Quantification of within- and between-farm dispersal of Culicoides biting midges using an immunomarking technique

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Summary

1. Culicoides biting midges (Diptera, Ceratopogonidae) are vectors of arboviruses that cause significant economic and welfare impact. Local-scale spread of Culicoides-borne arboviruses is largely determined by the between-farm movement of infected Culicoides.

2. Study of the dispersal behaviour of Culicoides by capture–mark–recapture (CMR) is problematic due to the likelihood of mortality and changes in behaviour upon capture caused by the small size and fragility of these insects, evidenced by low recapture rates. To counter the problem of using CMR with Culicoides, this study utilised an ovalbumin immunomarking technique to quantify the within- and between-farm dispersal of Culicoides in southern England.

3. Both within- and between-farm dispersal of Culicoides was observed. Of the 9058 Culicoides collected over 22 nights of trapping, 600 ovalbumin-positive Culicoides, of 12 species including those implicated as arbovirus vectors, were collected with a maximum dispersal distance of 3125 m.

4. This study provides the first species-level data on the between-farm dispersal of potential bluetongue, Schmallenberg and African horse sickness virus vectors in northern Europe. High-resolution meteorological data determined upwind and downwind flight by Culicoides had occurred. Cumulative collection and meteorological data suggest 15.6% of flights over 1 km were upwind of the treatment area and 84.4% downwind.

5. Synthesis and applications. The use of immunomarking eliminates the potential adverse effects on survival and behaviour of insect collection prior to marking, substantially improving the resolution and accuracy of estimates of the dispersal potential of small and delicate vector species such as Culicoides. Using this technique, quantification of the range of Culicoides dispersal with regard to meteorological conditions including wind direction will enable improved, data-driven modelling of the spread of Culicoides-borne arboviruses and will inform policy response to incursions and outbreaks.

Key-words: African horse sickness virus, arbovirus, bluetongue virus, capture–mark–recapture, Ceratopogonidae, epizootic-haemorrhagic disease virus, ovalbumin, protein marking, Schmallenberg virus, vector-borne disease

Introduction

Culicoides biting midges (Diptera, Ceratopogonidae) are vectors of arboviruses that cause significant economic and welfare impact to the livestock industry, most notably the incursion of multiple serotypes and strains of bluetongue virus (BTV) into Europe (Purse et al. 2015). The dispersal of hematophagous arthropod vectors is a key factor in determining the spread of arboviruses (Sellers 1980). Long-distance dispersal of Culicoides overwater has been linked to incursions of Culicoides-borne arboviruses hundreds of kilometres from the nearest virus source and where introduction by movement of viraemic mammalian hosts was discounted (Sellers, Pedgley & Tucker 1978; Alba, Casal & Domingo 2004; Burgin et al. 2012). In
contrast, studies examining overland spread of BTV by correlating wind-density maps and outbreak data (Hendrickx et al. 2008) and stochastic simulation of likely time of infection and wind speeds and direction (Sedda et al. 2012) have found limited evidence for long-distance dispersal of Culicoides. In both studies, the model prediction of distance for arbovirus spread was less than 5 km day\(^{-1}\) and was largely determined by the between-farm movement of infected Culicoides (Hendrickx et al. 2008; Sedda et al. 2012). Quantification of the range and probability of Culicoides dispersal within- and between-farms is essential to allow accurate estimation of the speed and extent of spread of Culicoides-borne arboviruses and to inform policy response during incursions and outbreaks (Defra 2012). Quantification of the range and probability of Culicoides dispersal within- and between-farms is essential to allow accurate estimation of the speed and extent of spread of Culicoides-borne arboviruses and to inform policy response during incursions and outbreaks (Defra 2012).

A variety of methods have been used to quantify Culicoides dispersal at a local scale. Movement from isolated areas of larval habitat has been used where these can be readily identified and demarcated (Kettle 1960; Williams 1962; Zimmerman & Turner 1984). Culicoides have also been marked using of radioactive isotopes (Davies 1965; Holbrook, Belden & Bobian 1991) and ingestion of food dye (Campbell & Kettle 1976), however these techniques utilised emerging adults and are thus reliant on the availability of larval habitat. The putative vectors of BTV in the northern Palaearctic; Culicoides obsoletus Meigen, Culicoides scoticus Downes and Kettle, Culicoides dewulfi Goetghebeur, Culicoides and C. mohave has also been carried out (Wirth (Brenner 2013). In this study, we utilise the ovalbumin immunomarking technique to mark pollinator, phytophagous and predaceous insects (Hagler 2011; Biddinger et al. 2013; Swezey et al. 2013). Immunomarking has also been used on Anaphes iole Girault (Hymenoptera: Mymaridae), minute parasitoid wasps that at 1 mm in length are smaller even than Culicoides (Hagler & Jackson 1998). Sanders & Carpenter (2014) have previously demonstrated that an ovalbumin marker persisted on Culicoides nubeculosus (Meigen) for 72 h post-exposure with no detrimental effects on survival. A preliminary field trial also demonstrated that ovalbumin-marked subgenus Avaritia Culicoides could be recovered from areas close (≤20 m) to marked larval development habitat (Sanders & Carpenter 2014). In this study, we utilise the ovalbumin immunomarking technique for the first time at a broader-scale to quantify the within- and between-farm dispersal of Culicoides in southern England.

**Materials and methods**

**STUDY SITE AND HABITAT IMMUNOMARKING**

Habitat immunomarking was conducted at a mixed sheep and beef cattle farm in southern England (51°43’39”N; 1°23’16”W) (Fig. 1a). Surrounding the farm were additional livestock holdings to the south and east, with arable crops to the west and a residential area and water reservoirs to the north (Fig. 1b). Two areas (total of ~60 m\(^2\)) of mixed cattle dung and straw bedding along both sides of an open-sided shed (Fig. 2) were sprayed 2–3 h prior to sunset on day 0 using a hand-operated knapsack sprayer (Silverline, Yeovil, UK) with 40 L of 20% (w/v) liquid egg white solution (Sports Supplements Ltd, Colchester, UK), resulting in an approximate application rate of 0.33 Lm\(^{-2}\). Insects were then collected for 3 or 4 days post-treatment. To prevent cross-contamination, different personnel and equipment
were used to mark habitat to those responsible for insect collection. Five egg solution treatments were conducted; two during 2013 and three during 2015, with a minimum period of 2 weeks between the last day of insect collection and the next treatment.

**METEOROLOGICAL CONDITIONS**

Meteorological conditions [temperature (°C), wind speed (ms⁻¹), wind direction (°), relative humidity (%), solar radiation (Wm⁻²) and precipitation (mm)] were monitored at 1–15 min resolution using an automatic weather station located 30 m from the marked area and analysed using the openAir (Carslaw & Ropkins 2012) package in R v. 3.2.2 (R Development Core Team 2015) with RStudio v. 0.99.467 (RStudio Team 2015).

**CULICOIDES COLLECTION**

Insect collections were completed overnight using ultraviolet miniature Center for Disease Control traps (model: 912; John W Hock, Gainsville, FL, USA) suspended at approximately 1.2 m from metal hooks (Gardman Ltd, Peterborough, UK). All traps were located at field margins and hedgerows beyond the reach of livestock. Insects were collected dry into disposable, single use 560 mL polypropylene cups that contained a small quantity of shredded paper towel based on a preliminary study to minimise reduce contact and therefore potential marker transfer between collected specimens (Fig. S1, Supporting Information). Collected insects were immobilised by exposure to cold (−20 °C) and subsequently stored at −20 °C prior to further analysis.

Fourteen trap locations were used in two trial replicates completed in 2013 (locations 1–14, Fig. 1). Trap locations two and three formed a short linear transect (T1) extending southeast from the marked area (Fig. 1). In three replicates in 2015, the 14 trap locations utilised in 2013 were supplemented with an additional 10 trap locations which formed three linear transects extending out from the egg-marked area to a maximum distance of 3876 m (Fig. 1). These three transects extended through the livestock pastures of adjacent farms and represent potential dispersal routes to the available livestock hosts of dairy cattle (T2), beef cattle (T3) and sheep (T4) respectively.

**OVALBUMIN-SPECIFIC ELISA**

An ovalbumin-specific ELISA modified from Hagler & Jones (2010) was utilised to detect ovalbumin-positive Culicoides.
Ovalbumin-positive controls were created by either soaking a 225 mm² piece of paper, which had been dipped in egg white solution, in 300 μL of tris buffered saline (TBS) (pH 7-4) (Alfa Aesar, Heysham, UK) with 0.001% Silvet L-77 (De Sangosse Ltd, Cambridge, UK) (TBS-SILWET) (Sanders & Carpenter 2014) or via the addition of 3 μL egg white solution per 300 μL TBS-SILWET. From this, a 10-fold serial dilution was created in order to monitor relative detection sensitivity between assay runs. 

Culicoides were identified morphologically to species or subgenus/group level using the keys of Campbell & Pelham-Clinton (1960) and reference wing images (The Pirbright Institute 2007). Culicoides identified from their morphology were then transferred individually using disposable cocktail sticks to tubes containing 300 μL TBS-SILWET and incubated at 4 °C overnight.

Subsequently, 80 μL of the sample solutions were transferred to the corresponding wells of an ELISA plate. Negative controls (80 μL TBS-SILWET) and positive controls (10⁶, 10⁻¹, 10⁻², 10⁻³ and 10⁻⁴ dilutions) were then added to the remaining wells of the ELISA plate. The ELISA plate was then incubated at 37 °C for 1 h; washed five times with phosphate-buffered saline with tween-20 (PBST) (Sigma-Aldrich, Gillingham, UK); 360 μL of phosphate-buffered saline with bovine serum albumin (PBS-BSA) (Sigma-Aldrich) added per well; incubated at room temperature for 1 h; washed twice with PBST; 80 μL 0.0125% rabbit anti-chicken egg albumin (Sigma-Aldrich) in PBS-BSA with 0-13% (w/v) Silvet L-77 (PBS-BSA-SILWET) added per well; incubated for 1 h at 37 °C; washed five times using PBST; 80 μL of 0.05% goat anti-rabbit IgG conjugated to horseradish peroxidase (Sigma-Aldrich) in PBS-BSA-SILWET added per well; incubated for 2 h at 37 °C; washed five times using PBST; 80 μL of 1-Step Ultra TMB-ELISA Substrate solution (ThermoFisher Scientific, Warrington, UK) added per well; incubated at room temperature for 10 min followed by the addition of 80 μL 2N sulphuric acid (Sigma-Aldrich) to stop the reaction. The ELISA optical density (OD) of each well was measured at 450 nm with a microplate reader.

All ODs were transformed using the standard normal variate (SNV) transformation (Sivakoff, Rosenheim & Hagler 2011). The threshold score was set using the ‘maximum negative control’ algorithm i.e. the highest negative control score observed across all ELISA plates in the study (Hooper & Woolson 1991; Sivakoff, Rosenheim & Hagler 2011). Specimens were classified as ovalbumin-positive if their score was greater than the selected threshold.

**Avaritia subgenus species identification**

Following the ovalbumin-specific ELISA, subgenus Avaritia specimens collected in replicates 3-5 were transferred individually to 200 μL of 50% Chelex® 100 resin (Bio-Rad, Hemel Hempstead, UK) solution with 50% proteinase K (Qiagen Ltd, Manchester, UK); homogenised at 25 Hz for 4 min using a tissue-lyser (Qiagen Ltd) and the homogenate incubated overnight at 37 °C to extract total DNA. A multiplex polymerase chain reaction (PCR) assay was used then to identify samples to species level (Nolan et al. 2007). Amplification reactions were performed in a total of 20 μL consisting of 0.6 μL nuclease free water, 10-0 μL Qiagen® Fast Cycling PCR Mastermix (Qiagen Ltd), 2.0 μL CoralLoad concentrate (Qiagen Ltd), 2.0 μL 50% D(+)-Trehalose (Sigma-Aldrich), 0.4 μL 50 mM magnesium chloride (Qiagen Ltd), 40 mM forward primers: 0-1 μL UOAchIF, 0-1 μL UOAchWF, 0-1 μL UOAsobF and 0-2 μL UOAscoF (Nolan et al. 2007) and 0-5 μL 40 mM reverse primer C1-N-2191 (Dallas et al. 2003) and 4 μL Chelex supernatant (DNA template). Positive and negative controls for the amplification reactions were carried out at every PCR round. The PCR thermal cycle included: an initial denaturation step at 95 °C for 5 min followed by 35 cycles of 96 °C for 5 s, 48 °C for 5 s, 72 °C for 15 s, followed by a final extension step at 72 °C for 1 min. Amplification success and banding patterns were assessed by electrophoresis of PCR products on 2% (w/v) pre-cast agarose gels containing SYBR® Safe (E-Gel® 96; ThermoFisher Scientific).

**Statistical analysis**

Generalised linear mixed models (GLMM) with a Binomial error distribution and a logit link function were used to investigate the influence of meteorological conditions and the spatial relationship between trap location and the egg-marked area in determining the presence of ovalbumin-positive Culicoides within-trap catches. Models were implemented in a Bayesian setting using the bglmer function in package ‘blme’ v. 1.0-2 (Dorie 2014) in R v. 3.1.2 (R Development Core Team 2015) with RStudio v. 0.99.467 (RStudio Team 2015).

The GLMMs were fitted by maximum likelihood with the Laplace approximation with flat covariance priors and normal fixed priors, with days since egg marking included as a random effect, and a total of five additional fixed predictors including: distance from egg-marked area, difference between trap bearing and mean wind direction, mean air temperature, mean relative humidity, mean precipitation, mean solar radiation, mean wind speed, maximum gust speed as linear functions were considered. The fixed predictors were zero-centred and scaled to yield a standard deviation of one. Final models were obtained using a backwards-stepwise-selection-based procedure (Zeileis, Kleiber & Jackman 2008), such that variables that did not contribute significantly to explaining variation in trap catch were successively eliminated on the basis of Akaile information criterion (AIC) (Akaike 1973). This continued until the removal of a variable caused an increase in AIC of two or more.

**Results**

A total of 9058 Culicoides were collected over 22 nights of trapping (Fig. 3, Table S1). In 2013, 140 trap collections were completed with 7526 Culicoides collected from 14 trap locations over 10 nights of trapping. In 2015, 288 trap collections were completed with 1532 Culicoides collected from 24 trap locations over 12 nights of trapping (Table S1). Culicoides collections were dominated by female Culicoides with only 1% of Culicoides collected being male (Table S1). In total, 14 Culicoides species were identified: Culicoides achravi Kettle and Lawson, Culicoides albicans (Winnertz), C. chiopterus, Culicoides circuncrisspus Kieffer, Culicoides clastrieri Callot, Kremer and Deduit, C. devulfi, Culicoides festivipennis Kieffer, C. nubeculosus, C. obsoletus, Culicoides pictipennis (Staeger), C. pulicaris, C. pancutatus, Culicoides riethi Kieffer and C. scoticus (Figs 3 and 4, Table S1). Trap catches were dominated by subgenus Avaritia species, which accounted for 74%-0% of the total number of Culicoides.
of the other Culicoides species collected, C. punctatus, C. pulicaris, C. achrayi and C. festivipennis were the most abundant in trap catches accounting for 8.6%, 7.1%, 6.9% and 2.9% of the total number of Culicoides collected, respectively (Table S1). The remaining six species accounted for only 0.6% of the total number of Culicoides collected (Table S1). Subgenus Avaritia specimens collected in replicates three, four and five, which were identified to species level were dominated by C. obsoletus (52.5%), with 35.5%, 9.4% and 2.6% C. scoticus, C. dewulfi and C. chiopterus collected respectively (Fig. 4, Table S1).

A total of 600 Culicoides from ten species were observed to be positive for ovalbumin; C. achrayi, C. chiopterus, C. dewulfi, C. circunscriptus, C. festivipennis, C. nubeculosus, C. obsoletus, C. pulicaris, C. punctatus and C. scoticus (Figs 3 and 4, Table S2). The maximum distance from the egg-marked area that Culicoides were collected and observed to be ovalbumin-positive was 3125 m (Table 1, Figs 5 and 6). Ovalbumin-positive Subgenus Avaritia Culicoides were observed at the maximum distance of 3125 m from the egg-marked area as early as day 1, however positive C. pulicaris individuals were not collected at this distance until day 3 (Table 1). In replicates three, four and five of the trial where subgenus Avaritia individuals were identified to species level, C. dewulfi, C. obsoletus and C. scoticus were recorded at 3125 m at day 1; however, C. chiopterus was not recorded at this distance until day 2 (Table 1). Ovalbumin-positive C. nubeculosus individuals were collected at a maximum distance of 2542 m by day 4, and C. punctatus individuals at a maximum of 710 m by day 3. Ovalbumin-positive specimens from all other species were collected ≤406 m from the egg-marked area (Table 1).

All traps at a distance of <500 m collected ovalbumin-marked individuals. In 2015, all transects extending beyond 1 km had traps that collected positive individuals, with the greatest number collected along transect three (Figs 5 and 6, Table S2).

Mean air temperature, mean relative humidity, mean solar radiation, mean wind speed and distance from the egg-marked area were all found to be significant determinants of the likelihood of ovalbumin-positive Culicoides being collected (Table 2). Within the range of meteorological conditions experienced during this study (Fig. 7, Table S3), increasing mean air temperature (P ≤ 0.001), mean relative humidity (P ≤ 0.01) and mean solar radiation (P ≤ 0.01) were found to significantly increase the likelihood of ovalbumin-positive Culicoides being collected within a trap catch (Table 2). Increasing mean wind speed (P ≤ 0.001) and increasing distance from the egg-marked area (P ≤ 0.01) were found to significantly decrease the likelihood of ovalbumin-positive Culicoides being collected within a trap catch (Table 2).

Wind direction was not found to be a significant determinant of the probability of collecting ovalbumin-positive Culicoides, demonstrated by the observation of both upwind and downwind flight from the treatment area (Fig. 7). Of the ovalbumin-positive Culicoides collected at more than 1 km from the treatment site on the first trapping session of each replicate, one was observed to have flown upwind and 11 downwind. Subsequent collections were not independent of the wind direction of previous sessions within the replicate, and therefore flight direction cannot be determined but cumulative data for wind direction and collection show that of individuals collected at distance greater than 1 km from the treated area, 15.6% were upwind and 84.4% downwind of the treated area (Fig. 7).

In addition to Culicoides, 27 female Culex mosquitoes were collected of which 18 were found to be positive for ovalbumin from trap locations two, three, four, five, six, seven and eight.
seven, 10 and 13. Ovalbumin-positive *Culex* were collected up to a maximum of 698 m on day 3 from the egg-marked area, the maximum distance from the egg-marked area ovalbumin-positive *Culex* was recorded on day 1 post-egg-spraying was 314 m.

**Discussion**

This study uses a habitat marking approach to provide the first, species-level data on the within- and between-farm dispersal of Palaearctic *Culicoides*. Previous estimates of the dispersal potential of Palaearctic *Culicoides* had been made from CMR studies with very limited data from the recapture of a few marked individuals (Kirkeby et al. 2013; Kluiters, Swales & Baylis 2015). In contrast, this study used an immunomarking technique to mark insects in and around a potential larval development site and collected 600 marked *Culicoides*.

The majority (94%) of marked *Culicoides* were collected within 1 km of the treatment area, where trap density was greatest; suggesting most individuals which had emerged from or visited the marked habitat subsequently engaged in short appetitive flights to locate hosts, mates or breeding habitat. The fact that *Culicoides* were intercepted within traps at close range before they were able to disperse may lead to an underestimation of the potential for

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**Table 1.** Maximum dispersal distance of collected *Culicoides* by day and species/species group (*–* indicates specimens positive for ovalbumin not collected on this day in any replicate or trap catch)

| Species          | Maximum observed dispersal distance (m) |
|------------------|----------------------------------------|
|                  | Day 1       | Day 2       | Day 3       | Day 4       |
| *Subgenus Avaritia* |            |            |            |            |
| *Culicoides* species* |            |            |            |            |
| *C. chiopterus*† | 3125        | 3125        | 3125        | 3125        |
| *C. dewulfi*†    | 3125        | 247         | 1377        | 6           |
| *C. obsoletus*†  | 3125        | 1385        | 3125        | 3125        |
| *C. scoticus*†   | 3125        | 1385        | 3125        | 1377        |
| *C. aacrayi*     | 314         | –           | 314         | –           |
| *C. circumscriptus* | 406        | –           | –           | –           |
| *C. festivipennis* | 406        | 314         | 314         | –           |
| *C. nubeculosus* | 698         | –           | 6           | 2542        |
| *C. pictipennis* | 314         | –           | –           | –           |
| *C. pulicaris*   | 406         | 406         | 3125        | 3125        |
| *C. punctatus*   | 406         | 307         | 710         | 6           |

*Based on data from replicates 1–5. Includes number of *Culicoides* displayed in *C. chiopterus*, *C. dewulfi*, *C. obsoletus* and *C. scoticus* rows.

†Species identifications based on multiplex PCR assay of specimens collected in replicates three, four and five only.
long-distance movement. Despite the low density of traps, 6% of the ovalbumin-positive insects were collected in traps 1.3–3.1 km from the marked area, with *C. dewulfi*, *C. obsoletus* and *C. scoticus* demonstrating a maximum-recorded dispersal distance of more than 3 km over just one night. Ovalbumin-positive *Culicoides* were collected from all three transects, in areas used for grazing both cattle and sheep, with the greatest number collected in the trap near the greatest density of cattle suggesting individuals were engaged in active host-seeking behaviour. *Culicoides pulicaris* and *C. nubeculosus* individuals were collected at the maximum distance only 3 or 4 days post-marking, with *C. punctatus* and other species not recorded at distances greater than a kilometre from the marked area. This contrasts with the previous dispersal data of *Pulicaris* group where individuals were recaptured at
1750 m from the release site two or three nights after release (Kirkeby et al. 2013). This may reflect the lower abundance of these species in this study area and therefore the reduced probability of a marked individual being captured. Topographical features of the study area including hedgerows, trees and a river did not appear to influence the direction of Culicoides dispersal. The within- and between-year variation in difference in Culicoides abundance at the site may be due to increased drying of the larval habitat in the summer of 2015 in comparison with 2013 when the site had been flooded over the winter.

In previous studies, the low number of individuals recaptured (40 in total) at all distances obscures the estimation of local movement of the majority of Culicoides individuals (Kirkeby et al. 2013; Kluiters, Swales & Baylis 2015). Using the SNV transformation of ELISA data did not affect the maximum dispersal distance of marked Culicoides. It did, however, have a significant effect on the proportion of trap catch recorded as positive for ovalbumin marker, with 27.1% and 6.8% positive using the conventional threshold and SNV transformation respectively (Table S4). Whilst this may result in a conservative calculation of the number of marked individuals caught, the data are more robust and can be used with greater confidence.

Whilst the immunomarking technique used within this study offers considerable insights into behaviours that were previously precluded by the small-size and fragility of Culicoides, the technique does have limitations. The potential transfer of the marker protein between marked and unmarked individuals in a trap catch is of concern as a source of false positives (Hagler, Machtley & Blackmer 2015). When tested in the laboratory under optimal conditions, the transfer of ovalbumin between colony C. nubeculosus was limited, with evidence of transfer in only one replicate at a high ratio of marked to unmarked individuals (15 marked : 50 unmarked) (Fig. S1). In this study, marker transfer between Culicoides would not affect interpretation of the distance travelled by marked insects. However, in studies where the number of marked individuals would have a significant bearing on interpretation of results, the potential for marker transfer should be considered.

As the number of individuals marked is unknown, the proportion of the population that engaged in dispersal and estimates of survivorship and migration (Lysyk & Axtell 1986; Hanski, Alho & Moilanen 2000) cannot be estimated from the data in this study. For arbovirus vectors, even rare dispersal events may have significant

**Table 2.** Regression coefficients with 95% Wald confidence intervals and ΔAIC for the fixed effects of the final Bayesian general linear mixed models with a Binomial error distribution used to describe the presence of ovalbumin-marked Culicoides (all species) (random effects: days since egg treatment)

| Parameters                          | Estimate (95% CI)     | ΔAIC |
|-------------------------------------|-----------------------|-----|
| Intercept                           | –3.48 (–4.39 to –2.57)** |     |
| Mean air temperature                | 2.83 (1.50–4.17)***    | 18.34 |
| Mean relative humidity              | 1.33 (0.43–2.22)**     | 5.71 |
| Mean solar radiation                | 2.13 (0.81–3.47)**     | 9.04 |
| Mean wind speed                     | –2.30 (–3.63 to –0.97)*** | 12.22 |
| Distance from egg-marked area       | –1.73 (2.85 to –0.62)** | 9.66 |

**p ≤ 0.01, ***p ≤ 0.001.

Fig. 7. Locations of ovalbumin-positive Culicoides collections and windroses displaying cumulative mean windspeed (ms⁻¹) as a function of wind direction (°) summarised by replicate. [Colour figure can be viewed at wileyonlinelibrary.com]
impacts on epidemiology. The 6% of the marked individuals collected on adjacent pastures at greater than 1 km imply Culicoides on lowland farms represent well-connected populations with significant mixing between farms. Within replicates three to five, traps at a distance greater than 1 km collected 59% of marked individuals. The rapid spread of BTV and particularly SBV across northern Europe (Carpenter, Wilson & Mellor 2009; Elbers et al. 2012; Balenghien et al. 2014) has been used to infer a substantial movement of Culicoides between farms. The present data support previous estimates of dispersal distance of 3–5 km per day of infected Culicoides (Ducheyne et al. 2007; Hendrickx et al. 2008; Sedda et al. 2012; Sedda & Rogers 2013). There is, however, evidence of population-scale differences in phylogeny in Culicoides (Jacquet et al. 2016) that suggests the presence of barriers to the dispersal of Culicoides at a greater scale than represented in this study.

Collection of ovalbumin-positive Culicoides was not significantly influenced by wind direction, with upwind and downwind flight observed in both cumulative and session-by-session analysis of collections and wind direction. Since collection may not occur at the night during which dispersal occurred, only the first trapping session can be used to confirm upwind or downwind flight of individuals, and here we present evidence for upwind flight and downwind flight of individual Culicoides to distances greater than 1 km in a 24 h period. Whilst Kluiters, Swales & Baylis (2015) suggest upwind and downwind flight from the release point occurred in their study, the wind speed and direction used data were at too low a resolution (24 h average) to test this hypothesis given the likely influence of the local topographic complexity on wind direction (Whitman & Doran 1993). Here, all replicates took place under favourable conditions for Culicoides flight, with light winds (<3 m s⁻¹) (Fig. 7). Increasing wind speed increased the likelihood of capture at distance from the marked area, suggesting downwind transport may also play a role in local dispersal. Increased lunar radiation correlated with increased probability of detecting ovalbumin-positive Culicoides likely due to the increased Culicoides activity on moonlit nights (Linhares & Anderson 1990; Bishop et al. 2000).

Conclusion

The ovalbumin-immunomarking technique was demonstrated to be a highly effective tool for the study of Culicoides dispersal behaviour with evidence it may be suitable for other hematophagous vectors e.g. Culex and Phlebotominae. The use of the immunomarking technique in areas of increased topographical complexity, potentially involving multiple marker proteins to label different hosts or habitats within a single location (Hagler & Jones 2010), combined with investigations of gene flow between populations using landscape genetics (Manel & Holderegger 2013) and deep sequencing of viral genomes, could illustrate how Culicoides and the arboviruses they transmit move through the environment and quantify the relative impact of barriers to dispersal. Increased confidence in the estimates of within- and between-farm dispersal of Culicoides will enable improved data-driven modelling of the spread of Culicoides-borne arboviruses (Graesbøll et al. 2016) and will inform policy response to incursions and outbreaks at both the national and transnational level.

Authors’ contributions

C.J.S. and L.E.H. collected the data, S.C., L.T., V.B. and M.E. assisted in fieldwork, L.E.H. analysed the data; C.S. and L.E.H. led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

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Data accessibility

Data used in the production of this manuscript are available from the Dryad Digital Repository https://doi.org/10.5061/dryad.s403s (Sanders et al. 2017).

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Supporting Information
Details of electronic Supporting Information are provided below.

Table S1. Distance (m) of trap locations relative to the egg solution-marked area and number of Culicoides collected.

Table S2. Number of Culicoides collected positive for ovalbumin during immune-marking studies.

Table S3. Meteorological conditions recorded during field trials.

Table S4. Comparison of different threshold methods for analysis of enzyme-linked immunosorbent assay (ELISA) optical density results.

Fig. S1. The relative level of contamination of individuals within groups of naive Culicoides nubeculosus exposed to different numbers of ovalbumin-positive C. nubeculosus.