DISCUSSION

4.1. Inter-annual variations in the timing of migration and correlation with spring temperature

In both species sedge warblers *Acrocephalus schoenobaenus* and reed warblers *A. scirpaceus* the migration peaks occurred earlier in recent years. The best predictor of peak date was spring (March) temperature, post-nuptial migration culminating earlier in years with warm springs. This fits the prediction that spring temperatures are a better determinant of autumnal migration timing than summer or autumnal temperatures. Evidence for this phenomenon remains quite rare (Sokolov et al. 1999, Sokolov 2006).

Since both species advanced their migration date in years with warm springs, neither species takes advantage of climate warming to spend more time in their temperate breeding grounds. This supports the hypothesis that the advance in departure dates is explained by fitness benefits from early arrival at wintering grounds, rather than by temporal constraints on food availability en route. However, we acknowledge that

the latter conclusion is based on the assumption of a possible differential change in phenology of the food availability between the 2 species (Bibby & Green 1981).

The data and methods we used suffered from 3 main caveats or limits. First, at Site S76, the averaged duration of capture sessions per year was (~50%) smaller than the total duration of fall migration, and the dates of capture sessions advanced throughout the years. However, we are confident that the observed advance in peak date is the expression of a biological phenomenon and not the result of a bias due to data or methodology caveats. Simulations proved that migration peaks occurring outside the migration period actually monitored were satisfactorily retrieved by the models used for estimating peak date. The linear effects of year and temperatures were far better determinants of among-year variations in peak date than mean capture-session dates, which suggests that the capture sessions overlapped enough with the main peak to allow the models to fit the main modality in most years. Second, migration timing was (reasonably) assumed to be unimodal (the method used could estimate 1 peak date per year only), and among-age differences in migration timing (e.g. Caillat et al. 2005) were ignored. Future studies may therefore explore the use of multimodal modelling methods of migration timing. And among-year variations in migration peak date may be analysed separately per age class. It would be particularly interesting to know what degree of among-year variations in migration timing could be accounted for by the interaction between age-differential timing (cf. bimodality) of migration and among-year variations in reproductive success (cf. age ratio). Finally, the number of fitted models (14) to explain among-year variations in migration peak date was high compared to the number of statistical units (43 species-years). As stressed by Burnham & Anderson (2002), this increases the risk of selecting a 'best' model that indeed has a weak explanatory power. Future tests of our predictions with datasets from other sites and species will indicate how robust and general our conclusions are.

4.2. Stopover duration and correlates

Among-site, -year and -species variations in SOD estimates were best explained by DMG, August temperatures and linear year effect. DMG did not depend on temperature. This suggests that the advance in migration timing is sufficient for the birds not to experience any decrease in food availability at stopover sites (which would have resulted in a decrease in DMG and SOD with increasing spring temperature; Hedenström et al. 2007). Even if an advance in the timing of

food abundance en route is not the driving force of the advanced timing of migration, this advance seems sufficient to track the possible changes in the phenology of autumn food resources and to avoid any decrease in the experienced food abundance. If the advance in peak date with increasing T_{03} had not been sufficient, birds (and especially SW) would potentially have experienced a decrease in autumn food resources and we should have observed a decrease of DMG and SOD with T_{03} . This decrease is not observed, suggesting that birds efficiently adapted to the newly experienced conditions. This interpretation lays on the assumption that there is a modification of food resource phenology. Even if this assumption is not verified, our results still support that recent climate warming did not affect the stopover duration.

The fact that the relationship between SOD and DMG is positive is in agreement with theoretical work on time-minimising migrants using a discrete environment (Alerstam & Lindström 1990, Hedenström et al. 2007). The significance threshold reached in SW only brings to the fore the influence of diet involved in our second hypothesis: SW was predicted to be more constrained than RW and, as a result, to respond more strongly to variations in its experienced DMG, because when the resource peaks somewhere, it is already scarce further south. This result is in agreement with previous findings by Schaub & Jenni (2001) that between-site variation of SOD is much greater for SW than for species with a more reliable food resource. However, when analysing both species with a single model, the interspecific difference was not significant. The much higher variability of DMG in SW than in RW may mask this difference.

4.3. Observed temporal trend: Phenotypic plasticity or natural selection?

For both species, migration timing is better predicted by a physical variable (March temperature) than by a linear year effect alone. Among-year adjustment of migration timing to temperature conditions suggests that the observed changes in migration behaviour would be essentially plastic (Jonzén et al. 2006). Since no temporal linear change in migration date is found after correction for temperature, then directional selection for earlier migration either does not operate, or is too weak to be detected. The influence of August temperatures on migration timing (through its interaction with March temperature) and on SOD further support that SOD and migration date are adjusted to experienced conditions thanks to phenotypic plasticity rather than by natural selection—August temperatures not being correlated to year.

However, both species show a significant decrease in stopover duration with year that is explained neither by DMG nor by temperatures. This result is more difficult to interpret. The linear year effect might integrate some other factors that we did not include in this analysis. Since DMG did not depend on year or on temperature, we do not think that site quality decreases with year.

Particularly for SW, the DMG might be the integrative clue used by individuals to plastically adapt their stopover duration to local conditions. DMG is much more variable in this species, and this may be due to its specialised diet (Bibby & Green 1981, Schaub & Jenni 2001). As our results suggest that the advance in migration timing does not occur in response to the temporal shift in food availability, mistiming between migration and appearance of the aphid peak is possible in the coming years. The future evolution of the system will allow further testing of this hypothesis. We will need to quantify the temporal variations in aphid abundance and their relationships with spring and autumn temperatures, as well as with the diet and stopover duration of both warblers in relationship to aphid availability.

Acknowledgements. We warmly thank all ringers who collected data, and F. Jiguet and J.-M. Moisselin (Meteo France) for providing temperature data. Ringing licenses were delivered by CRBPO, Muséum National d'Histoire Naturelle. We are grateful to 4 anonymous reviewers for their constructive comments, and to R. Choquet and O. Gimenez for their help in data analysis.

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Submitted: April 28, 2007; Accepted: October 30, 2007

Proofs received from author(s): December 14, 2007