Migrating Curlews on Schedule: Departure and Arrival Patterns of a Long-Distance Migrant Depend on Time Rather than on Wind Conditions

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Research

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Abstract

Background: Choosing the appropriate time to depart for spring migration is crucial to achieving a successful subsequent breeding season among migratory bird species. We expected Eurasian Curlews (*Numenius arquata*) to start their migration during favourable weather conditions and to adjust their flight heights to prevailing wind conditions.

Methods: We equipped 23 curlews with Global Positioning System data loggers to record the spatio-temporal patterns of their departure from and arrival at their wintering site in the Wadden Sea, as well as the first part of their migration. We obtained data for 42 curlew migrations over a period of 6 years. Departure and arrival dates were related to 73 meteorological and bird-related predictors using the least absolute shrinkage and selection operator (LASSO) to identify drivers of departure and arrival decisions.

Results: Curlews migrated almost exclusively to the western part of Russia for breeding. They left the Wadden Sea mainly during the evening hours from mid- to late April and returned between the end of June and mid-July. There was no difference in departure times between the sexes, but males tended to return from their breeding sites later than females. Flight speeds of the birds increased significantly with increasing tailwind component, suggesting that they timed their migration according to favourable wind conditions. However, curlews left the Wadden Sea during various wind and weather conditions, with significant numbers leaving during headwind conditions, in contrast to the apparent wind-driven start of migration. Curlews migrated at very low altitudes during tailwind conditions, but flew significantly higher during headwind conditions, at altitudes of up to several kilometres. Departure dates varied by <4 days in individual curlews that were tagged over consecutive years.

Conclusions: Our results suggest that the start of migration in a long-distance migrant mainly depends on the date and is independent of weather conditions. Given the high repeatability of the departure day among subsequent years, this clearly suggests the existence of an internal clock prompting the start of migration. Further insights into the timing of migration in immatures and closely related birds might help us to understand the genetic mechanisms triggering temporal migration patterns.

Background

Migration is an essential part of the life cycle of a wide range of species, with potentially important consequences for their fitness [1–5]. Birds show the most extensive and far-ranging migrations [6–8]. Careful timing of migration is essential to maximize the chance of survival during migration (e.g. to avoid severe weather) [9–12, 5] and to coincide with beneficial environmental resources in the destination areas, as a prerequisite to ensuring fitness [13–14].

We studied the departure and arrival patterns of a long-distance migrating shorebird, the Eurasian Curlew (*Numenius arquata*), at one of its most important wintering sites on the East Atlantic Flyway, the Wadden Sea. Despite strong population decreases in the flyway population as a whole [15], the numbers of curlews wintering in the Wadden Sea have remained stable for several decades [16]. The Wadden Sea
population holds around 200,000-260,000 individuals, accounting for around 40% of the total flyway population [16–17]. However, information on the migration patterns of curlews wintering in the Wadden Sea is scarce (but see [18] for a preliminary study). A previous study from south-west England investigated the arrival and departure patterns of curlews based on a dataset of re-sightings of colour-marked individuals [19]. In the current study, we equipped curlews with Global Positioning System (GPS) data loggers that recorded the times of arrival and departure of each individual bird in the Wadden Sea. This allowed individual departure and arrival patterns to be precisely related to meteorological data, and allowed the repeatability of temporal patterns across subsequent years to be assessed. We proposed the following five hypotheses. (1) Tailwinds increase flight and migration speeds [6; 20–21], and we therefore expected a positive linear relationship between flight speed and tailwind components (TWC). (2) Individuals of a range of bird species have been shown to time their departure according to favourable wind and weather conditions (e.g. a tailwind and no or little precipitation; [12, 22–24]. In line with our first hypothesis, we therefore expected curlews to time their departure from and arrival at their wintering grounds according to favourable wind and weather conditions, especially in relation to tailwind conditions and lack of precipitation. (3) We expected flight heights (as recorded by GPS tags) to increase during non-tailwind conditions to allow the birds to reach air layers with improved wind conditions [25–26]. (4) We hypothesized that birds breeding further from their wintering site would depart earlier because they needed more time for travelling. In this context, we expected males to arrive at the wintering sites later than female curlews, because, as for other shorebird species, females are known to desert their chicks earlier than males [27]. (5) Finally, we predicted significant effects of departure date and tailwind component on the distance to and duration of the first stop-over event. Curlews departing later were expected to stop-over for only a short period of time and then continue quickly to their breeding grounds [23]. Previous studies indicated that headwind conditions could significantly shorten the distance to the first migration stop-over and increase the stop-over duration to allow birds to refuel before continuing [e.g. 28].

In the current study, we analysed data for tagged curlews from the Wadden Sea area over multiple years and extended the preliminary GPS-tagging study presented in [18].

**Methods**

**Study area**

Curlews were caught along the eastern Wadden Sea coast of the German federal states of Schleswig-Holstein and north-eastern Lower Saxony between 54°36´N and 53°42´N, and between 7°54´E and 8°54´E (Fig. 1). Meteorological parameters were recorded by the automatic recording station of the Research and Technology Centre, located in Büsum, federal state of Schleswig-Holstein (54°7´55˝N; 8°52´37˝E; yellow circle in Fig. 1). Flight speeds and altitudes of GPS-tagged individuals were recorded within an area stretching from the Wadden Sea coast to the Baltic Sea and from south Denmark to the northern part of the federal State of Lower Saxony (red box in Fig. 1).
Catching of curlews and deployment of GPS tags

A total of 26 adult curlews were caught at their high-tide roosts using mist nets, between 2014 and 2020. Three of the GPS devices malfunctioned or the birds were predated before departure from their wintering grounds, and data for 23 adult wintering curlews (11 females and 12 males) were therefore available for this study. Some curlews migrated multiple times before the device stopped working, and we were therefore able to record the departures and arrivals of the same individuals in up to 5 subsequent years. We recorded a total of 42 departures from and 33 arrivals at the wintering sites. All individuals were equipped with solar-powered GPS-GSM data loggers attached by breast harnesses [18, 29]. The devices recorded time, date, geographical position, and flight speed at pre-programmed intervals of 1–15 min. Twelve individuals were equipped with “Skua” data loggers (Ecotone, Poland) weighing 17 g, and the other 11 individuals were equipped with OT-20 (3 individuals), OT-15 (7 individuals), and OT-10 (1 individual) data loggers, weighing 20, 15, and 10 g, respectively (Ornitela, Lithuania). Our study took place over a period of 7 years and we therefore aimed to use progressively lighter data loggers in line with technical developments (particularly logger weight reduction) over this time period. The mean body masses of female and male curlews were 957.7 ± 74.3 g and 827.8 ± 92.1 g, respectively. Even the heaviest data loggers used in this study therefore accounted for only about 2.4% of body mass, which was below the threshold of 3% suggested to avoid confounding effects of the devices [30]. The Skua devices only sent part of the data to a server via a GSM connection, which could then be downloaded directly, and the rest of the data were transmitted to base stations set up next to the high-tide roosts. The full dataset for the Skua devices was thus only obtained after the birds had returned to their wintering sites in the Wadden Sea. The highest temporal resolution achieved by these devices was 15 min. In contrast, the OT devices transmitted the whole dataset to an online portal, and the recording intervals were programmed according to a flexible schedule, generally ranging from 1–15 min, based on the battery status of the device. We programmed “geofences” (i.e. defined areas in which the devices recorded data constantly in 1-min intervals) for all OT devices. The geofence covered the red box shown in Fig. 1, but excluded high-tide roosts, to save battery power. The high temporal resolution of the GPS fixes allowed flight height measurements to be derived within the red box area shown in Fig. 1, and also further east.

All data recorded by the GPS devices were finally stored in the online portal Movebank (www.movebank.org).

In addition to equipping each bird with a GPS device, all individuals were ringed and weighed, bill and wing lengths were measured, age was determined, and sex was determined by taking a breast feather for genetic sexing in the lab (Tauros Diagnostics, Berlin).

Data analyses

We visualized the GPS data for each curlew using the Geographical Information System ArcGIS (version 10.3) [31]. The time (UTC) and day of the year at which the birds left their wintering grounds heading north-east towards their breeding sites were determined. This departure was evident from the GPS tracks,
and was always associated with a clear increase in flight speed (and flight heights in OT devices). The time and day of year of arrival at the wintering grounds was determined in a similar manner.

Meteorological data were recorded at 1-min intervals by an automatic recording station located at the Research and Technology Centre in Büsum, and weather conditions were related to the curlews’ departure and arrival patterns (see Statistical analysis section below). The recorded parameters were: temperature (°C), wind speed and maximum wind speed (m/s), wind direction (degrees), precipitation (mm), global radiation (W/m²), UVA and UVB (W/m²), photosynthetically active radiation (W/m²), air pressure (mbar), and air humidity (%). For all parameters, we computed the mean values of the 1-min recordings over a period from 90 min before to 90 min after the departure/arrival of each individual curlew. The mean values were then related to the departure/arrival day (see Statistical analysis section below) and used to test for differences in weather conditions between departures and arrivals. To account for the circular nature of the wind direction, means were calculated as the direction of a circular vector using the R-package circular [32]. Wind-rose plots of the wind direction during the departure and arrival of curlews were created using the R-package openair [33].

To relate the departure and arrival dates to linear distance from the breeding area, it was necessary to determine the location of the breeding area for each curlew. The likely breeding area was identified in GIS (white triangles in Fig. 4). It could be distinguished from stop-over locations by being the most distant point from the wintering location, and at the same time as a location where the birds stayed for several weeks (more than at each stop-over location) with very limited movement behaviour. The approximate coordinate of the nest site was calculated by computing the mean geographical position of all positions in the potential breeding site. Finally, the linear distance between this position and the wintering site was calculated in GIS.

Departure dates were also related to the linear and flown distances (calculated in GIS) to the first stop (red circles in Fig. 4), the flight time to the first stop, and the duration of the first stop. The same was applied for the arrival dates using the last stop before the wintering site (orange circles in Fig. 4). The mean locations of the nearest stops to the wintering sites were calculated in the same way as for the breeding site. In some cases, birds migrated from their wintering sites in the Wadden Sea to other Wadden Sea areas (always < 30 km distance), probably to join other birds shortly prior to departure. These locations were not regarded as first stop-over events, and the departure from the last site in the Wadden Sea was used for the analyses instead.

Finally, we computed the relative deviation between the flown and linear distances (%) as an indicator of the curvature of the actual flight track. This was expected to increase during headwind conditions, because birds might try to avoid headwinds by choosing different flight angles. Only flight tracks with log-intervals of ≤ 5 min were used for this to keep the flight tracks comparable.

We determined the mean departure direction of each individual across the red box shown in Fig. 1 and related it to the mean recorded wind speed and wind direction to compute the TWC. This is known to have a significant impact on the migration speed of birds [20–21, 34], and was therefore expected to affect the
departure and arrival decisions of the curlews in the current study. According to [35], we used the following formula: \( TWC = v \times \cos x \), where \( v \) is the wind speed in \( \text{ms}^{-1} \) and \( x \) is the angular deviation between the opposite flight direction of the curlew (i.e., tailwind direction) and the wind direction (in degrees). In addition to using \( TWC \) as an additional predictor of departure and arrival decisions, we also related it to mean flight speed to demonstrate if the birds were able to increase their speed during tailwind conditions, and to mean flight height within the red box in Fig. 1.

**Statistical analysis**

All statistical analyses were carried out using the open source software R, version 3.6.3 [36]. Plots were visualized using the R package `ggplot2` [37]. All regressions were performed using generalized additive models (GAMs) [38–39] using the R package `mgcv` [40]. Individual was included as a random intercept in the GAMs to avoid pseudo-replication due to multiple observations of the same individual. For each model, we selected an appropriate probability distribution for the variable of interest; if different probability distributions were reasonable (e.g. in the case of possibly overdispersed count data), we selected the most appropriate distribution based on the Akaike Information Criterion [41].

We related the departure day to meteorological data to see if departure decisions were affected by the weather. We considered the meteorological data under six conditions, including (1) at the moment of departure and (2) the mean conditions 90 min before to 90 min after departure. We also contrasted (3) the moment of departure and (4) the mean conditions 90 min before to 90 min after departure with the same measure on the 4 days before departure, to determine if curlews experienced suboptimal conditions prior to their departure, which might have delayed their departure decision. Finally, we also contrasted (5) the moment of departure and (6) the mean conditions 90 min before to 90 min after departure with the same measure at the same time of day on the same date for the average of the 4 previous years to determine if the curlews experienced suboptimal meteorological conditions compared with the average conditions on similar dates. In all cases, the meteorological conditions were compared by dividing the weather data at the moment of departure (respectively 90 min before to 90 min after departure) by the average conditions at the same moment (same time interval) for the 4 preceding days/years. If the meteorological variable of interest had a zero value, the difference was calculated instead of the quotient.

In addition to the above meteorological parameters, we also used the following additional predictors to model the departure decision (defined by day of the year): number of migrations for each individual bird, sex of the individual, catching location in the Wadden Sea, year, time of day, breeding latitude, linear distance to breeding area, departure direction, \( TWC \), time to first stop-over, duration of first stop-over, flown distance to first stop, and linear distance to first stop. For \( TWC \), we used the six different combinations of meteorological data given above. All other predictors were kept constant.

We therefore used a total of 73 predictors to model the departure decision (i.e. 55 constant and contrasted meteorological predictors, 6 constant and contrasted combinations of \( TWC \), and 12 constant predictors related to the individual curlews or the first stop-over event). The same predictors were used to model the arrival of the birds in their wintering grounds.
The effects of the 73 predictors on day of the year (outcome variable) were tested using the least absolute shrinkage and selection operator (LASSO) [42–43] technique for predictor selection. This technique is known to handle a large number of possible predictors without being prone to statistical problems e.g. compared with stepwise methods (c.f., below). Notably, LASSO has been combined with cross-validation to select promising predictors based on their predictive capacity [42–44]. In contrast to the common stepwise methods (e.g., forward or backward selection procedures), LASSO-based results are not sensitive to the order of the performed tests [44–45]. However, it is necessary to bear in mind that the chance of detecting a significant relationship between a predictor and the considered outcome variable increases with the number of investigated predictors.

**Results**

**Phenology of departures and arrivals**

The first curlew departed from the wintering grounds on April 7 and the last on May 16. Most individuals departed between mid- and late April (Fig. 2). Females tended to depart earlier, but the sex difference was not significant (GAM: $t = 1.23$, df = 41, $p = 0.23$). The first curlews arrived at the wintering sites on June 3 and the last on July 24, with most birds arriving between mid-June and the end of July (Fig. 2). There was a clear but not significant tendency for females to arrive earlier than males (GAM: $t = 1.97$, df = 31, $p = 0.067$; mean day of year females: 177.9 ± 13.1, mean day of year males: 189.9 ± 11.7).

Departures mainly occurred during the late evening, shortly before sunset, with no significant difference between the sexes (GAM: $t = -0.29$, df = 41, $p = 0.78$; Fig. 3a). The time of day for arrivals differed from that for departures (Table 1, Suppl. 1a), with significantly more arrivals during the nighttime and also occasionally during daytime. As for departing curlews, there was no sex difference in terms of arrivals (GAM: $t = 1.7$, df = 31, $p = 0.09$; Fig. 3b).
### Table 1

GAMs comparing migration parameters, and wind variables between departing and arriving curlews.

|                                | Estimate Std. | Error  | t-value | p-value |
|--------------------------------|---------------|--------|---------|---------|
| Time of day                    | −3.08         | 1.32   | −2.33   | 0.023   |
| Rel. diff. between flown and linear distance to nearest stop | −2.06         | 1.75   | −1.8    | 0.244   |
| Linear distance to nearest stop | −432.4        | 106.2  | −4.07   | < 0.001 |
| Time to nearest stop           | −189.47       | 118.16 | −1.6    | 0.114   |
| Duration of nearest stop       | 258.7         | 433.5  | 0.6     | 0.553   |
| Mean flight speed              | −18.53        | 4.12   | −4.49   | < 0.001 |
| Tail wind component            | −0.33         | 0.67   | −0.5    | 0.618   |
| Mean flight altitude           | −930.2        | 164.9  | −5.64   | < 0.001 |
| Mean wind speed                | −1.29         | 0.45   | −2.89   | 0.005   |
| Mean wind direction            | −2.2          | 27.43  | −0.08   | 0.936   |

Estimate Std = estimated standard deviation. GAM outputs are illustrated in Suppl. 1.

[Table 1 should be placed approximately here]

GPS tracks of curlews and relationships with nearest stop-over sites

After their departure from the wintering grounds in the Wadden Sea, all curlews headed towards their breeding sites, which were located exclusively in north-western Russia (except for one individual that was breeding in Finland; see white triangles in Fig. 4). The most distant breeding site was 3,840 km from the wintering site, east of the Ural Mountains. The mean linear distance of all flight tracks was 2,339 ± 612 km. The relative differences between the flown and linear distances to the breeding sites were similar for arriving and departing curlews (Table 1; Suppl. 1b).

The mean linear distance between the nearest stop-over site and the wintering site for departing curlews was 775.8 ± 376 km, which represents 33.2% of the linear distance to their breeding sites (i.e. birds performed about 1/3 of their overall migration during their first migration bout). Curlews selected a straight flight path to reach their first stop-over site, with the distance flown on average only 45.7 km longer than the linear distance (5.6%).

Many curlews crossed the Baltic Sea directly, while others stopped over on the Danish islands or the southern Baltic Sea coast (see red circles in Fig. 4). The nearest stop-over site for arriving curlews was significantly closer to the wintering site than that for departing curlews (mean: 342.4 km; Table 1; Suppl. 1b).
but the distance was highly variable (± 503.1 km). In contrast, both time to the nearest stop-over site and duration of the nearest stop were similar for departing and arriving curlews (Table 1; Suppl. 1d, e).

Finally, there was no significant relationship between flight time to the nearest stop-over and stop-over duration, suggesting that curlews did not take longer to recover after longer migration bouts.

Flight speed and flight height in relation to TWC

The flight speeds of both departing (GAM: t = 8.42, df = 32, p < 0.001) and arriving curlews (GAM: t = 5.07, df = 27, p < 0.001) were positively and highly significantly related to TWC (Fig. 5), suggesting that birds were able to increase their migration speeds with wind assistance. Interestingly, flight speeds during departure were significantly higher (mean: 73.8 ± 18.5 km/h, range: 41.9–115.6 km/h) than speeds during arrival (mean: 55.3 ± 11.9 km/h, range: 40.8–84.0 km/h), as indicated by the small overlap of confidence intervals in Fig. 5 (see also Table 1; Suppl. 1f). There was no difference in the TWC relationships between departing and arriving curlews (Table 1; Suppl. 1 g).

Curlews departed at significantly higher altitudes during headwind compared with tailwind conditions (GAM: t = −9.52, df = 19, p < 0.001), but there was no significant relationship between TWC and flight altitude in arriving curlews (Fig. 6; GAM: t = −1.25, df = 13, p = 0.25). As for flight speeds, flight altitudes were significantly higher and more variable during departure (mean: 1,113.3 ± 592.0 m, range: 175.2–2337.7 m) compared with during arrival (mean: 182.3 ± 164.4 m, range: 37.0–639.2 m) (Table 1; Suppl. 1 h).

There was no significant relationship between TWC and the relative difference between the own and linear distances, indicating that the straightness of the flight pass was not impacted by wind conditions. Finally, there was also no correlation between TWC and migration distance to the first stop or stop-over duration, suggesting that headwind conditions had no effect on flight distances and length of the first stop-over.

Departure/arrival decisions

The LASSO model showed that only one of the 73 predictors significantly influenced the departure day of curlews: curlews departed significantly later with increasing linear distance to their breeding sites (Fig. 7; GAM: t = 2.63, df = 37, p < 0.05). Wind direction at the time of departure contrasted with the mean wind direction at the same time of day, and day of year 4 years prior to departure remained a predictor in the final LASSO model, but had no significant impact on departure day (GAM: t = −0.77, df = 37, p = 0.45). Interestingly, LASSO did not select any other meteorological predictors, bird-related variables, or variables associated with the nearest stop-over event. Mean wind direction and wind force in the period from 90 min before to 90 min after departure were highly variable (Fig. 8a), which explains the absence of any significant relationships with day of departure.

According to LASSO, none of the 73 predictors was related to arrival day. As for departures, wind direction and force were highly variable (Fig. 8b). Wind force (but not wind direction) differed significantly between departure and arrival flights (Table 1; Suppl. 1i, j). During arrival, most of the higher wind forces were associated with north-easterly winds (Fig. 8b), which might have assisted some returning curlews, but
eventually had no significant impact. This also led to a lack of any significant difference in tailwind conditions between departing and arriving curlews (Suppl. 1 g).

In addition, there was no significant relationship between departure/arrival date and stop-over duration, indicating that curlews that migrated later did not have shorter stop-overs.

Repeatability:
Among the 42 curlews with departure information, we recorded multiple departures in subsequent years for 12 individual birds (2–5 consecutive years). It was therefore possible to assess the repeatability of the departure day in the same individuals in different years. The mean absolute difference in departure days in subsequent years was only 3.68 ± 2.97 days (n = 19). However, the variability in arrival days of returning curlews was more than twice as high (7.17 ± 4.83, n = 12). The departure days recorded in each individual’s first year were significantly related to the departure days in subsequent year(s) (Fig. 9; GAM: t = 5.29, df = 18, p < 0.001), while there was no significant relationship for arrival dates (GAM: t = −0.5, df = 11, p = 0.62).

There was also high site fidelity in terms of the locations of breeding and wintering sites, but less fidelity for the location of the nearest stop-over sites (Fig. 4). The repeatability of flight directions was also high, with a mean variability of only 9.7 ± 8.2° for departures and 13.3 ± 10.2° for arrivals among subsequent years for the same individuals.

Discussion
Wind and weather effects and flight heights

Our data confirmed that curlew flight speed increased with increasing TWC, in accordance with previous studies showing similar relationships [6, 20]. Based on the clear benefit of faster flight speeds during tailwind conditions, we hypothesized that curlews would mainly select days with suitable tailwind conditions for their departure from (and arrival at) their wintering grounds in the Wadden Sea. This relationship was demonstrated in previous studies of songbirds [e.g. 12, 22], while long-distance migrating shorebirds were found to encounter unfavourable and potentially fatal wind conditions if they failed to time their departure precisely [10]. However, in contrast to our assumption, the current study revealed virtually no effect of wind or other weather parameters on the departure/arrival day for curlews, even when the actual weather during departure was compared with the mean weather conditions 4 days and/or 4 years prior to departure. Although curlews rarely departed during hours with precipitation (in accordance with other bird species) [12, 23–24], this predictor had no significant influence in our final model. The current findings were therefore not in agreement with previous studies, which found that the probability of departure of curlews from a pre-breeding staging site was reduced during precipitation [46]. In contrast, our findings were in accord with observations of departing curlews in China, where there was also no significant influence of wind force or wind direction [46]. The same holds true for the closely related whimbrel (*Numenius phaeopus islandicus*), for which there were no correlations with wind conditions for either direct flight or one-stop-over migration [47].
Although weather parameters had no impact on departure/arrival decisions, we found a significant negative correlation between TWC and flight altitude in curlews. Although there were no available data for wind speeds (and directions) of different air layers for this study, this pattern clearly suggests that curlews tried to find more favourable wind conditions at higher altitudes if they encountered headwinds at lower altitudes. In temperate latitudes, the prevailing westerly wind conditions in the higher air layers suggest wind assistance when ascending [48]. This behaviour has previously been recorded for nocturnal songbird migrants [25–26], as well as for diurnal long-distance migrants using radar techniques [49]. [50] found intensive songbird migration in air layers up to 3 km altitude in temperate regions, when the birds encountered headwind conditions close to the surface. The authors demonstrated that migrating birds benefited from the wind conditions in higher air layers by ascending, exclusively during their spring migration. This might explain why there was no significant relationship between flight altitude and TWC in arriving curlews during their autumn migration. The current results clearly suggest that curlews depart (and stay) at lower altitudes when wind conditions close to the surface are beneficial, and use higher air layers during spring migration when they encounter headwinds.

Interestingly, curlews arrived at significantly slower flight speeds and lower altitudes compared with departing curlews. Meteorological reasons for this can be excluded, given that the wind conditions and TWC were similar for departing and arriving individuals. One likely reason is that the linear distance to the nearest stop-over in arriving curlews was far smaller than for departing individuals, which might explain why departing curlews ascended to higher altitudes and had faster flight speeds compared with arriving birds.

In conclusion, the absence of clear relationships between departure decision and wind/weather parameters suggests that curlews behave conservatively and follow an individual time schedule driven by an internal genetic trigger [14, 51–52] (see discussion on repeatability below). If their individual departure day happens to coincide with headwind conditions, the birds seem to account for this by flying at higher altitudes. However, further studies are needed to determine if curlews have knowledge of the wind conditions at higher altitudes when they encounter headwinds on the ground.

Temporal patterns and location of breeding sites

According to our LASSO analysis, distance to the breeding site was the single (highly significant) predictor affecting the day of departure of curlews from the Wadden Sea. Counterintuitively and in contrast to our formulated hypothesis, birds with the longest migration distances showed the latest departure and arrival days. Given the long migration distance, we expected these curlews to leave earlier to allow them to complete their journey to their breeding grounds in time to start breeding. However, shorebirds have been shown to encounter higher food availability and have faster chick growth rates if they start nesting soon after snowmelt [53–54], suggesting that curlews should aim to arrive at their breeding sites as soon as the areas are free from snow and ice. This could in turn explain why curlews that breed further from their wintering grounds (e.g. in the eastern parts of Russia in this study) might wait longer to allow the snow and ice in their Arctic breeding grounds to melt, and/or to ensure that they
encounter optimal arthropod densities. Similarly, [19] showed that colour-ringed curlews breeding in Fennoscandia departed later than birds breeding further west. Although this study dealt with a different sub-population, the results matched the patterns found in the current study.

We found that curlews wintering in the Wadden Sea departed within a very short time window (i.e. mostly between mid-April and mid-May). This contrasts with birds wintering in south-west Britain, which had already started to depart during February and March [19]. However, in contrast to British curlews that breed in north-western Europe [19], all but one of the curlews reported in the present study bred in Russia, i.e. much further east. The more condensed departure window in our study might thus also be a consequence of the relatively late availability of breeding sites due to snow and ice melt. This emphasises the need for the birds to have an internal clock to ensure their timely departure.

The main window of arrival of birds in the Wadden Sea was late June to mid-July, which is about 2 weeks later than reported for birds breeding in central or northern Europe [19]. Although desertion of offspring by females is common in shorebirds and has been shown for curlews [27], we found no sex differences in departure patterns, and only a non-significant tendency for females to arrive earlier. The reason for this is unclear. It is possible that some birds failed to rear chicks successfully, leading to the earlier arrival of at least some males.

Departures of songbirds usually occur at night and around sunset [22, 51] and the same is true for many long-distance migratory wader species, given that birds are thought to take advantage of favourable atmospheric factors during the night and to calibrate their orientation systems before they start in the evening hours [55–56]. Our results support these patterns, with more curlews departing during the early evening or early nighttime. However, an earlier study of curlews departing from a final pre-breeding stop-over site in China showed high variability in terms of the time of day for departures [46]. The reason for these different findings remains unclear. Given the higher number of individuals arriving during morning and afternoon, our results suggest the existence of a strong temporal trigger regulating departure decisions, but a weaker such trigger for arrivals.

Correlations with first and last stop-overs

We expected that the departure day and TWC would be significantly related to the distance to the nearest stop-over site and the stop-over duration; however, no such correlation was found. Curlews did not stage for shorter periods if they departed later, nor did they stage for longer if they encountered headwind conditions during the first part of their migration to allow more time for re-fuelling. This finding is in accordance with studies of songbirds, which likewise showed no or only weak relationships [57–58]. Similar results were also found for whimbrels, with no impact of wind conditions on stop-over patterns; however, in contrast to our study, they tended to skip a potential stop-over when they departed later [47].

Repeatability
In a recent review, [52] presented evidence for genetic control of the timing of bird migration. However, the authors also found much individual variation in this genetic programme as a result of interactions with environmental and social factors, as well as due to individual learning. The curlews in the current study also demonstrated a clear genetic circannual rhythm, given that individual curlews showed very little variation in departure and arrival patterns among subsequent years, and did not time their departure according to the weather. The current results thus provide a robust example of the concept of an internal clock, responsible for timing bird migration [14, 51]. In contrast to these results however, black-tailed godwits (Limosa limosa limosa) tracked across subsequent years exhibited a much broader window of inter- and intraspecific timing of migration [59]. The authors suggest that this broad window of repeatability indicated weak selective forces with respect to migration timing. This assumption would suggest that such forces are strong for curlews, although for the specific mechanisms affecting the high inter- and intraspecific repeatability of migration timing evidence is currently lacking. In contrast, [8] hypothesized a conservative annual-cycle strategy in long-distance migrants that was thought to minimise risks and reduce carry-over effects. Our study may support this hypothesis, given that curlews clearly showed constant, conservative patterns in terms of their departure decisions, independent of the weather conditions at their wintering sites. These findings are in line with other Arctic breeding birds, such as bar-tailed godwits (Limosa limosa baueri) from New Zealand, which also showed high repeatability [60].

Conclusions

The results of the current study demonstrated constant temporal departure and arrival patterns in curlews, with no significant impact of meteorological conditions. The high repeatability of the departure day in the same individuals in subsequent years suggests a genetic trigger regulating the timing of migration. These results clearly show that curlews departed during any weather conditions, including headwinds, which seemed to be accounted for by flying at higher altitudes. However, the main factor driving the timing of migration seems to be the date, suggesting the existence of an internal clock prompting the start of migration. Follow-up studies are needed to explore the timing of migration in immature curlews, ideally by tagging birds from the same clutch and their parents. Further insights into the timing of migration in closely related birds might help us to understand the genetic mechanisms triggering temporal migration patterns.

Abbreviations

GPS
Global Positioning System
GSM
Global System for Mobile Communications
TWC
tailwind component
Declarations

Ethics approval and consent to participate

Permission to attach GPS tags was issued by the Ministerium für Energiewende, Landwirtschaft, Umwelt, Natur und Digitalisierung of the federal state of Schleswig-Holstein (file numbers V 312-7224.121-37(42-3/13) and V 241-35852/2017(88-7/17)) as well as by the Niedersächsisches Landesamt für Verbraucherschutz und Lebensmittelsicherheit of the federal state of Lower Saxony (file number 33-19-42502-04-17/2699). Catching and treatment of birds complied with current EU and German laws. All authors consent to participate in this study.

Consent for publication

Not applicable.

Availability of data and materials

The tracking data of curlews are available in www.movebank.org.

Competing interests

The authors declare that they have no competing interests.

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Author’s contributions

PS designed the study, conducted the fieldwork, performed the data analyses and wrote most parts of the manuscript. MM designed and performed the statistical tests and wrote parts of the method section. KHV recorded and analysed the meteorological data. PB wrote parts of the manuscript. SG designed the study and compiled the movement data. All authors read and approved the final manuscript.

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