Assessing the Effects of Climate on Host-Parasite Interactions: A Comparative Study of European Birds and Their Parasites

Anders Pape Møller1*, Santiago Merino2, Juan José Soler3, Anton Antonov4, Elisa P. Badás5, Miguel A. Calero-Torralbo3, Florentino de Lope5, Tapio Eeva6, Jordi Figuerola7, Einar Flensted-Jensen8, Laszlo Z. Garamszegi9, Sonia González-Braojos10, Helga Gwinner10, Sveinn Are Hanssen11,12, Dieter Heylen13, Petteri Ilmonen13, Kurt Klarborg14, Erkki Korpimäki16, Javier Martínez15, Jose Martínez-de la Puente7, Alfonso Marzal15, Erik Mattheysen12, Piotr Matyjasiak16, Mercedes Molina-Morales17, Juan Moreno12, Timothy A. Mousseau18, Jan Tøttrup Nielsen19, Péter László Pap20, Juan Rivero-de Aguilar2, Peter Shurulinkov4, Tore Slagsvold21, Tibor Szép22, Eszter Szöllősi23, Janos Török23, Radovan Vlčav4, Francisco Valera3, Nadia Ziane25

1Laboratoire Ecologie, Systematique et Evolution, Unité Mixte de Recherche 8079 Centre National de la Recherche Scientifique-Université Paris-Sud XI-AgroParisTech, Batiment 362, Université Paris-Sud XI, F-91405 Orsay, France, 2Department of Ecologia Evolutiva, Museo Nacional de Ciencias Naturales, Consejo Superior de Investigaciones Científicas, C/José Gutiérrez Abascal 2, Madrid, Spain, 3Department of Ecologia Funcional y Evolutiva, Estación Experimental de Zonas Áridas-Consejo Superior de Investigaciones Científicas, Ctra. Sacramento s/n, La Cañada de San Urbano, Almería, Spain, 4National Museum of Natural History - Sofia, Bulgarian Academy of Science, Sofia, Bulgaria, 5Department of de Anatomia, Biologia Celular y Zoología, Universidad de Extremadura, Badajoz, Spain, 6Section of Ecology, Department of Biology, FIN-20014 University of Turku, Finland, 7Department de Ecologia de Humedales, Estación Biológica Doñana (Estacion Biologica de Doñana-Consejo Superior de Investigaciones Científicos), Sevilla, Spain, 8Cypraves 1, DK-9700 Bønderup, Denmark, 9Department of Ecología Evolutiva, Estación Biológica Doñana (Estacion Biologica de Doñana-Consejo Superior de Investigaciones Científicos), Sevilla, Spain, 10Max Planck Institute for Ornithology, Eberhard-Günther-Straße, Haus Nr. 11, Seewiesen, Germany, 11Norwegian Institute for Nature Research – Norwegian Institute for Nature Research, Polar Environmental Centre, Tromsø, Norway, 12Evolutionary Ecology Group, Department of Biology, University of Antwerp, Antwerpen, Belgium, 13Division of Genetics and Physiology, University of Turku, Finland, 14Skovvej, DK-9490 Pandrup, Denmark, 15Department of Parasitología, Universidad de Alcalá, Alcalá de Henares, Spain, 16Faculty of Biology and Environmental Sciences, Cardinal Stefan Wyszynski University in Warsaw, Woycickiego 1/3, Warsaw, Poland, 17Department of Zoología, Facultad de Ciencias, Universidad de Granada, Granada, Spain, 18Department of Biological Sciences, University of South Carolina, Columbia, South Carolina, United States of America, 19Espedal 4, Tolne, DK-9870 Sindal, Denmark, 20Evolutionary Ecology Group, Hungarian Department of Biology and Ecology, Babeș-Bolyai University, Cluj Napoca, Romania, 21Department of Biology, Centre for Ecological and Evolutionary Synthesis, University of Oslo, Oslo, Norway, 22Institute of Environmental Sciences, College of Nyíregyháza, H-4400 Nyíregyháza, Sóstói út 31/b, Hungary, 23Behavioural Ecology Group, Department of Systematic Zoology and Ecology, Eötvös Loránd University, Pázmány Péter stny 1/C, Budapest, Hungary, 24Institute of Zoology, Slovak Academy of Sciences, Dúbravská cesta 9, Bratislava, Slovakia, 25Department of Biology, Faculty of Science, Badji Mokhtar University, Boite Postal 12, Annaba, Algeria

Abstract

Background: Climate change potentially has important effects on distribution, abundance, transmission and virulence of parasites in wild populations of animals.

Methodology/Principal Finding: Here we analyzed paired information on 89 parasite populations for 24 species of bird hosts some years ago and again in 2010 with an average interval of 10 years. The parasite taxa included protozoa, feather parasites, diptera, ticks, mites and fleas. We investigated whether change in abundance and prevalence of parasites was related to change in body condition, reproduction and population size of hosts. We conducted analyses based on the entire dataset, but also on a restricted dataset with intervals between study years being 5–15 years. Parasite abundance increased over time when restricting the analyses to datasets with an interval of 5–15 years, with no significant effect of changes in temperature at the time of breeding among study sites. Changes in host body condition and clutch size were related to changes in temperature between first and second study year. In addition, changes in clutch size, brood size and body condition of hosts were correlated with change in abundance of parasites. Finally, changes in population size of hosts were not significantly related to changes in abundance of parasites or their prevalence.

Conclusions/Significance: Climate change is associated with a general increase in parasite abundance. Variation in laying date depended on locality and was associated with latitude while body condition of hosts was associated with a change in temperature. Because clutch size, brood size and body condition were associated with change in parasitism, these results suggest that parasites, perhaps mediated through the indirect effects of temperature, may affect fecundity and condition of their hosts. The conclusions were particularly in accordance with predictions when the restricted dataset with intervals of 5–15 years was used, suggesting that short intervals may bias findings.

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Introduction
The commonest interactions between species occur between parasites and their hosts [1,2]. These interactions depend on parasite prevalence and abundance, rate and mode of transmission, effect of parasites on fecundity and mortality of hosts and the level of anti-parasite defence by hosts [3]. Each of these different steps in the interaction can potentially be affected by environmental conditions including climatic conditions [4]. Several studies have indicated that parasite diversity is higher at low latitudes [5–7], and negative impacts on avian [8] and human hosts [5,9,10] are higher at low latitudes. This is partly linked to latitudinal clines in climate, but also clines in host immunity and host population density [11–13], and consequently climate may have direct effects on interactions between hosts and parasites by affecting the abundance of parasites, or by indirectly affecting phenology of parasites and hosts [14].

Many studies have pondered the effects of climate change on parasites, host-parasite interactions and ultimately veterinary and public health [15–22]. Increasing temperature is the main factor linked to climate change effects on living organisms although changing precipitation and wind may also play a role. Many parasites are advancing their date of emergence [4,23–25], and some that were previously active only in the summer are now active year-round [26,27]. A longer and earlier period of reproduction by parasites may increase the number of parasite generations per year, as a longer breeding season of the host provides parasites with a selective advantage [28,29]. However, a longer reproductive season may also allow hosts to better defend themselves against parasites and hence achieve higher reproductive success because timing of reproduction is less constrained by migration [30,31]. Most models of the effect of climate change on host-parasite interactions predict that the distribution of parasites will move northwards, while the overall geographical range of the parasite will not change, nor will total population size [4]. Host populations may also expand as a consequence of climatic change, although that will partly depend on their immunity and hence their ability to cope with novel parasite assemblages [32].

Empirical assessments of effects of climate change on host-parasite interactions are scarce, and, therefore, there is high uncertainty in predicting the consequences. Moller [33] analyzed prevalence and intensity of several species of parasites exploiting the barn swallow Hirundo rustica, showing that parasites that do not complete their life cycles on hosts were more strongly affected by climate change than those that do. In addition, a directly transmitted mite seemed to have reduced its virulence in response to climate change, while cell-mediated immunity of swallows decreased over time. A recent meta-analysis concluded that the incidence of blood parasites and avian malaria increased during recent decades [34]. Likewise, changes in prevalence of a brood parasitic cuckoo were related to change in temperature [35–36]. In addition, several studies have shown changes in phenology of both hosts and parasites [34]. Finally, some studies of hosts and parasites have demonstrated changes in parasite-mediated mortality and resistance of hosts [34,36,37]. For example, host races of the cuckoo Cuculus canorus rely on migratory hosts that now arrive well before the arrival of the parasitic cuckoo. Therefore such host races have declined dramatically in abundance in recent years [36]. While there is evidence of changes in host-parasite interactions related to climate change, there is considerable unexplained heterogeneity among parasite taxa.

In an attempt to fill this gap, we established a European network of scientists interested in host-parasite interactions exploiting existing historical data on abundance and impact of parasites on their bird hosts. By returning to exactly the same host populations in 2010, and recording parasitological, host demographic and host density data as already done in exactly the same way in an earlier study year, we developed a paired design to test for climate-driven change in host-parasite interactions. We explicitly tested for an effect of interval between first and second study year on temperature and parasite and host variables while statistically controlling for the time elapsed between studies. Such within-population comparisons are known to be particularly powerful because they allow separation of within from between population variance [38].

Methods
Ethics Statement
This study was carried out in strict accordance with current laws of all the countries where the study was performed and following the recommendations of the ‘Guidelines to the Use of Wild Birds in Research’ [41]. All efforts were
made to ameliorate suffering of animals and minimize handling time. No birds were injured or killed during the study.

Blood sampling was carried out at locations in Belgium, Bulgaria, Denmark, Finland, Hungary, Norway, and Spain. The sampling consisted of collection of a tiny blood sample from the brachial vein. This procedure is a standard procedure performed by numerous bird banders throughout the world for the study of avian malaria. No birds showed signs of negative consequences of blood sampling. We have recorded as high survival rates from sampled birds as from other birds. Collection of blood was specifically authorized in each location under the permits issued for this work.

**Study Populations**

In January 2010 we requested 58 European scientists, who had published on bird-parasite interactions to participate in the project. Participants were mainly selected based on a recent review of the effects of bird parasites on the fitness of their hosts [8]. Three requests were made and eventually 37 scientists participated in the study. Some scientists did not participate because they were not working on parasites any more, or they were committed to other projects during 2010. The geographical distribution of the study sites is shown in Fig. 1. We deliberately collected all recent samples during a single breeding season (2010) to reduce potential bias related to variation in variables among years. Although this approach may reduce variation in estimated differences due to the second year being the same for all populations, geographic variation in climatic conditions during 2010 was sufficiently large among study populations to ensure conclusions that are independent of the particular climatic conditions of 2010. Geographic variation in temperature in 2010 did not differ from that of the first study year (Levene's test, \(F = 0.45, \text{ d.f.} = 1, 88, P = 0.50\)), although the first study year differed among populations. We asked all participants to use exactly the same methods in 2010 as during the first year of their study, but also that the same person conduct or at least supervise the study, thus ensuring that all studies were consistent in methodology over time to avoid inter-observer variability.

**Parasites**

We distinguished between four functional parasite groups based on their taxonomic and transmission status: protozoans including blood parasites, feather parasites, dipteran parasites and non-dipteran parasites. Protozoans are generally vector-transmitted. Feather parasites such as chewing lice and feather mites live in the plumage of birds. Dipteran parasites include louseflies and blowflies. Non-dipteran parasites included blood-sucking fleas, ticks and mites. When more than one taxon of parasite within one of these groups were investigated in a target host species and population, we estimated mean values, which were used for subsequent analyses. Variance explained by the four functional groups of parasites was significantly larger than the residual variance (that was explained by different kinds of parasites within the four functional groups) for parasite abundance (\(F = 4.31, \text{ d.f.} = 1, 88, P = 0.04\)).
d.f. = 3.44, \( P = 0.01 \) and prevalence \( (F = 4.14, \text{d.f.} = 3.82, \ P = 0.009) \), which further supports the use of mean values.

We requested information on prevalence (the fraction of adults or nests harboring a given parasite) and the mean abundance of parasites exactly as done in the first year of the study and again in 2010. We requested that participants record information on demographic traits and population density of hosts as described below.

Information on the methods used to quantify parasites can be found in the references in Moller et al. [8] and Merino et al. [43]. In brief, we quantified prevalence and intensity of infection (mean abundance of parasites per host/host nest). Most studies either used two independent methods to quantify parasites or quantified parasites repeatedly for the same individuals/nests to allow estimation of repeatability. All studies used statistically significant and highly repeatable estimators of parasite prevalence and abundance.

### Temperature Trends

We used the E-OBS gridded dataset (version 5.0) maintained by the European Climate Assessment and Dataset (ECAandD) (http://eca.knmi.nl/) to estimate temperatures at the study sites [44]. We calculated mean monthly temperature for each 0.25°x0.25° degree squares, covering the study sites from the daily mean temperature of the E-OBS gridded dataset. Change in temperature during the month best covering the laying, incubation and nestling period of each bird host species was estimated as the temperature in 2010 minus the temperature in the first study year divided by the interval between the two years. This change in temperature over time \( (\text{°C/year}) \) is referred to as change in temperature throughout the remainder of this paper.

### Life History and Population Density of Hosts

We requested that all participants record for each host the date of laying of the first clutch, clutch size, reproductive success (no. fledglings) and body condition (estimated as body mass at the start of the breeding season, if possible). We also requested a local estimate of population density of hosts (such as the proportion of nest boxes occupied for nest box studies, the total number of occupied nest boxes, the number of individual birds captured, colony size, or in a few cases population density for open nesting species). The entire dataset is reported in File S1 in Tables S1–S2, while Table S3 in file S1 presents correlations between prevalence and response variables in the supplementary material (ESM).

### Statistical Analyses

All variables were \( \log_{10} \)-transformed to achieve approximately normal distributions. The effects of time between sampling of host populations and parasite intensity and prevalence were explored by Repeated Measures ANOVAs with laying date, clutch size, brood size, body condition of adults, population density, parasite loads and parasite prevalence estimated for the two study years as within subject factors and change in parasite abundance and prevalence as between subject factors. Parasite functional group, latitude and temperature change \( (\text{°C/years}) \) were also included in the models as additional between factors to statistically control for the effect of these factors on changes in host population characteristics. A significant interaction between repeated measure and between factors would be consistent with an effect of the latter on change in host population characteristics.

Statistical analyses assume that each observation provides equally precise information about the deterministic part of the process variance [38]. Sample sizes differed among species due to differences in abundance of hosts, and such variation may affect the precision of estimates and hence the outcome of the statistical analyses [45,46]. Therefore, we weighted all analyses by sample size to ensure that individual observations contributed relative to their precision.

The findings might depend on duration of the interval between first and second study year. Therefore, we conducted a second series of analyses based on datasets with an interval of 5–15 years, and these results are reported in Tables S4–6 in File S1. All analyses were performed with Statistica 10 [47]. Values reported are means (SE).

### Results

#### Summary Statistics

The mean change in temperature per year across the 89 data sets was \( +0.085 \text{°C} \) (SE = 0.029), with a range from \(-0.65 \text{°C}\) to \(+1.07 \text{°C}\), differing significantly from zero which would be expected if temperature on average was the same in the first and the last year of study \( (t\text{-test, } t = 2.90, P = 0.0047) \). A similar increase in temperature over time was found for studies based on intervals of 5–15 years \( (+0.076 \text{°C} \text{SE} = 0.020) \), with a range from \(-0.55 \text{°C}\) to \(+0.50 \text{°C}\). Temperature in the first year was lower than in the second year \( (t = 2.52, P = 0.013) \), and that was even more so for studies with an interval of 5–15 years \( (t = 4.30, P = 0.0001) \). The number of years between the first study year and 2010 was on average 10 years \( (\text{SE} = 0.63) \), range 1 to 39 years with 78% within the interval 5–15 years. We also detected a large range in change in temperature across studies from \(-2.94\text{°C}\) to +4.04\text{°C}, on average +0.40\text{°C}, for the mean period of study. Because change in temperature between the first year of study and 2010 decreased slightly with the number of years of study \( (F = 4.02, \text{d.f.} = 1.87, r^2 = 0.04, P = 0.048, \text{estimate} \text{SE} = -0.053 \text{(0.027)}) \), implying that temperatures decreased more in studies of long duration, we used temperature change among study years divided by number of years elapsed between the first year and 2010 in the subsequent analyses. There were no additional effects of latitude or longitude on change in temperature \( (\text{latitude: } F = 0.05, \text{d.f.} = 1.85, P = 0.83; \text{longitude: } F = 0.39, \text{d.f.} = 1.85, P = 0.54) \), suggesting that these variables were not confounding effects on change in temperature.

Total sample size was 9935 hosts (or host nests depending on study) in the first year and 7956 in 2010, or in total 17,891 hosts/nests. A total of 37.2% of the 89 parasites studied were ectoparasites, and 76.4% of the 89 parasites lived permanently on the host, with the remaining species living apart from the host at ambient temperature at least part of the annual cycle. Prevalence and abundance of parasites in the first study year was independent of the duration of the study period \( (\text{GLM: prevalence: log-likelihood } \chi^2 = 2.12, \text{d.f.} = 1, P = 0.14; \text{abundance: log-likelihood } \chi^2 = 0.42, \text{d.f.} = 1, P = 0.51) \).

### Temporal Change in Host Populations

Host populations on average started to reproduce later during 2010 than during the first study year \( (\text{Fig. 2}) \). Moreover, temporal change in laying date differed significantly among localities.
(Table 1), which may be related to differential effects of climatic change at different latitudes (Table 1, Fig. 3). Change in clutch size decreased with increasing change in temperature (Table 1; Fig. 4A). Change in brood size differed among localities (Table 1), but not when considering studies with intervals between 5–15 years (Table S4 in File S1). Change in body condition tended to decrease with increasing temperature (Table 1; Fig. 4B). Host populations were less abundant in 2010 and this did not depend on host identity when studies with intervals between 5–15 years were considered (Table 1, Table S4 in File S1).

**Table 1.** Repeated measures ANOVAs with laying date, clutch size, brood size, body condition and population density of hosts in two separate study years as within subjects factors and locality identity, host identity, latitude, interval in years and temperature change (°C/year) as between subjects factors.

| Repeated measure | Locality | Host identity | Latitude | Interval in years | Temperature change (°C/year) |
|------------------|----------|---------------|----------|-------------------|-------------------------------|
| Laying date      | 8.69     | 1.26          | 0.007    | 7.38 19.7         | 0.006 1.17 15.11 0.406 5.78 1.25 0.024 0.02 1.25 0.900 2.49 1.25 0.127 |
| Clutch size      | 0.86     | 1.29          | 0.363    | 0.83 20.9         | 0.654 0.80 17.12 0.673 0.27 1.28 0.606 0.27 1.28 0.606 6.84 1.28 0.014 |
| Brood size       | 0.18     | 1.28          | 0.673    | 3.53 19.9         | 0.029 0.42 16.12 0.948 3.02 1.27 0.093 0.00 1.27 0.973 0.00 1.27 0.983 |
| Body condition   | 0.57     | 1.22          | 0.458    | 1.00 12.10        | 0.505 1.22 14.8 0.400 0.14 1.21 0.710 0.18 1.21 0.675 5.33 1.21 0.031 |
| Population density | 22.88   | 1.38          | 0.0001   | 1.54 22.16        | 0.190 5.51 22.16 0.001 0.41 1.37 0.525 0.01 1.37 0.926 0.39 1.37 0.537 |

Each effect was estimated in separate models. P-values smaller than 0.05 are shown in bold.

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Figure 2. Temporal change in laying date of hosts, body condition of hosts, population size of hosts, brood size of hosts and clutch size of hosts between the first study year and 2010. Box plots show means, standard errors and 95% confidence intervals.
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Temporal Change in Parasite Populations

We detected 18 gains and 8 losses of parasites among studies, and for studies with 5–15 years intervals there were 10 gains and 10 losses. Abundance of parasites was larger in 2010 than in the first study year. However, none of the variables explained change in abundance of parasites (Table 2; Fig. 5A). Variables that explained change in parasite prevalence depended on interval between study years. Locality was associated with parasite prevalence when considering studies with 5–15 years intervals (Table S5 in File S1), while parasite identity did so when all studies...
Table 2. Repeated measures ANOVAs with parasite abundance and prevalence in two separate study years as within subjects factors and locality identity, parasite identity, latitude, interval in years and temperature change (°C/year) as between subjects factors.

| Repeated measure          | Locality | Parasite identity | Latitude | Interval in years | Temperature change (°C/year) |
|---------------------------|----------|-------------------|----------|-------------------|-----------------------------|
|                           | F        | d.f.              | P        | F        | d.f.              | P                  |
| Parasite load             |          |                   |          |          |                   |       |
|                           | 1.15     | 1,29              | 0.293    | 0.81     | 17,12             | 0.660              |
|                           |          |                   |          |          |                   |       |
| Parasite prevalence       |          |                   |          |          |                   |       |
|                           | 0.19     | 1,43              | 0.668    | 0.89     | 20,23             | 0.597              |
|                           |          |                   |          |          |                   |       |

Each effect was estimated in separates models. P-values smaller than 0.05 are shown in bold.

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Figure 5. Change in parasite abundance and parasite prevalence over time for different categories of parasites. (A) Change in parasite abundance and (B) change in parasite prevalence between study years for different parasite taxa. Box plots show means, standard errors and 95% confidence intervals. The right Y-axis in (A) is the number of non-dipteran parasites.

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were considered (Table 2). Non-dipteran parasites were the only group that tended to decrease in prevalence between sampling years although this category did not differ significantly from tendencies for other categories (Table 2; Fig. 5B).

**Discussion**

We posed the question whether a general increase in temperature had a significant effect on abundance of parasites and hosts, and whether this led to altered effects of parasites on host fitness. We recorded 18 gains and 8 losses of parasites although a more balanced frequency of 10 gains and 10 losses were found for the restricted dataset. Hosts generally started to reproduce later in recent years, and we could link this change in clutch size and change in body condition to change in temperature, although obviously other unknown factors could have changed as well. Change in parasite abundance and prevalence was not significantly related to change in temperature. Change in parasite abundance, but not change in parasite prevalence, explained a significant proportion of variance in host life-history variables. Specifically, increasing parasite abundance was associated with clutch size, brood size and body condition, as was the increase in temperature. This suggests that changes in parasite abundance occurred through indirect effects such as effects of temperature on hosts. This raises the possibility that other aspects of climate change such as precipitation or wind could affect insect vectors and larval nematodes. Finally, population size of hosts was not significantly correlated with abundance and prevalence of parasites or change in temperature.

Change in abundance of parasites was linked to change in temperature during the breeding season of hosts, which in general is also the breeding season of the parasites. This general finding is in accordance with several studies that have documented changes in prevalence and/or abundance of parasites over time [34], or in response to change in climate [33,35,36]. We caution that change in parasite prevalence or abundance over time may be linked to many factors other than climate change. Long-term studies of the same population of hosts and parasites during decades in areas where temperature has increased dramatically are clearly more powerful, and hence more likely to find predicted changes [33] than studies based on two years.

Increasing temperatures advance the emergence of resting stages of parasites [4,23–25,33,40] or their vectors [33,34,49]. This can affect the fitness costs of parasitism for hosts because parasites inflict greater fitness costs when weather is poor [50–52], and earlier start of reproduction by hosts is likely to increase the risk of adverse weather. Likewise higher temperatures can increase reproductive rates of parasites and thus the rate of evolution by parasites relative to that of hosts [4,23,24,48], even when hosts also increase their reproductive rate [33]. Finally, climate may affect defenses of hosts and thereby increase the level of virulence of parasites [4,13,37,53,54]. These scenarios predict changes in parasite populations associated with climate change as shown by our results when restricting the analyses to studies with an interval of 5–15 years. In addition, we found support for the prediction that temperature explained change in clutch size and body condition of hosts, and both these fitness components were associated with parasite abundance. This provides evidence of an indirect effect of parasites on fitness components of hosts associated with an effect of temperature on the same fitness components.

The support for our predictions when restricting the analyses to studies with intervals of 5–15 years suggests that a certain amount of time must have passed to result in changes. We found associations between change in temperature and change in clutch size and change in body condition, respectively, although similar effects were absent for laying date and brood size. Factors such as those related to host characteristics, and whether parasites spend part of the year away from hosts, may be important.

We found an increase in abundance of parasites in 2010 compared to the first year when restricting the analyses to datasets with an interval of 5–15 years. There was no confounding effect of population density of hosts because host population density on average decreased during the course of the study, thereby reducing parasite transmission rates. Most of the parasite species were endoparasites living permanently within their hosts, hence only indirectly being affected by changing temperatures. We can only speculate that the detected change in the abundance of parasites for populations may be related to the indirect effects of temperature on host population density (Results) and/or an increase in immunity and anti-parasite responses of hosts [14,55,56].

The parasite species have all at least sometimes been shown to negatively impact fitness components of their hosts [8]. We detected that species, which experienced an increase in the abundance of parasites during the study period, advanced their laying date compared to species in which parasite abundance decreased. However, this effect disappeared when considering the restricted dataset that only included studies with an interval of 5–15 years. With this restricted dataset we found evidence of change in body condition, clutch size and brood size of hosts being negatively linked to change in temperature and parasite abundance, while changes in body condition and clutch size were also associated with change in temperature. These associations were not detected when considering parasite prevalence. While parasite abundance reflects the impact of parasites on hosts at the level of individual hosts, prevalence constitutes a measure of parasite populations at the level of host populations. Our results are therefore in accordance with the assumption that parasitism negatively influences reproductive success of their hosts, and with
Table 3. Repeated measures ANOVAs and within subjects effects (parasite abundance/prevalence in the first and the second year) with laying date, clutch size, brood size, body condition and population density of hosts in two separate study years as within subjects factors and interactions between the repeated measure and latitude, temperature change, variation in parasitism (i.e. change in abundance and prevalence of parasites) and identity of parasite group as between subjects factors. P-values smaller than 0.05 are shown in bold.

|                 | Parasite abundance | Parasite prevalence |
|-----------------|--------------------|---------------------|
|                 | F      | d.f. | P   | F      | d.f. | P   | F      | d.f. | P   | F      | d.f. | P   |
|                 | Parasite abundance | Parasite prevalence |
| Laying date     | 18.37  | 1,19 | 0.001 | 16.32  | 1,19 | 0.001 | 2.95  | 1,19 | 0.102 | 10.73  | 1,19 | 0.004 | 7.24  | 3,19 | 0.002 |
| Clutch size     | 0.67   | 1,20 | 0.422 | 0.07   | 1,20 | 0.456 | 1.27  | 1,20 | 0.274 | 3.77   | 1,20 | 0.066 | 1.57  | 3,20 | 0.227 |
| Brood size      | 0.13   | 1,19 | 0.723 | 0.04   | 1,19 | 0.849 | 0.09  | 1,19 | 0.118 | 2.68   | 1,19 | 0.118 | 0.91  | 3,19 | 0.456 |
| Body condition  | 5.65   | 1,13 | 0.033 | 7.18   | 1,13 | 0.019 | 10.74 | 1,13 | 0.006 | 6.86   | 1,13 | 0.021 | 3.67  | 3,13 | 0.041 |
| Population density | 9.38  | 1,20 | 0.006 | 12.36  | 1,20 | 0.002 | 0.01  | 1,20 | 0.924 | 0.93   | 1,20 | 0.347 | 1.94  | 3,20 | 0.156 |

Interactions are indicated by *.

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the hypothesis that the effect of climate change on host reproduction is at least partially mediated by changes in parasitism due to climate change.

While changes in the distribution and abundance of parasites may impact their hosts, whether this has any population consequences for the host depends on density-dependent mortality and fecundity [57–59]. Populations which have declined the most are those that have not responded to climate change [9,35,60,61]. We found no evidence for an effect of climate change on population size of birds as mediated by parasites. Change in population density of hosts depended on latitude, and it differed among parasite groups. These findings suggest that climate change has not affected population size of hosts directly through effects of change in parasitism or change in temperature.

In conclusion, while climate change may affect hosts by affecting phenology, reproduction and host body condition, an increase in temperature was associated with a general increase in parasite abundance. Clutch size, brood size and body condition of hosts were associated with a change in temperature and with a change in parasitism, respectively, suggesting that parasites may affect fecundity and condition of their hosts.

Supporting Information

**Table S1** Data files and supplementary analyses of determinants of parasitism and host life history. Table S2. Summary information on parasitism and study populations. Table S3. Correlations between parasitism and life history traits. Table S4. Analyses of host populations based on data with an interval of sampling of 5–15 years. Table S5. Analyses of determinants of parasitism based on data with an interval of sampling of 5–15 years. Table S6. Analyses of parasitism in relation to life history based on data with an interval of sampling of 5–15 years.

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**Author Contributions**

Conceived and designed the experiments: APM SM. Performed the experiments: APM SM, JJS AA, EPB, MACT, FdL, TE, JF, EFJ, LG, SGB, HG. Analyzed the data: APM, JJS. Wrote the paper: APM SM, JJS.

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