

Challenging claims in the study of migratory birds and climate change

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ABSTRACT

Recent shifts in phenology in response to climate change are well established but often poorly understood. Many animals integrate climate change across a spatially and temporally dispersed annual life cycle, and effects are modulated

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by ecological interactions, evolutionary change and endogenous control mechanisms. Here we assess and discuss key statements emerging from the rapidly developing study of changing spring phenology in migratory birds. These well-studied organisms have been instrumental for understanding climate-change effects, but research is developing rapidly and there is a need to attack the big issues rather than risking affirmative science. Although we agree poorly on the support for most claims, agreement regarding the knowledge basis enables consensus regarding broad patterns and likely causes. Empirical data needed for disentangling mechanisms are still scarce, and consequences at a population level and on community composition remain unclear. With increasing knowledge, the overall support ('consensus view') for a claim increased and between-researcher variability in support ('expert opinions') decreased, indicating the importance of assessing and communicating the knowledge basis. A proper integration across biological disciplines seems essential for the field's transition from affirming patterns to understanding mechanisms and making robust predictions regarding future consequences of shifting phenologies.

Key words: bird migration, climate change, phenology, annual life cycle, match-mismatch, endogenous control, phenotypic plasticity, microevolutionary change, population trends, integrative biology.

CONTENTS

I. Introduction	2
II. Changes in spring migration	3
(1) Claim 1: birds advance their spring migration in response to climate change	3
(2) Claim 2: phenological response to climate change depends on migratory distance	4
(3) Claim 3: climate change affects migration distance and routes	4
III. Mechanisms underlying observed changes	5
(1) Claim 4: mechanisms controlling the timing of migration are generally hardwired	5
(2) Claim 5: changes in the timing of migration are mainly due to phenotypic plasticity	6
(3) Claim 6: phenotypic variation in timing of arrival is mainly due to weather conditions <i>en route</i>	6
(4) Claim 7: responses to climate change are constrained by the annual cycle	7
IV. Consequences of climate change	8
(1) Claim 8: migratory birds suffer from increased trophic mismatch on the breeding grounds	8
(2) Claim 9: climate change causes population declines in migratory birds	9
(3) Claim 10: climate change affects community composition	9
V. Claims, knowledge and support	10
VI. Conclusions	12
VII. Acknowledgements	13
VIII. Appendices	13
(1) Data on timing of bird migration	13
(2) Perspectives: searching for generality in the phenology of migratory birds	14
IX. References	14
X. Supporting Information	19

I. INTRODUCTION

Recent climate change has shifted patterns of abiotic factors playing key roles in structuring species abundance and geographical dispersion, and their variability over time (Gitay *et al.*, 2002; Stenseth *et al.*, 2002; Walther *et al.*, 2002; IPCC, 2007). A failure of species to respond sufficiently to the new environmental conditions may result in population declines or even species extinctions (Thomas *et al.*, 2004; Parmesan, 2006). The timing of life-history events with respect to a fluctuating environment can be crucial for survival and successful reproduction, and classical life-history theory predicts that individuals should optimize the match in space and time with important resources (Lack, 1968; Roff, 2002). Hence, it is not surprising that shifts in the seasonal timing of life-history events (i.e. phenology) are

among the most well-established effects of climate change (Parmesan & Yohe, 2003; Rosenzweig *et al.*, 2008). It is, however, the combined effects of spatial changes in species distributions and changes in timing, relative to those of interacting species, that will determine the viability of populations in light of climate change and other human impacts, such as habitat and landscape changes. There is so far no general understanding of how phenological impacts of climate change integrate over the course of a seasonal life cycle, and how this translates to individual life histories and population fluctuations. Furthermore, transitional phases in the life histories of animals may involve movements between distant areas, making it hard to pinpoint phenological events and climatic influences in time and space.

Here we focus on migratory birds, which have fascinated humans since ancient times and nourished a vigorous

research field over several centuries (Berthold, 2001). Bird migration has presumably evolved repeatedly as a response to seasonality in environmental conditions, resulting in a lifestyle where the spatial and temporal schedule of events over the annual cycle represents an overall optimal match with the environment (Newton, 2008). There is a long tradition for using birds as model organisms for studying impacts of climate on animals, and changes in the timing of migration and breeding in birds are indeed among the best documented responses to climate change (Parmesan & Yohe, 2003; Møller, Fiedler & Berthold, 2004b; Rubolini *et al.*, 2007b). We see this as a good area for seeking in-depth understanding of phenological changes.

In this rapidly developing, but still fragmented field of research, a number of claims and assumptions have been made, not all of which may be as well justified as we believe. In order to improve our understanding of the total picture and avoid the pitfalls of affirmative science, we review and critically synthesize this literature. We focus particularly on issues pertaining to timing of spring migration—i.e. the transition from the non-breeding period *via* migration to the arrival on breeding grounds. Climate-change effects are most consistent and well studied for this phase, and fitness effects are highly likely through the close association with onset of reproduction. We here list what we consider the more important key statements, discuss their basis and subjectively score their support and how much has been done to check their validity. We identify gaps in knowledge and make suggestions on how to proceed in order to improve our scientific knowledge and predictive power regarding these issues.

II. CHANGES IN SPRING MIGRATION

(1) Claim 1: birds advance their spring migration in response to climate change

Recent decades have seen a wealth of studies on long-term trends in timing of spring migration and arrival at breeding grounds in relation to changing climate. Extensive reviews show a general pattern of advancement (Lehikoinen, Sparks & Zalakevicius, 2004; Gordo, 2007; Rubolini *et al.*, 2007b; Lehikoinen & Sparks, 2010), irrespective of the specific metric considered (Appendix 1). However, estimated rates of advancement vary according to a number of factors, including species or taxonomic group, life-history traits and geographic region (Butler, 2003; Lehikoinen *et al.*, 2004; Gordo, 2007; Rubolini *et al.*, 2007b; Végvári *et al.*, 2009; Lehikoinen & Sparks, 2010). Also, trends are not constant over time, and non-linear responses to temperature have been suggested (Gordo & Sanz, 2006; Askeyev, Sparks & Askeyev, 2009). Most studies do not consider different responses between sexes or age groups (see online supporting information, Appendix S1).

Within species or populations, advancement is typically most pronounced early in the season (Lehikoinen *et al.*,

2004; Jonzén *et al.*, 2006; Lehikoinen & Sparks, 2010). Two meta-analyses of a large number of European time series (Lehikoinen *et al.*, 2004; Rubolini *et al.*, 2007b) show a mean advancement of approximately 0.4 days year⁻¹ for first arrival dates, compared to 0.1–0.2 days year⁻¹ for mean or median migration dates. This might reflect a larger impact of climate on early migrating individuals (Vähätalo *et al.*, 2004), as well as sampling biases and statistical properties for first arrival dates (Appendix 1), but the general pattern is consistent with studies using more reliable metrics. Even though there is non-negligible variation both among regions and species or populations (Peñuelas, Filella & Comas, 2002; Lehikoinen *et al.*, 2004; Rubolini *et al.*, 2007b; Lehikoinen & Sparks, 2010), the patterns of overall advancement emerging from studies of North American and Australian birds are broadly similar to those in Europe (Mills, 2005; Murphy-Klassen *et al.*, 2005; Beaumont, McAllan & Hughes, 2006; Miller-Rushing *et al.*, 2008b).

Three lines of observational evidence link observed changes to changes in climatic conditions. (1) Many studies have shown clear and predictable relationships between bird migration and climate variables, including both local weather variables (see also Claim 6, Section III.3) and climatic indices such as the North Atlantic Oscillation (NAO) index (Stenseth *et al.*, 2003; Lehikoinen *et al.*, 2004; Gordo, 2007; Lehikoinen & Sparks, 2010). These findings typically persist after accounting for potentially confounding effects of long-term trends (Jonzén *et al.*, 2006, 2007; Rubolini *et al.*, 2007a; Saino *et al.*, 2007), and reversal in temperature trends between cooling and warming periods during the last century is matched by corresponding delays and advances in migration phenology (Lehikoinen *et al.*, 2004). (2) Geographical and between-species patterns of variability in phenological trends support the notion of climate-induced changes. Species differ in their movements and ecology, and climate change is not uniform (Gitay *et al.*, 2002; Stöckli & Vidale, 2004; IPCC, 2007; Karlsen *et al.*, 2009; Sparks *et al.*, 2009). Hence, we do not expect trends to be equal across species and geographic regions (Ahola *et al.*, 2004; Both, Bijlsma & Visser, 2005; Hüppop & Winkel, 2006; Askeyev *et al.*, 2007, 2010; Both & te Marvelde, 2007). (3) Studies of individually marked populations are able to link variability among individuals and years to climatic conditions experienced in wintering areas or during migration (Saino *et al.*, 2004; Gunnarsson *et al.*, 2006). Good individual-level data on arrival are, however, hard to obtain for most species (Appendix 1), and we are only aware of two studies examining phenotypic plasticity in the timing of passerine migration in the wild, showing that individual barn swallows (*Hirundo rustica*) advance arrival dates in years with beneficial environmental conditions on the wintering (Saino *et al.*, 2004) or staging (Ballbontín *et al.*, 2009) grounds.

In sum, there is strong evidence that migratory birds have advanced their spring migration and arrival during the last decades in a manner consistent with climate change, but there is large and poorly understood variability among species, and mechanisms underlying change need to be elucidated further. It should also be noted that most data are

from passage areas or close to breeding grounds; we often know very little about departure dates from wintering areas in, for instance, Africa.

(2) Claim 2: phenological response to climate change depends on migratory distance

Long-distance migrants, particularly those crossing the Sahara desert, are often claimed to show a smaller phenological response to climate change than short-distance migrants (Berthold, 1991; Both & Visser, 2001; Butler, 2003; Pulido & Widmer, 2005). This assertion is partly based on the expectation of strong stabilizing selection on the timing of arrival (Claim 8, Section IV.1) having resulted in rigid endogenous control of migration schedules synchronized by, e.g. photoperiod for long-distance migrants (Berthold, 1996; Gwinner, 1996), while the proximity of wintering and breeding areas for short-distance migrants allows adjustment of departure and arrival schedules according to weather conditions at their destination (Claim 6, Section III.3).

Rigid endogenous control could result in reduced flexibility of migration timing in response to climatic factors, i.e. canalization of the phenotype (Pulido & Widmer, 2005). To the best of our knowledge, explicit comparisons of endogenous control between short- and long-distance migrants have not been performed, although for a few species there is experimental evidence suggesting greater between-individual variability in duration and intensity of migratory activity in short-distance migrants (Gwinner, 1968; Berthold *et al.*, 1972). Stronger stabilizing selection on migratory traits of long-distance migrants has been repeatedly advocated, but evidence is indirect (Pulido & Widmer, 2005), and the strength of selection on arrival dates is very rarely estimated (Møller *et al.*, 2009a). Some studies indicate larger interspecific variability in trends among European short-distance compared to long-distance migrants (Jonzén *et al.*, 2006; Rubolini *et al.*, 2007b; Rubolini, Saino & Møller, 2010), but this could also be due to the difference in overall timing, since weather (and hence migration or arrival dates) can be more variable early in spring (Loxton & Sparks, 1999; Newton, 2008; Wilson, 2009). There is no consistent pattern of short-distance migrants responding more strongly to favourable conditions (i.e. high temperature and NAO values) (Forchhammer, Post & Stenseth, 2002; Hüppop & Hüppop, 2003; Hubálek, 2004; Stervander *et al.*, 2005; Jonzén *et al.*, 2006; Gienapp, Leimu & Merilä, 2007). Moreover, because long-distance migrants encounter a larger spectrum of environmental conditions *en route*, the comparison between short- and long-distance migrants can be biased, due to issues such as statistical power and possibly opposite effects in different parts of the migration route.

The evidence for differential trends in spring migration phenology according to migration distance is somewhat mixed; while clearly most studies have reported larger advancement in short-distance migrants (Butler, 2003; Lehikoinen *et al.*, 2004; Rubolini *et al.*, 2007b, 2010; Miller-Rushing *et al.*, 2008b; Lehikoinen & Sparks, 2010), others did not find any difference (Hüppop & Hüppop, 2003;

Zalakevicius *et al.*, 2006), or even greater advancement in long-distance migrants (Stervander *et al.*, 2005; Jonzén *et al.*, 2006). Meta-analyses indicate overall larger advances for short-distance migrants in North America (Gienapp *et al.*, 2007), but show inconsistent results for Europe (Gienapp *et al.*, 2007; Rubolini *et al.*, 2007b). Studies also differ in terms of temporal scope and species composition, and environmental constraints on the progress of migration (Both & te Marvelde, 2007; Gordo, 2007; Newton, 2008). In addition, relevant trends in abiotic and biotic conditions (Menzel, 2000; Stöckli & Vidale, 2004; Sparks *et al.*, 2009) are likely to be different for short-distance and the later long-distance migrants.

In sum, even though differences are likely to exist, no firm patterns of differential level of response to climate according to migratory distance can be established so far. Case studies and meta-analyses should more properly account for confounding factors and effects of study design.

(3) Claim 3: climate change affects migration distance and routes

It has frequently been hypothesized that changes in spring arrival dates are due to changes in the timing or speed of migration (Gwinner, 1986; Pulido, 2007a). However, range shifts (Thomas & Lennon, 1999; Böhning-Gaese & Lemoine, 2004; Hitch & Leberg, 2007; La Sorte & Thompson, 2007; Maclean *et al.*, 2008; Zuckerberg, Woods & Porter, 2009; Brommer & Møller, 2010) and some route changes (Sutherland, 1998; Fiedler, 2003; Newton, 2008) are well documented, and may provide additional pathways for adaptive change in phenology (Pulido & Berthold, 2004; Pulido, 2007a; Coppack *et al.*, 2008; Visser *et al.*, 2009b).

Climate change is likely to affect movement patterns in at least three ways. Firstly, the propensity of individuals to migrate can change over time. Reviews show ample observational evidence (from case studies of diverse bird orders) of migratory birds becoming more sedentary, i.e. wintering closer to their breeding grounds and abandoning the more distant parts of their former wintering range (Sutherland, 1998; Fiedler, 2003; Newton, 2008). Improved feeding conditions, due to mild winters and in some cases possibly additional feeding by humans, is the most likely explanation. Selection for shorter migration distance can result in rapid evolution of residency, as suggested by a common garden experiment with blackcaps (*Sylvia atricapilla*) (Pulido & Berthold, 2010).

Secondly, changes in both breeding and wintering ranges are well documented (Sutherland, 1998; Böhning-Gaese & Lemoine, 2004; Brommer, 2004; Hitch & Leberg, 2007; La Sorte & Thompson, 2007; Maclean *et al.*, 2008; Newton, 2008; Zuckerberg *et al.*, 2009). At intermediate and northern latitudes, observed range shifts are mostly (breeding range) or almost always (winter range) northwards or towards areas of initially harsher climate, suggesting they are driven by recent climate amelioration. Although rates of distributional change depend on whether the change is measured at the boundaries or in the interior of a species' geographic distribution, and

southern range margins of northerly breeding species might not have shifted (Brommer & Møller, 2010), there are indications that shifts in winter ranges of migratory birds occur at rates at least as high as shifts in summer ranges (Thomas & Lennon, 1999; Devictor *et al.*, 2008; Maclean *et al.*, 2008). Furthermore, there are more indications of shortening than lengthening of migratory routes (Sutherland, 1998; Fiedler, Bairlein & Koppen, 2004; Newton, 2008; Visser *et al.*, 2009b). On the other hand, species distribution models coupled with climate-change scenarios suggest a larger and more consistent shift in northern breeding ranges (Huntley *et al.*, 2007; Barbet-Massin *et al.*, 2009; Doswald *et al.*, 2009), while wintering areas in the Southern Hemisphere may become shifted to the south (Barbet-Massin *et al.*, 2009); i.e. to higher latitudes, as in the Northern Hemisphere. Several species known for their trans-Saharan migrations now overwinter in Spain and other parts of southern Europe (examples in Newton, 2008). Geographical barriers may prevent gradual range shifts—for instance, the Sahara desert restricts available wintering areas for Palearctic long-distance migrants (Pulido & Berthold, 2004; Pulido, 2007a).

Thirdly, migration routes are shaped by biogeographic history and can be less than optimal in terms of complexity and travelled distance (Sutherland, 1998; Newton, 2008; Bensch *et al.*, 2009). For most species of long-distance migrants, there is poor knowledge of the migratory connectivity (i.e., the linkage of populations between wintering, migration and breeding; see online supporting information, Appendix S2). In the absence of severe geographical constraints there could be potential for adaptive change in migration routes. For instance, microevolution of a novel migratory direction towards W–NW within a time scale of a few decades was demonstrated for blackcaps breeding in Central Europe and previously wintering along the Western Mediterranean (Berthold *et al.*, 1992).

Phenological consequences of spatial change in migration patterns are so far little studied, but are likely to be important (Pulido, 2007a), not the least in light of range shifts predicted by climate envelope models (Huntley *et al.*, 2007). While a reduction in migratory distance seems likely to result in advancing arrival (Coppack & Pulido, 2004; Pulido, 2007a), we are not aware of any attempts to model rigorously the implications for migration phenology. This task is likely to be hard, since it is not clear how to separate and model various pathways for phenological change in any particular observed case, and phenological responses are likely to be constrained by seasonal interactions between stages of the annual cycle (Claim 7, Section III.4).

III. MECHANISMS UNDERLYING OBSERVED CHANGES

(1) Claim 4: mechanisms controlling the timing of migration are generally hardwired

It has been claimed that the main reason underlying observations of insufficient adjustment of the timing of spring

arrival in long-distance migrants to changing phenology on the breeding ground is the control of migration by a rigid endogenous migration programme (Both & Visser, 2001; Coppack & Both, 2002).

This argument is based on empirical findings under laboratory conditions, with some support from field studies. For a number of study species, the timing of gonadal growth, moult, fat deposition and migratory activity has been shown to be controlled by the circannual rhythms of an endogenous programme (Gwinner, 1986, 1996; Berthold, 1996), which under controlled laboratory conditions appear largely insensitive to perturbations in environmental factors such as precipitation, temperature, and (to some extent) food availability (Berthold, 1984; Gwinner, 1996). These rhythms are strongly influenced by photoperiod, i.e. day length *per se* and changes in day length, and recent studies suggest high heritability (Helm & Visser, 2010) and linkage to geographically distinct annual scheduling (Helm, Schwabl & Gwinner, 2009). Photoperiod has been suggested to be the most reliable predictive cue used by birds to initiate migration both on the breeding and on the wintering grounds (Kok, Van Ee & Nel, 1991; Coppack & Both, 2002; Coppack & Pulido, 2004). This could be particularly so for long-distance migrants wintering in the tropics, and observations of smaller variation in spring arrival times for these species compared to short-distance migrants (Mason, 1995; Lehikoinen *et al.*, 2004; Jonzén *et al.*, 2006; Rubolini *et al.*, 2007b) could indicate stronger endogenous control (see also Claim 2, Section II.2).

On the other hand, the timing of migration may be constrained by the environment in addition to endogenous mechanisms. Long-term studies on the American redstart (*Setophaga ruticilla*) show that departure schedules from wintering grounds are affected by habitat occupancy *via* differences in food abundance, which is linked to rainfall (Marra, Hobson & Holmes, 1998; Studds & Marra, 2005, 2007). Migration requires fuel deposition and for some species completion of winter moult, and a poor food supply can delay the rate of both and hence departure dates (Saino *et al.*, 2004; Newton, 2008). Day length varies little near the equator, and intraspecific variation in the phenological response of moult and migration (and for migratory restlessness in non-migratory populations; Helm & Gwinner, 2006) to photoperiod has been described for many species (Berthold, 1996). In the blackcap, additive genetic variation for this response has been found, though it is unclear how large it is (Coppack, Pulido & Berthold, 2001). A comparison of the timing of autumn migration for two species under laboratory conditions suggests that small phenotypic variability in long-distance migrants is not due to differences in additive genetic variation, but possibly a result of environmental forcing resulting in similar phenotypes (environmental canalization; Appendix S2, Pulido & Widmer, 2005).

In some groups of large, long-living birds (cranes, geese, swans and storks), migratory routes and stopover sites are predominantly culturally transmitted in autumn. This has

been hypothesized to facilitate rapid adaptive changes in wintering areas or migration routes (Sutherland, 1998; van Noordwijk *et al.*, 2006; Newton, 2008). Although these species also likely possess a spatio-temporal programme controlling their annual cycle (Chernetsov, Berthold & Querner, 2004; Newton, 2008), it can be overridden or modified by environmental input and individual experience. Similarly, in species with a rigid circannual organization (and apparently hard-wired behaviour), field evidence suggests that the endogenous programme strongly determines migratory route and phenology mainly on the first migratory trip from the breeding to the non-breeding area. Thereafter, birds appear to rely more on their experience and on environmental information (Berthold, 2001; Mouritsen, 2003; Pulido, 2007*b*; Thorup *et al.*, 2007; Chernetsov, Kishkinev & Mouritsen, 2008).

Hence, although there is strong support for endogenous control of the timing and spatiotemporal progress of bird migration in at least some species, it is unclear how rigid this control is in the wild. A lack of response to climate in migration phenology need not imply a rigid program, but could also be due to constraints pertaining to the annual cycle (Claim 7, Section III.4) or genetic architecture (Appendix S2).

(2) Claim 5: changes in the timing of migration are mainly due to phenotypic plasticity

Adaptive adjustment of a population to climatic changes can be a combination of evolutionary change and individual phenotypic adjustment (i.e. phenotypic plasticity, including trans-generational maternal effects). The relative importance of the two processes has been debated and is critical for rates and limits of phenotypic change. In principle, changes due to phenotypic plasticity will be faster, but might be rather limited. Adaptive evolution is likely to be slower, but can in theory continue until additive genetic variation is depleted (Pulido & Berthold, 2004).

In general, it has been assumed that phenotypic plasticity is sufficient for explaining the observed changes in spring arrival. A number of empirical results suggest the importance of phenotypic plasticity: (1) in barn swallows, population variability in arrival dates is partly a consequence of individual responses to environmental conditions in winter quarters (Saino *et al.*, 2004) and *en route* (Balbontín *et al.*, 2009). (2) Established correlations between weather *en route* and at arrival (Claim 6, Section III.3) also are supported for populations of intensively studied species (Ahola *et al.*, 2004; Both *et al.*, 2005; Hüppop & Winkel, 2006; Both & te Marvelde, 2007). (3) Meta-analyses of rates of phenotypic change in response to climate change suggest phenotypic plasticity rather than adaptive evolution (Gienapp *et al.*, 2007, 2008; Hendry, Farrugia & Kinnison, 2008), and changes in the timing of breeding seem to be due to phenotypic plasticity (Pulido & Berthold, 2004; Charmantier *et al.*, 2008). (4) Laboratory studies indicate plasticity in timing related to photoperiodic cues. Several long-distance migrants showed earlier onset of migratory activity when subjected to photoperiods of more

northern areas, suggesting that a change in wintering latitude could induce profound changes in the annual schedule (Coppack & Pulido, 2004; Coppack *et al.*, 2008).

However, this does not exclude the possibility of evolutionary change (Pulido & Berthold, 2004; Gienapp *et al.*, 2007; Pulido, 2007*b*), and it has been claimed that observed changes in spring arrival may reflect an evolutionary response to changing environmental conditions (Jonzén *et al.*, 2006, 2007). Various studies show that rapid evolutionary change is possible and expected (Brown & Brown, 2000; Pulido & Berthold, 2004, 2010; Møller, 2007*b*; Pulido, 2007*a*). In order properly to demonstrate adaptive evolutionary responses to climate we need to show that (1) there is phenotypic variation, (2) this variation is inherited, (3) fitness is largely determined by phenotypic variation, and (4) this variation in fitness is caused by changes in meteorological variables (Gienapp *et al.*, 2008). Phenotypic variation in migratory traits is evident from field studies, but these primarily concern variation in arrival or migration (not departure) dates among sex- and age-classes, populations and species. In order to assess the relative influence of genetic and environmental effects on phenotypic variation, we need information on between- and within-individual variability, which still is scarce (van Noordwijk *et al.*, 2006; Pulido, 2007*b*). Heritability of migratory traits has been established (Pulido & Berthold, 2003; Pulido, 2007*b*), but only for few populations and species, and mostly in laboratory experiments. Hence, it remains unclear how important genetic variation is in the expression of phenotypic variation of migratory traits in the wild. Selection pressures in terms of consequences of arrival date for survival, mating opportunities, access to food and territories are well identified (Claim 8, Section IV.1; Kokko, 1999), but there is a lack of empirical data on actual consequences of climate change. Selection on the timing of migration has been demonstrated in the wild (Brown & Brown, 2000; Møller, 2007*b*), but we are only aware of one study (Møller, 2007*b*) linking selection on arrival time with climatic variables, and trying to separate between fitness effects of the timing of arrival and the timing of breeding.

In sum, empirical support has so far favoured phenotypic plasticity as the main mechanism underlying recent changes in the timing of migration. However, adaptive responses to selection pressures are also expected and observed, though evolutionary change is hard to demonstrate. More needs to be done to evaluate the relative importance of both processes.

(3) Claim 6: phenotypic variation in timing of arrival is mainly due to weather conditions *en route*

The effect of weather conditions experienced during migration on trends and between-year variability in arrival dates is important to assess, since it sheds light on the potential for migratory birds to cope with climate change by individual plasticity or evolutionary change in the timing of migration (Pulido, 2007*a*). For instance, the advancing spring arrival of the song thrush (*Turdus philomelos*) to the southeast Baltic over the past 40 years was mainly explained by increasing

tailwind frequency and temperatures *en route* (Sinelschikova *et al.*, 2007).

Apart from the recent focus on correlations between arrival date and local weather (Claim 1, Section II.1), it is well known from the literature that weather variables affect observable migration intensity (Richardson, 1978; Berthold, 1996; Newton, 2008), as well as flight speed and stop-over decisions during migration (Hedenström & Ålerstam, 1995; Hedenström, 2008). Many species apparently respond rather similarly and possibly in a simple threshold manner, to a limited number of weather variables governing departure and flight conditions—such as wind, precipitation, temperature, and visibility of celestial cues (Åkesson *et al.*, 2001; Erni *et al.*, 2002; Cochran *et al.*, 2005; Newton, 2008).

The question of whether the cumulative effect of weather *en route* is sufficient to explain the observed changes in migration timing needs to be considered in light of variability in temporal trends along the migratory route, and species- and individual-specific migration strategies and constraints. Increasing temperatures at intermediate and high latitudes are expected to improve conditions for migration and increase food availability (hence potentially decrease stop-over durations) due to advancing phenologies of vegetation and food resources (Gordo, 2007), while effects of climate change in the tropics are less clear (Hulme *et al.*, 2001; Herrmann, Anyamba & Tucker, 2005; Bell, 2007; Robson & Barriocanal, 2010). Due to spatially correlated weather and the general lack of knowledge of wintering areas and pre-migratory movements in most species (Newton, 2008), it is hard to separate the effects of earlier departure (Marra *et al.*, 1998; Gordo *et al.*, 2005; Jonzén *et al.*, 2006; Saino *et al.*, 2007; Studds & Marra, 2007; Gordo & Sanz, 2008) and increased migration speed (Marra *et al.*, 2005). The role of spatiotemporal correlation patterns of environmental variables along the flyway has only recently been studied in detail (Marra *et al.*, 2005; van der Graaf *et al.*, 2006; Saino & Ambrosini, 2008), and might explain differential trends in timing along different flyways (Tøttrup *et al.*, 2008). Recent case studies suggest the importance of fine-tuning the progress of migration in response to the environment *en route* (Saino & Ambrosini, 2008; Tøttrup *et al.*, 2010), and such effects could mask phenotypic plasticity in the onset of migration (Both, 2010).

Adjustment of migratory movements based on short-range meteorological correlations has frequently been advocated for short-distance migrants (Hötter, 2002; Newton, 2008), while it has been argued that long-distance migrants have no available cues for predicting conditions at the breeding grounds when they are in the wintering area or at distant staging sites (Cotton, 2003; Lehikoinen *et al.*, 2004; Gordo *et al.*, 2005). However, due to global atmospheric circulation patterns such as those underlying the NAO (Appendix S2), meteorological conditions in breeding areas of Northern Europe and those in the central and eastern Mediterranean correlate negatively, implying close to zero correlation (and hence poor opportunity to predict weather conditions at destination) at intermediate latitudes (Saino & Ambrosini,

2008). Moreover, temperature anomalies in the Sahel (a major wintering/stop-over region for long-distance migrants (Newton, 2008) just before the start of spring migration are correlated with those in Europe in the following two months (Saino & Ambrosini, 2008). Hence, migrants might be able to tune both the onset and progression of migration based on *en route* prediction of meteorological conditions ahead. This calls for further consideration of the spatiotemporal organization of migratory journeys and integration with optimal migration theory (see online supporting information, Appendix S3).

It is beyond doubt that the timing of arrival can be affected by weather conditions *en route*, but data are still insufficient to evaluate properly the relative magnitude of this effect for individuals, except for a few special cases (see, e.g. Shamoun-Baranes *et al.*, 2003; Thorup *et al.*, 2006; Strandberg & Ålerstam, 2007). More research is also needed to separate between various direct (e.g. tailwinds) and indirect (e.g. enhanced food availability) effects of local weather *en route*, and to assess effects of migratory strategy and distance (Claim 2, Section II.2).

(4) Claim 7: responses to climate change are constrained by the annual cycle

Since the link between winter habitat conditions and timing of arrival to breeding grounds was made explicit at the level of individuals using colour marking and stable isotopes (Marra *et al.*, 1998; Gill *et al.*, 2001; Gunnarsson *et al.*, 2005), there has been increasing awareness of constraints imposed by the annual cycle on migration and arrival phenology (Carey, 2009). Particular interest has been directed at long-distance migrants, whose annual schedule incorporates very different environments, and who are especially time-constrained due to prolonged migration periods, also requiring fuel deposition on top of the usual requirements for breeding and moulting (Hedenström, 2006, 2008).

Constraints imposed by the annual cycle can partly be understood in terms of interactions between the wintering, migration and breeding stages (seasonal interactions; Appendix S2). Effects of factors in one stage can cascade to the next stage, i.e. carry-over effects (Marra *et al.*, 1998; Runge & Marra, 2005; Pulido, 2007*a*; Newton, 2008; Møller, Flensted-Jensen & Mardal, 2009*b*). More generally, since migratory birds use different areas at different stages of their annual cycle, and climate change varies across space and time, they might not be able to adapt to all important environmental changes encountered. This can have a genetic component, reflecting the inability of populations to evolve particular trait combinations due to the lack of sufficient genetic variation in a multi-trait character space and/or due to unfavourable genetic correlations (Appendix S2). Also, constraints can be due to the inability to change a trait if particular ecological conditions at the preceding stages are not met, or if the change induces a penalty at a later stage. If, for instance, conditions on the breeding grounds favour earlier arrival, but conditions at the wintering sites or along the migratory route have not changed, birds may not be able to advance their arrival since they will not find the food

needed to build up energy reserves required for migrating earlier or faster (Strode, 2003; Hüppop & Winkel, 2006; Both & te Marvelde, 2007). On the other hand, if there is environmentally induced change in one life-cycle stage but not in later stages, a mismatch between environmental conditions (e.g. food availability; Claim 8, Section IV.1) and the life cycle can arise. As an example, pied flycatchers (*Ficedula hypoleuca*) breeding in Finland are arriving earlier on the breeding grounds, due to improved conditions during migration, but do not breed earlier since conditions on the breeding grounds have not advanced—hence experiencing less favourable conditions upon arrival and a prolonged pre-breeding period (Ahola *et al.*, 2004). Changes in arrival on the breeding grounds could be constrained by a range of abiotic factors during migration (Claim 6, Section III.3) or wintering (Marra *et al.*, 1998; Saino *et al.*, 2004; Norris, 2005; Studds & Marra, 2007), and may also be determined by shifting winter ranges (Coppack & Pulido, 2004; Pulido, 2007a; Claim 3, Section II.3).

Apart from constraints pertaining to post-breeding moult, the issue of flexibility in scheduling of the annual cycle has received little attention. In general, flexibility in timing is expected to relate inversely to migratory distance and body size (Hedenström, 2006, 2008), as well as to the number and spacing of stages and substages in the annual cycle (Wingfield, 2008). Climate change may affect the latter, for instance by earlier arrival allowing extra clutches or more time between clutches (Møller, 2007a; Najmanová & Adamík, 2009; Møller *et al.*, 2010a). On the other hand, ontogenetic effects on the timing of migration are also little explored. An effect of hatching date on the timing of subsequent spring migration has been suggested for pied flycatchers (Both, 2010), and spring arrival of Arctic terns (*Sterna paradisea*) correlated positively with breeding date the previous year (Møller *et al.*, 2009b). For birds migrating long distances non-stop, the link between breeding latitude and departure dates from wintering quarters, as well as subsequent return migration, can be remarkably strong (Conklin *et al.*, 2010).

Overall, potential constraints on phenological responses due to the interaction of events across periods of the annual cycle, in particular carry-over effects, are well identified, but there is little empirical knowledge about their consequences for responses to climate change. Further progress also needs to be made regarding ways to integrate such constraints in modelling efforts (Appendix S3).

IV. CONSEQUENCES OF CLIMATE CHANGE

(1) Claim 8: migratory birds suffer from increased trophic mismatch on the breeding grounds

A mere change in the timing of migration does not necessarily mean a proper response in context of the birds' breeding environment (Visser & Both, 2005; Goodenough, Hart & Stafford, 2010; Saino *et al.*, 2010). If timing of arrival to breeding grounds changes too rapidly or too slowly, the result could be an increased temporal mismatch with food

resources (Inouye *et al.*, 2000; Sanz *et al.*, 2003; Strode, 2003; Pearce-Higgins, Yalden & Whittingham, 2005; Visser & Both, 2005). This might lead to reduced fitness (Both *et al.*, 2006; Visser, Holleman & Gienapp, 2006), and to a non-adaptive response to climate change (Both & Visser, 2001; Coppack & Both, 2002).

For species breeding at intermediate and high latitudes, arriving early in spring could be costly, due to low food abundance and periods of cold weather causing mass mortality (Brown & Brown, 2000; Newton, 2007) or temporary return migration (Vepsäläinen, 1968; Richardson, 1978). Arriving too late may also have fitness consequences due to poor matching with peak food abundance (Both *et al.*, 2009) and difficulties in acquiring high-quality breeding sites and/or partner(s) (Alatalo, Lundberg & Glynn, 1986; Smith & Moore, 2005; Sergio *et al.*, 2007; Newton, 2008). Fitness consequences of the timing of arrival are therefore expected to be high, with penalties both for being too early or too late, resulting in stabilizing selection pressures (Møller *et al.*, 2009a). If the temporal mismatch with phenology of underlying trophic levels increases, we expect this selection pressure to change over time (Møller, 2007b), either favouring birds arriving earlier than average (if the population advances too little) or later than average (if the population advances too much) (Visser, Both & Lambrechts, 2004).

There are, however, few studies explicitly addressing effects of climate change on trophic mismatches for migratory species. Two studies showed increased mismatch due to breeding dates advancing less (Both *et al.*, 2009) or more (Pearce-Higgins *et al.*, 2005) than the main food peak for nestlings. In a recent study, an index of likely mismatch was calculated from temperature trends in wintering and breeding areas, and correlated positively with population declines in the Nearctic (Jones & Cresswell, 2010). Moreover, a study of 117 migratory bird species in Northern Europe (Saino *et al.*, 2010) showed that birds now arrive at higher degree-days than in the past, suggesting that ecological mismatch at time of arrival has increased despite earlier arrival dates. Species for which larger increase of heat accumulation at arrival has been recorded also showed the largest population declines. Fluctuating fitness consequences of arrival date have been demonstrated for a barn swallow population without accurate data on food phenology (Møller, 2007b). For pied flycatchers breeding in the Netherlands, selection for early breeding also became progressively stronger over the years (Both & Visser, 2001), but the response to climate change appears to be insufficient, probably because spring arrival did not advance (Both & Visser, 2001; Both *et al.*, 2005). Unequal temperature change during the course of spring could uncouple the cues for the start of breeding (early spring temperature) and the reproductive consequences determined by a food peak later in the season (depending on temperatures after start of breeding) (Visser *et al.*, 2004), particularly if the time lag is long. Long-distance migrants may be particularly vulnerable, due to a longer sequence of decisions preceding and determining the start of reproduction, beginning with

preparations for migration on the wintering grounds. The value of temperature as a cue does, however, depend on propagation of phenological effects through the food chain (Visser & Both, 2005; Bretagnolle & Gillis, 2010; Thackeray *et al.*, 2010), and direct effects of temperatures might have been overlooked (Visser & Sanz, 2009; Visser, Holleman & Caro, 2009a).

Despite the strong interest in possible trophic mismatches, support for this claim is still limited. Few empirical studies have addressed the claim in detail. A proper assessment of whether responses are adaptive or not requires estimating fitness consequences not just of breeding date, but also of arrival date and spring migration timing. Since there are yet almost no available data of sufficient detail, we have only just begun to understand these issues. Mismatches may also extend to other ecological relationships, including interspecific competition and host-parasite interactions (Claim 10, Section IV.3), and climatic effects on single predator-prey interactions need to be considered in the context of a web of interacting species (Tylianakis *et al.*, 2008; Bretagnolle & Gillis, 2010).

(2) Claim 9: climate change causes population declines in migratory birds

Strong population declines have been reported for long-distance migrants both in Europe and North America over recent decades (Robbins *et al.*, 1989; Berthold *et al.*, 1998; Holmes & Sherry, 2001; Sanderson *et al.*, 2006; Heldbjerg & Fox, 2008), and climate change is likely to be one of multiple causes (Sanderson *et al.*, 2006; Lemoine *et al.*, 2007a; Valiela & Martinetto, 2007; Møller, Rubolini & Lehikoinen, 2008; Newton, 2008; Van Turnhout *et al.*, 2010). Partly, population declines can result from a failure to respond phenologically to changing spring conditions (Møller *et al.*, 2008; Saino *et al.*, 2010; Claim 8, Section IV.1). On the other hand, population declines will affect some phenological metrics (Appendix 1) and genetic (as well as phenotypic) variability for natural selection to act on (Claims 4 and 5, Sections III.1 and III.2).

Climate change is likely to have great impacts on population dynamics of birds through effects mediated by local weather. Although conditions either in breeding areas or wintering areas can dominate (Newton, 1998), population regulation in migratory birds is a complex issue due to the multitude of factors involved at various stages in the annual cycle (Newton, 2004, 2006b). Only for a few species do we know how environmental conditions in different seasons affect both survival and reproduction (Sillert, Holmes & Sherry, 2000). On the wintering grounds, habitat destruction (Sanderson *et al.*, 2006) and habitat deterioration due to climatic fluctuations such as droughts (den Held, 1981; Peach, Baillie & Underhill, 1991; Baillie & Peach, 1992; Szép, 1995; Szép *et al.*, 2006) are probably the main factors causing population declines. Rainfall in Africa is likely the most important single climatic factor determining survival in many Palearctic long-distance migrants, and overwinter condition and survival of American redstarts

is clearly influenced by rainfall and moisture gradients (Marra *et al.*, 1998; Studds & Marra, 2005, 2007). For most species we know far too little about their ranges, movements and ecology in winter quarters to predict clearly how changing wintering conditions will affect population dynamics. Our knowledge from the breeding season is far better, and increased trophic mismatches could result in drastic population declines, as shown for pied flycatchers in the Netherlands (Both *et al.*, 2006). Consequences of mismatch can be expected to depend on the seasonality of the habitat and differential climatic effects (Schaefer *et al.*, 2006; Both *et al.*, 2010), but although some general patterns of population trends according to habitat have been suggested, they are not established (Robbins *et al.*, 1989; Sanderson *et al.*, 2006; Møller *et al.*, 2008; Newton, 2008).

Limiting factors associated with population trends have so far mostly been localized to the breeding season for North American birds and to the winter season for European birds (Newton, 2008), although the combination of effects over the annual cycle remains poorly investigated (Faaborg *et al.*, 2010a,b). There are recent indications of climate change becoming more important than land-use changes (Lemoine *et al.*, 2007a; Møller *et al.*, 2008; Newton, 2008), but such factors are likely to interact (Brook, Sodhi & Bradshaw, 2008; Darling & Côté, 2008; Brotons & Jiguet, 2010). Habitat associations, diet and a number of life-history traits are likely candidates for explaining population trends (Sanderson *et al.*, 2006; Thaxter *et al.*, 2010; Van Turnhout *et al.*, 2010). Furthermore, through seasonal interactions (Claim 7, Section III.4; Appendix S2), survival can affect population dynamics at later stages through delayed density dependence (Ratikainen *et al.*, 2008). Effects at one stage can be compensated at another stage; for instance increased mortality during migration can be compensated by increased survival or reproduction at wintering or breeding grounds due to decreased intraspecific competition (Newton, 2008). Efforts should be made towards better incorporation of seasonal interactions and spatiotemporal complexities in the study of population regulation in migratory birds (Webster & Marra, 2005).

In sum, climate change has been shown to affect population trends, and is likely to be one important cause for current population declines. Very few studies have addressed how multiple factors affect declines, and there is so far insufficient knowledge to predict whether populations of long-distance migrants in general are more vulnerable to climate change than those of short-distance migrants or resident species (see also Claims 2, 4 and 7, Sections II.2, III.1 and III.4).

(3) Claim 10: climate change affects community composition

Considering the observations of range shifts (Claim 3, Section II.3) and concerted population declines (Claim 9, Section IV.2), it has not surprisingly been claimed that climate change can affect community composition (Böhning-Gaese & Lemoine, 2004), and in particular the balance between migratory and resident species in temperate and

northern breeding bird communities (Lemoine & Böhning-Gaese, 2003).

Two main pathways for climate effects have been suggested (Böhning-Gaese & Lemoine, 2004): one through range shifts and regional population trends, and one through interspecific interactions on the breeding grounds. While there is empirical support for the former mechanism, the effect of changing interspecific interactions on the breeding grounds has been little studied. Although effects of climate change on predation risk (Niehaus & Ydenberg, 2006; Bretagnolle & Gillis, 2010) and parasitism levels (Møller, de Lope & Saino, 2004a; Møller, 2009; Saino *et al.*, 2009; Merino & Møller, 2010; Douglas *et al.*, 2010; Møller *et al.*, 2010b) have also been addressed, the main focus has been on competition. Specifically, mild winters are expected to lower the proportion of migrants and increase the survival of residents, resulting in migrants experiencing increased competition for resources upon arrival (Lemoine & Böhning-Gaese, 2003) and hence population decline. This hypothesis has so far not been properly evaluated, but analysis of time series of two or more competing species could be helpful (Newton, 1998; Sætre, Post & Král, 1999; Stenseth *et al.*, 2004; Ahola *et al.*, 2007). Recent reviews suggest the importance of explicit consideration of species interactions for understanding effects of climate change at the community level (Bretagnolle & Gillis, 2010; Brotons & Jiguet, 2010).

Few empirical studies have addressed changes in community composition with respect to climate. Case studies from central Europe (Lemoine *et al.*, 2007a; Reif *et al.*, 2008b) confirm population increases in southern species and decreases in northern species, as expected from range shifts. Interactions with habitat change and land-use change were, however, demonstrated (Lemoine *et al.*, 2007a; Reif *et al.*, 2008a), and such changes can potentially confound climate-change effects (Brotons & Jiguet, 2010). One of these studies (Lemoine *et al.*, 2007a) showed the expected pattern of population declines for long-distance migrants and increases for residents and short-distance migrants, and similar results were found for Dutch breeding birds (Van Turnhout *et al.*, 2010). On the other hand, an analysis of trends from 21 sites across Europe (Lemoine, Schaefer & Böhning-Gaese, 2007b) showed a pattern of decreases in species richness for short-distance migrants and a weak increase in long-distance migrants. Population trends for 68 passerine species breeding in the Czech Republic were mainly explained by food type and general life-history strategy ('r-selected' versus 'K-selected'), and not by different migratory strategies, but populations of resident species fluctuated more than those of short-distance migrants when controlling for phylogeny (Reif *et al.*, 2010). Hence, empirical support for this claim is so far inconclusive and partly rests upon a mixture of changes in quantitative species composition and community-level attributes (Brotons & Jiguet, 2010), as well as species distribution models substituting space for time (La Sorte *et al.*, 2009) by using species-climate relationships at biogeographic scales (Lemoine & Böhning-Gaese, 2003; Schaefer, Jetz &

Böhning-Gaese, 2008). Projections based on climate-change scenarios do not yet indicate clearly whether community change will be mainly due to range shifts (Huntley *et al.*, 2007) or changes in migratory activity (Schaefer *et al.*, 2008); a recent study indicates that the composition of the French breeding bird community is indeed tracking spatiotemporal shifts in temperature, but is lagging behind changes in climate (Devictor *et al.*, 2008). If range shifts are species-specific, changes in community composition seem inevitable, though novel combinations of environmental factors resulting in species assemblages without a present-day analogue may or may not be expected from the magnitude of change projected for the 21st Century (Huntley *et al.*, 2007; Stralberg *et al.*, 2009).

Overall, it is clear that climate change can affect the composition of breeding bird communities through range shifts and population trends. It is poorly known to what extent climate-induced community changes are shaped by interspecific interactions, and effects of parasitism, diseases and predation need to be addressed. We can not currently predict whether migrants and residents will be affected differently, nor how different phenological responses between similar species (Sparks & Tryjanowski, 2007) may transfer to community structure. Changes in community composition outside the breeding season are largely unexplored.

V. CLAIMS, KNOWLEDGE AND SUPPORT

Phenological effects of climate change are well documented for birds, the field has been developing quickly, and birds are particularly well-studied organisms—easily raising expectations for consensus views or consistent 'expert opinions'. We therefore performed a simple assessment of our ability to provide such. As with most typical 'expert assessments' of a field, research questions (here, 'claims') are qualitatively different and inter-related, authors are a non-random and non-interchangeable sample of the research community, and the idea of quantitative analysis was conceived at a late stage. Hence, authors (all respondents being active researchers in the field) simply scored their opinion regarding the amount of research effort so far invested in the claim (a rough measure of the amount of 'knowledge', hence hereafter referred to as the 'knowledge basis') and whether the claim holds in general ('support'), individually and separately for each claim, on a continuous scale from 0 (least) to 10 (most). Agreement (concordance) among observers was quantified using correlation-based measures (see online supporting information, Appendix S4).

There was overall stronger support for claims regarding patterns than for possible mechanisms and consequences (Figs 1, 2A). As expected, support correlated positively with the knowledge basis, both when considering mean scores (Fig. 2A; Spearman's rank correlation on mean scores; $\rho = 0.65$, 95% bootstrap C.I.: -0.09 – 1.00 , $N = 10$) and individual ratings (Kendall's $\tau = 0.47$, 95% bootstrap C.I.:

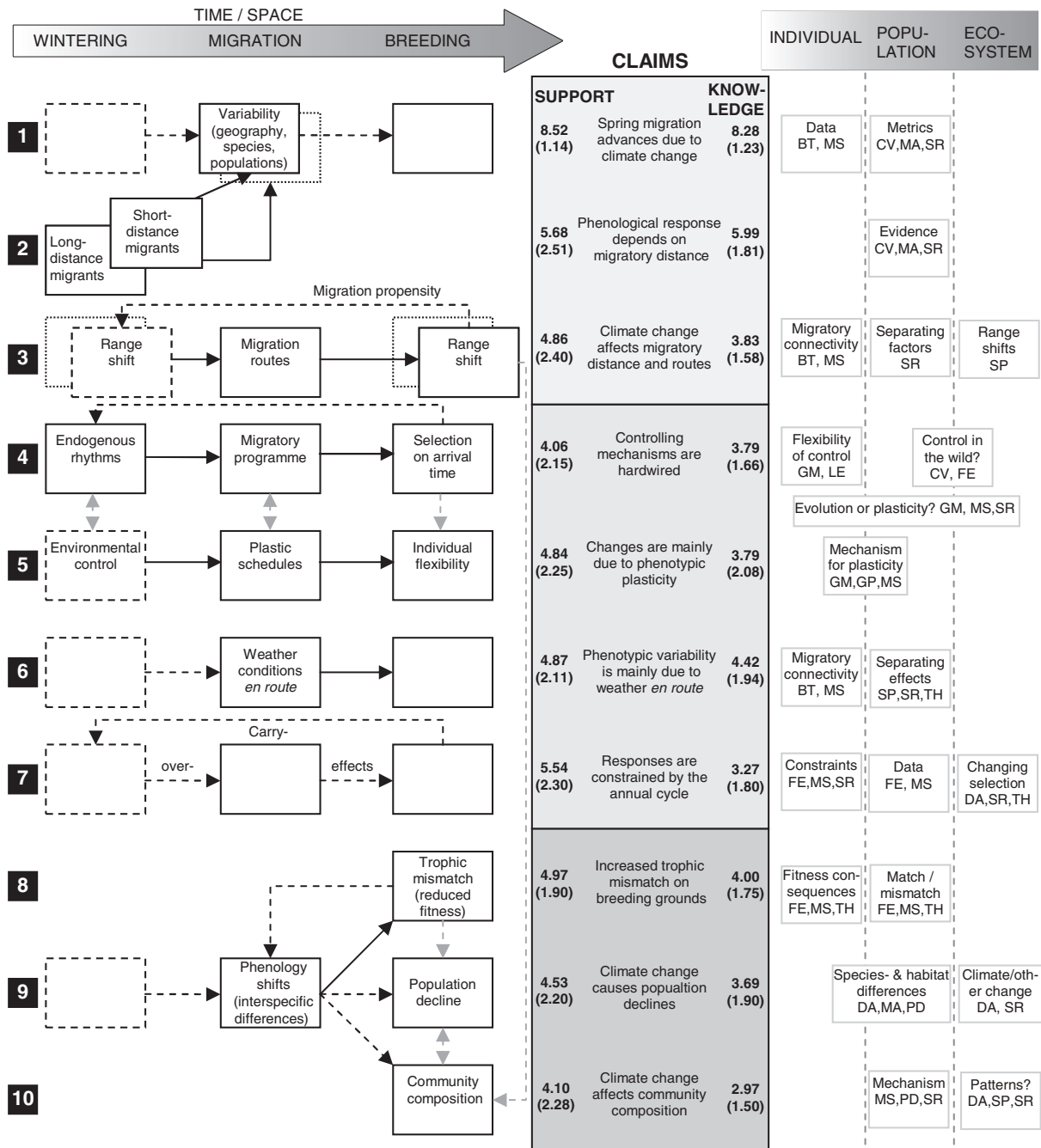


Fig. 1. Claims investigated, overall support, knowledge basis and research challenges. Claims are schematically illustrated to the left; shifts in annual cycle stages are indicated by overlapping boxes, and some links between claims are indicated by arrows. Dashed lines indicate lack of data or uncertain links. In the central column, reported values for support and knowledge basis are the sample mean and standard deviation (in parentheses) of $N = 18$ scores on a scale from 0 to 10 (see main text for details). Some general approaches for meeting research challenges at different levels of biological organization are indicated to the right (BT = biotelemetry; CV = cross-validation; DA = more data; FE = field experiments; GM = molecular genetics/genomics; GP = population genetics; LE = laboratory experiments; MA = meta-analysis; MS = model systems; PD = population dynamics; SP = spatial distribution models; SR = statistical refinement; TH = theory development).

0.38–0.56, $N = 180$). Many claims appear to be rather poorly investigated, and some of these scored higher for support than expected on basis on their knowledge score (Fig. 2A). The level of knowledge about an issue is

often poorly communicated to those outside the research community. This is potentially problematic—the less we know about a matter, the less we tend to agree (Fig. 2B) when asked for our ‘expert opinion’, and the less we agree,

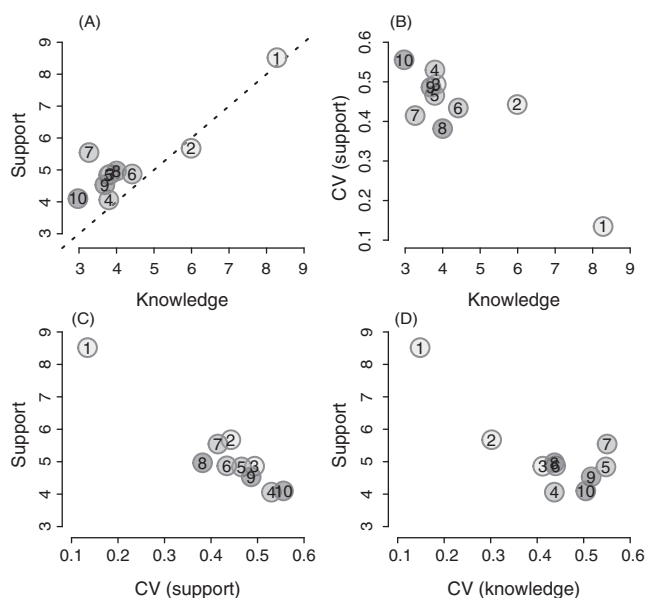


Fig. 2. Selected scatterplots based on summary statistics for $N = 18$ scores for ‘knowledge’ and ‘support’ for the 10 claims, showing how (A) support and (B) between-researcher variability in support is related to the amount of knowledge, as well as how support relates to between-researcher variability in (C) support and (D) knowledge. Plot symbols are annotated with claim numbers, and their shading indicates whether the claim was classified as ‘pattern’, ‘mechanism’ or ‘consequence’ (see also Fig. 1). The dotted line in A shows the ‘45° line’ where mean scores for knowledge and support are equal.

the less overall support we give a claim in a consensus assessment (Fig. 2C,D).

Analysis of concordance in scorings showed overall weak agreement between the 18 researchers regarding support (Lin’s multiple-group correlation $\rho_c = 0.34$, 95% bootstrap C.I.: (0.03–0.49). In less than one of five cases were two researchers more likely to agree than to disagree regarding which of two claims were most supported (Kendall’s $\tau = 0.17$, 95% bootstrap C.I.: 0.14–0.20). Agreement regarding the knowledge basis was moderate (multiple-group $\rho_c = 0.57$, 95% C.I.: 0.04–0.62). However, agreement was largely restricted to observed patterns (claims 1–3); when excluding these, there was no concordance in the scoring of support ($\rho_c = 0.06$) and knowledge basis ($\rho_c = 0.06$). Disagreement between researchers from different disciplines and lines of research is one likely reason for this, and there is a need for unravelling the mechanistic basis of the observed patterns in order to assess potential consequences of climate change. Between-researcher differences in the use of the scoring scale do not appear important; when comparing Kendalls coefficient of concordance to Lin’s correlation (for all claims), we obtained similar results for support ($W = 0.33$, 95% bootstrap C.I.: 0.06–0.64), but somewhat lower concordance for effort ($W = 0.43$, 95% bootstrap C.I.: 0.06–0.79).

VI. CONCLUSIONS

(1) In general, patterns of phenological change in spring migration seem to be well established for birds breeding in Europe and Northern America, with exception of the departure pattern of long-distance migrants from their wintering grounds. Attention could be directed towards mechanisms (including effects of range shifts and migratory routes) by which change can occur, and towards consequences of climate change for individuals, populations and communities. There are, however, interesting biological patterns (e.g. differential migration; Appendix S1) and methodological issues (Appendix 1) that need further attention, and more effort should be put into making the best possible use of the large amounts of data being collected and compiled.

(2) Whether organisms will be able to cope with rapid climate change or not will depend on the rates of adaptive response. Hence, one of the pressing issues is to resolve mechanisms of adaptive change—for instance, to what extent changes in the timing of migration are due to microevolutionary change, changes in migration conditions, or behavioural plasticity in the response to the environment (Sheldon, 2010). This will require collaboration bridging gaps across biological disciplines such as genetics, physiology and ecology (Pertoldi & Bach, 2007; Chown *et al.*, 2010; Visser *et al.*, 2010). The molecular genetic basis regulating the timing of migration remains unknown, but recent progress has been made towards identifying genes potentially regulating the timing of reproduction in partial migrants (see Liedvogel *et al.*, 2009). If good individual-level data are available, quantitative genetics models can be applied, and advanced methods for disentangling variance components such as those sorting under the term ‘animal models’ (Postma, 2010) seem promising. There is also a need for reconciling field studies with experimental approaches, and the huge literature of older experimental work (Berthold, 1984, 1996, 2001) on the behaviour and physiology of migratory birds should be more properly considered in terms of climate-change issues. More knowledge on the evolution of movement patterns and flexibility of the annual cycle in light of long-term climate variability is also needed, and can be achieved along lines of palaeobiological, phylogeographic and phylogenetic comparative studies. It should not be forgotten that regular migration is only one strategy for large-scale movement; seemingly different alternative strategies such as nomadism and irruptive migration may share the same set of basic control mechanisms (Newton, 2006a, 2008; Jonzén *et al.*, 2011).

(3) In order properly to address population-level consequences of climate change, migration needs to be considered in its ecological and evolutionary context, which includes the temporal structure of all important life-cycle stages, as well as frequency- and density-dependent effects within and across trophic levels. Theoretical advancements geared towards climate change (Appendix S3), as well as a fuller and more proper integration

of diverse frameworks (such as match-mismatch, optimal migration behaviour, genetic change, population dynamics and seasonal interactions), can provide helpful steps in this direction. Technological developments (Robinson *et al.*, 2010) will increasingly allow scientists to follow individual birds through the annual cycle, filling in parts of the enormous gap in knowledge on winter distributions, migratory routes and migratory connectivity of individuals and populations. Important mechanistic issues are, however, likely to remain unclear for a while yet, and species distribution models coupled with global change scenarios may be helpful for gaining new insight even if these tools are based on sweeping assumptions and not yet based on a detailed mechanistic understanding (Wiens *et al.*, 2009; LaSorte & Jetz, 2010). More generally, there is a need for integration across lines of research and cross-checking of results derived using different data sources or approaches.

(4) An assessment of concordance among the group of researchers revealed poor agreement regarding the support for most claims, but a somewhat better agreement regarding the knowledge basis, enabling consensus regarding broad patterns and likely causes. With increasing knowledge, the overall support for a claim increased and between-researcher variability in support decreased. Although we as researchers would expect this, the outside world is often considerably more interested in the state of the world than in the state of our knowledge, and often expects us to provide robust consensus views and consistent expert opinions. Our overall low capability to provide a consensus view suggests that even for well-studied organisms, there can be substantial challenges in reaching the required level of knowledge.

(5) Society's call for actions facing climate change has in large been met by biologists in a phenomenological manner, focused on changes in distributions, timing, population levels and ecosystem fluxes. For well-studied organisms such as migratory birds, modern biology (being quantitative and technologically advanced) here meets the wealth of detailed biological knowledge acquired through centuries of natural history research. Integrative biology in this interface is challenging in terms of balancing detail and generality (Appendix 2), but rewarding in terms of providing scientific understanding on how organisms balance environmental changes across the life cycle, and how genetic, physiological, behavioural and ecological aspects of the organism interact in environments subject to changing evolutionary forces. Hence, efforts to assess the knowledge basis and synthesize the literature may be important first steps for more unified and robust attempts at predicting climate-change effects.

VII. ACKNOWLEDGEMENTS

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VIII. APPENDICES

(1) Data on timing of bird migration

The choice of a metric for timing of bird migration has largely been based on availability of data. Dates of first arrival to the breeding area and mean/median migration dates from bird observatories along the migratory route are the most frequently recorded (Lehikoinen *et al.*, 2004). Many sources of noise and bias have been identified in these data (Sparks, Roberts & Crick, 2001; Lehikoinen *et al.*, 2004; Knudsen *et al.*, 2007). The latent phenological distribution of arrival, which can be seen as a population-level life-history trait, is not directly observable and is typically ignored in phenological studies. The shape of this distribution will be modified by migration conditions (with individual phenotypically plastic responses) and mortality during migration, resulting in the realized distribution of arrival.

The most frequently used metric is first arrival date, usually derived from observational data collected by amateurs. This is likely to be only weakly representative of the underlying realized phenological distribution, and the metric is sensitive to variable sampling effort and expected to vary greatly by chance if the distribution is thin-tailed (Sparks *et al.*, 2001; Miller-Rushing, Inouye & Primack, 2008a; Moussus, Juillard & Jiguet, 2010). Sampling effort is usually not controllable and may induce systematic bias due to increasing effort over time, and noise due to higher effort during weekends (Sparks, Huber & Tryjanowski, 2008). First arrival dates are directly negatively related to population size, since the probability of one bird migrating early increases with the number of individuals (Sparks *et al.*, 2001; Miller-Rushing *et al.*, 2008b). The relationship between first arrival dates and population size or observation effort is typically non-linear and depends on distribution shape and location.

Other commonly used metrics of phenology are sample mean or median arrival dates, or other sample quantiles. Typically, these metrics are derived from ringing or observation data at bird observatories. These estimates are usually unbiased with respect to population size, and sampling effort is usually standardized. Some sampling problems do, however, persist, especially for uncommon species and low or high sample quantiles, and as the data usually record birds on stop-over rather than migration, the representation of the data at some localities can be questioned. Bias correction is hard, and the best option may be robust modelling of the arrival distribution (Knudsen *et al.*, 2007), in order to account for varying observation effort and weather effects. The geographic origins and destinations of birds observed at bird observatories are largely unknown, and complications in both interpretation and modelling may arise if the data consist of mixtures of populations or population segments,

particularly if ranges shift (Thomas & Lennon, 1999) or populations show different trends (Miller-Rushing *et al.*, 2008b).

Individual-level data may allow a better approximation of 'true' arrival. Migratory movements can be tracked for the larger species by use of expensive satellite transmitters or geolocators, but sample sizes are usually small. For intensively studied populations, arrival of individuals can be estimated from male singing activity or first capture dates, but species may be unobtrusive or wide-roaming upon arrival, and arrival is probably best estimated for colonial species (Brown & Brown, 2000; Møller, 2008). Such data do, however, allow for better examination of the underlying arrival distribution (Møller, 2008), and factors such as imperfect detectability and mortality during migration can be addressed by using capture-recapture methodology. A potentially important aspect is to define the scale of arrival at breeding grounds. Many birds might gather to feed and/or move around a wider area to recognize different options, before moving to the actual breeding territories (Newton, 2008).

(2) Perspectives: searching for generality in the phenology of migratory birds

Despite the large number of studies on climate-change effects on the phenology of migratory birds, the literature suggests that the search for general patterns has been difficult. Climatic effects and selection pressures vary across space, time and microhabitat, and the complex life cycle spans continental scales. This variability and gaps in knowledge add to the complexity of integrative biology properly considering ecological and evolutionary processes at all organizational levels of organisms, from genes to ecosystems. Hence, researchers are at risk of being either overly pessimistic ("populations respond idiosyncratically") or overly optimistic ("species respond similarly"). However, generality can be found both along the lines of reductionism and holism. Endogenous control mechanisms do indeed regulate the annual cycle, and modelling of births and deaths at various life-history stages sheds light on how climate change could determine population dynamics, thereby also affecting phenology through seasonal interactions. On the other hand, the large number of studies of a large number of species across a large number of environments shows a great and, as yet, underutilized potential for comparative studies, meta-analyses and studies of phylogenetic and life-history constraints. Scaling relationships would be helpful in predicting climate-change consequences—for instance, speed of migration, potential flight range and the time required for moult and breeding scale with body size (Hedenström, 2006, 2008; Rohwer *et al.*, 2009), and in the extension one might address the link between climate and body size, as well as other morphometric traits (Brown & Brown, 1998; Teplitsky *et al.*, 2008; Salewski, Hochachka & Fiedler, 2010; Van Buskirk, Mulvihill & Leberman, 2010), and ask whether small, short-lived species with high demographic turnover are likely to evolve more quickly than longer-lived species.

IX. REFERENCES

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X. SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article.

Appendix S1. Patterns of differential migration.

Appendix S2. Important concepts in the study of bird migration and climate change.

Appendix S3. Theory for studying bird migration and climate change.

Appendix S4. Correlation-based measures of concordance.

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