# No evidence for cumulative effects of small wind turbines on bat activity

*Jeroen Minderman1$, Mairi H. Gillis2, Helen F. Daly3 & Kirsty J. Park1*

1. Biological & Environmental Sciences, University of Stirling, Stirling, FK9 4LA, United Kingdom.
2. 228 Galashiels Road, Stow, Galashiels, Selkirkshire, TD1 2RA, [mgillis4a7i@gmail.com](mailto:mgillis4a7i@gmail.com)
3. 20/11 Duff Street, Dalry, Edinburgh, Midlothian, EH11 2HG, [hfdaly@outlook.com](mailto:hfdaly@outlook.com)

$***Corresponding author***. Current address: School of Biology, University of St Andrews, St Andrews, Fife, KY16 9ST, UK. Email: [jm340@st-andrews.ac.uk](mailto:jm340@st-andrews.ac.uk).

**Key words**: microturbines, domestic turbines, renewables, renewable energy, conservation, planning, wildlife impact

5365 words excl. appendix, 4642 excl. table/figure captions.

# Summary

1. While the effects of large wind farms on wildlife (particularly birds and bats) are generally well-studied, similar effects of small wind turbines (SWTs, in the UK usually up to 25m in hub height) remain relatively unknown. This is problematic, because rapid development and increasing accessibility of SWT technology have led to rapid growth of the number of SWTs installed worldwide.
2. Although recent studies showed that bat activity is negatively affected by the immediate proximity of single operating SWTs, the potential for wider landscape-scale effects of multiple SWTs installed in clusters remains unstudied. By contrast, such cumulative effects are commonly considered in impact studies of large wind farms.
3. We measured variation in bat activity at a sample of 34 SWT sites in the UK (free-standing SWTs between 6 and 25m in hub height), in order to test whether (1) accounting for variation in habitat, bat activity is generally lower in closer proximity of installed SWTs (e.g. 0-100m, compared to 100-200m, 200-300m, etc.), and (2) whether this effect is stronger in sites with multiple SWTs compared to sites with single turbines.
4. We focused our analysis on Pipistrelle bats because more than 98% of our recorded activity was from this genus. We show that although observed Pipistrelle activity was up to 10% lower within 100m of multiple turbine sites compared to activity at 100-200m, after accounting for confounding effects of habitat and environmental conditions, there is no effect of turbine proximity, irrespective of the number of turbines installed.
5. We conclude that within our sample there is no evidence of cumulative effects of SWTs on the activity of Pipistrelle bats on a landscape scale (i.e. over >100m distances). Our previous studies did show that single SWTs may adversely affect bat activity on smaller spatial scales (within 25m), so the combination of these findings suggest that any adverse effects of SWTs may be relatively localised, even if more than one SWT is present. It should be noted that this study focused on small wind turbines only, and that these findings should not be directly extrapolated to larger wind turbines.

# Introduction

Understanding and predicting how wildlife is affected by disturbance or habitat loss is becoming increasingly important as anthropogenic pressures on the natural world continue to grow (Tuomainen & Candolin 2011). This is particularly pertinent where incremental changes or disturbances could combine to produce cumulative effects. For example, whilst the rapid growth of the wind energy sector worldwide provides a vital contribution to sustainable energy production, a growing number of studies show that wind farms can have adverse effects on wildlife (Kuvlesky *et al.* 2007). Although the recognition of the potential importance of cumulative effects (defined as "*the additional changes caused by a proposed development in conjunction with other similar developments or as the combined effect of a set of developments, taken together*", SNH (2012)) is relatively recent and its quantification is in its infancy (Masden *et al.* 2010), both direct (collision mortality of birds and bats, e.g. Drewitt & Langston (2006), Rydell *et al*. (2010)) and indirect effects (disturbance or displacement, e.g. Larsen & Guillemette (2007), Pearce-Higgins *et al*. (2009)) of individual turbines or wind farms have been widely studied.

By contrast, similar effects of small wind turbines (SWTs, also referred to as micro-turbines or domestic turbines) on wildlife remain relatively unknown. These turbines are not only much smaller in size (in the UK, typical units are between 6 and 25m hub height) than those in wind farms, they are also commonly installed in a much wider range of habitats. Thus, their impacts on wildlife is likely to be different to that of large turbines, but up until recently this had not been quantified systematically. Our previous work found evidence for adverse effects of SWTs on bat activity (Minderman *et al.* 2012, Tatchley et al. submitted) as well as direct mortality (Minderman *et al.* 2014). These previous studies focused on a relatively limited spatial scale (e.g. within 25m of installed SWTs) and on single SWTs only. As a result, it is unclear whether any such adverse effects of SWTs on wildlife could be cumulative, i.e. whether the combined effects of multiple SWT installations could have greater adverse effects on wildlife.

This lack of understanding of the potential cumulative effects is particularly important for SWTs for two reasons. First, the number of SWTs installed is growing rapidly, both in the UK and worldwide and this growth is set to continue (WWEA 2013; RenewableUK 2014). As a result, the density of installed SWTs is likely to increase even if they are not installed in the same scheme or by the same owner. Secondly, the number of SWTs installed in groups is growing rapidly, with installations of 2-4 turbines now relatively common in the UK. Indeed, some installers now specifically promote the installation of multiple SWTs in so-called "wind crofts" **REF?**. As a result, where SWT installation requires planning permission (Park, Turner & Minderman 2013), decision makers are increasingly faced with the question whether multiple-turbine installations would have greater effects on wildlife, or if limiting the number of SWTs to be installed would be a feasible mitigation option. Currently, the evidence base for such decisions for SWTs specifically is entirely lacking.

Here, we aim to address this knowledge gap by quantifying and analysing the potential cumulative effect of SWTs on bat activity on a wider landscape scale (up to 500m from installed turbines). Specifically, using data collected at 34 throughout the UK we test the following predictions:

1. Bat activity is systematically lower in closer proximity (e.g. 0-100m from SWTs compared to 100-200m, 200-300m, etc.) of operating SWTs, controlling for the effects of habitat and environmental conditions.
2. The effect of SWT proximity on bat activity as tested in Prediction 1 is stronger in sites with multiple (2-4) SWTs installed compared to single SWT sites. Support for this prediction would indicate evidence of cumulative effects of SWTs.

# Methods

## Sites

Data were collected at 34 SWT sites in central and eastern Scotland. Sites were selected from an existing database of owners (Minderman *et al.* 2012; Park, Turner & Minderman 2013) based on how representative they were of common UK SWT installation settings in terms of habitat, turbine models and size. All turbines studied here were free-standing and between 6 and 25m (mean 16m) in hub height, and all were in rural settings but specific habitat in the surrounding area varied (see [Habitat data and variable selection](#habitat-data-and-variable-selection)). The number of SWTs installed in each site varied between 1 and 4 (median 1.6). SWTs installed in individual sites were the same size and specification with the exception of one site. More than two SWTs were installed in only 5 sites in our sample. We therefore limited the analyses presented here to a comparison of single and multiple turbine sites.

## Bat data and transects

Bat activity data were collected between 28 May and 01 September 2013 and 07 July and 04 September 2014. The time of data collection varied but started 30 minutes after sunset at the earliest and finished well before sunrise in all cases.

Bat activity was measured along transects by 1-2 observers walking the length of each transect at a slow (approximately 2.5km h-1) and constant pace, using EchoMeter EM3+ bat detectors (WildLife Acoustics, Mass., USA). A target of four transects was planned for all sites, running out from the turbine (or the central point between turbines in the case of multiple turbine sites) in four cardinal directions. However, because of physical constraints (e.g. walls, impassable fences or ditches, houses or buildings) the actual number of transects per site as well as their length varied (3-4, mean 3.7 per site, length: 300-500m). All transects were placed so that (1) the combination of all transects within each site covered all major habitats present, and (2) overall distance separating each transect was maximised. Transects were divided into 100m sections running out from the turbine centre point, giving up to five distance bands running away from the turbine.

One measure of ground level wind speed was taken at the end of each section (i.e. in each distance band) on each survey visit, and minimum daily temperature measures for each survey visit were obtained from the UK MIDAS weather station data at Grangemouth (N 56° 1' 5.15, W 3° 43' 5.88) (‘UK Meteorological Office’ 2006).

## Habitat data and variable selection

To account for expected confounding effects of habitat variation on bat activity along the transects, we used two key measures of habitat variability in each transect section: (1) edge density and (2) proportion of woodland. These two variables were selected on the basis of a preliminary analysis of the effect of a full set of 10 habitat variables on bat activity. This was done to avoid both over-parameterisation of the main statistical models presented here, as well as the inclusion of highly collinear habitat metrics. Full details of this preliminary analysis are given in Appendix 1. To obtain habitat data per transect section, 50m buffers were placed around digital maps of each transect route, resulting in approximately 100m x 100m transect sections. The exact area of each section varied because of non-linear transect sections, but this was accounted for in the analysis, see [Statistical analysis](#statistical-analysis). All habitat variables were quantified in each transect section using 1:1250 UK Ordnance Survey MasterMap Topography digital maps, using QGIS v. 1.8 (‘Quantum GIS Development Team’ 2014). Mean distance (m) to both buildings and water was calculated by constructing a raster map of distances between each raster cell and the nearest cell with buildings or water map data, and averaging these raster values across each transect section. Edge density (m m-2) represented the density of "edge" habitat in each transect cell, and was calculated as the total length of all linear habitat features ("line" data in the OS Topography Layer) divided by the area of the transect section. Thus, this is a description of the density of e.g. building-, woodland and water edges, hedgerows, roads and tracks, roadsides, field boundaries. Finally, the proportion of tree coverage in each transect section was the sum of all tree coverage (m2) (coniferous, non-coniferous and unclassified trees) in the OS Topography polygon data divided by the transect section area. All Pearson correlation coefficients between these four habitat measures were <0.2 with the exception of the correlation between edge density and distance to buildings which was 0.37.

## Data analysis and statistics

### Bat activity: probability of a pass per hectare surveyed

Bat activity was initially quantified as the number of bat 'passes' (defined as a sequence of at least two echolocation calls separated by less than a second) per transect section. However, we chose to analyse our data as bat activity presence or absence per transect section, per survey visit, for two reasons. First, the distribution of observed counts was highly skewed (many zeros and excessive variation) so that count-based statistical models did not provide any reasonable fit. Second, using bat 'passes' as a measure of activity provides a relative measure of activity and analyses of absolute pass count would therefore add little information. In addition, because the area covered by each transect section varied slightly (see [Bat data and transects](#bat-data-and-transects)), we here model the probability of detecting bat activity per section and hectares covered.

### Statistical analysis

We used generalised linear mixed effects models (GLMMs) (Gelman & Hill 2007) and an Information-Theoretic model selection (IT) approach (Burnham & Anderson 1998) to analyse our data.

#### Model structure

We modelled the probability of a bat pass per unit area on a given survey visit as the response variable with a GLMM with binomial errors and a complementary log-log link function. This link function allowed us to include transect section area as an offset in the model (thus accounting for slight variation in the size of each transect section). To account for the non-independence of repeated measurements from the same site and transect sections, all models included transect nested within site as a random effect. To test our two predictions (see [Introduction](#introduction)), we included two focal fixed factors; (1) transect section (distance bands; 0-100m, 100-200m, 200-300m, 300-400m and 400-500m from the SWTs) and (2) turbine number (single or multiple SWTs), as well as the statistical interaction between the two. In addition to these focal factors we included the two selected (see [Habitat data and variable selection](#habitat-data-and-variable-selection)) habitat variables (distance to buildings, distance to water, edge density and proportion of tree cover) as continuous covariates. Moreover, because bat activity is known to vary with weather conditions (particularly temperature and wind speed) across the season and through the night, we also included covariates for the minimum temperature over 24h (C), wind speed (m s-1), Julian day number, time to midnight (mins.) and time to midnight2 (to account for potential non-linear effects of the latter). Thus, accounting for any confounding effects of habitat and environmental conditions, a transect section effect would indicate a systematic difference in bat activity as a function of distance from SWTs (Prediction 1), and an interaction between this and turbine number would indicate that the strength of this effect depends on the number of SWTs installed (Prediction 2). To avoid over-parameterisation of the model, interactions between confounding effects were not considered. All inputs were standardised (centred to 0 and scaled to 2 SD) following Gelman (2008) to improve performance of parameter estimation and allow for direct interpretation of relative effect strength, but we calculate and present model predictions on the response scale. Summary statistics of unstandardised model inputs are given in Table 1.

#### Model selection, averaging and predictions

The full model set was constrained to (1) only include the time to midnight2 term when its linear term was also included, (2) include all random effects in all models, and (3) use the same link function and distribution for all models, yielding a N = 480 models. Model fit was assessed using the adjusted Akaike Information Criterion (AICc) and we took the top models within 4 AICc points of the 'top' model as our candidate set. To account for model selection uncertainty within the candidate set, we interpret the averaged parameter estimates and their standard errors calculated using the 'zero method' (Burnham & Anderson 1998). Prediction intervals were calculated as the 95% quantiles (weighted by model weight) of predictions from N = 1000 simulated draws from the estimated parameter distributions for each of the models in the candidate set (Gelman & Hill 2007). For each model in the candidate set we also present the marginal and conditional R2 (Nakagawa & Schielzeth 2013).

We used R version 3.1.3 (2015-03-09) (‘R Development Core Team’ 2015) for all statistical analyses. GLMMs were fit using package *lme4* v. 1.1-7 (Bates *et al.* 2014). Model selection and averaging was performed using package *MuMIn* v. 1.13.4 (Barton 2015), and standardisation of model inputs and parameter simulation used package *arm* v. 1.7-07 (Gelman & Su 2014).

# Results

Over the two years of the study, we collected bat activity data in N = 1395 transect sections, during 78 survey visits. Most sites (N = 30) were surveyed on at least two occasions (2-6 surveys), but four sites were only visited once. Within this sample, N = 20 were single-turbine sites, and N = 14 were multiple-turbine sites (2-4 turbines). In total, we recorded N = 1867 bat passes, of which 98.4% (N = 1838) were Pipistrelle (*Pipistrellus pipistrellus* or *P. pygmaeus*) bats. Thus, here we only present analyses of Pipistrelle bat activity combined. Overall, Pipistrelle bat activity was detected on 466 section surveys (observed average probability of a Pipistrelle pass = 0.33).

The candidate set (AICc<4) of GLMMs for the probability of a bat pass per hectare contained 10 models. The top model retained 7 out of the 10 predictors in the full model, including distance band but neither the number of turbines nor an interaction between distance band and number of turbines. This model was = 0.244/0.155 = 1.574 (evidence ratio) times more strongly supported than second-best model, and = 0.244/0.033 = 7.394 times better supported than the saturated model (ranked 10th). The null model was ranked 446th and had no support ( < 0.001) (**Table 2**).

Although it was retained in the top model, four out of ten models in the candidate set did not retain an effect of distance band (predictor weight = 0.66). The relative importance of both the number of turbines and its interaction with distance band was even lower ( = 0.42 and = 0.1, retained in 5 and 2 models in the candidate set respectively). With the exception of minimum temperature ( = 0.32) all other predictors were retained in all models in the candidate set (Table 3).

Thus, although at multiple turbine sites the observed probability of a Pipistrelle bat pass appeared to be lower in the closest (0-100m) distance band compared to the 100-200m distance band, this difference was relatively small (0.29 vs. 0.37) (Figure 1, dark bars, and Table 3). Moreover, this effect is no longer apparent when considering model predictions that account for the effects of habitat- and environmental conditions. For example, in multiple turbine sites, predicted bat activity ranged from 0.2 to 0.34 in the nearest distance band (0-100m), and from 0.24 to 0.39 in the 100-200m distance band. Similar overlapping prediction intervals apply for both single turbine sites as well as the further distance bands (Figure 1, points and error bars).

# Discussion

We show that, when accounting for confounding effects of habitat and environmental conditions, Pipistrelle bat activity is not systematically lower within 100m of SWTs compared to longer distances (100-200m, 200-300m, 300-400m and 400-500m), irrespective of the number of turbines (single or multiple) installed. On this basis, we conclude that there is no evidence for cumulative effects of SWTs on a landscape (i.e. >100m) scale.

By contrast, we previously showed that bat activity can be systematically lower in the immediate vicinity (<25m) of operating SWTs (Minderman *et al.* 2012), and indeed subsequent experimental studies of bat activity before- and after installation of SWTs near hedgerows confirmed such adverse effects on activity (Tatchley *et al*. submitted). Combined with the present finding of a lack of similar effects on a larger scale (i.e. within 100m), this suggests that such adverse effects are relatively localised. Although the mechanism causing such avoidance behaviour is unclear, experimental studies in a laboratory setting have shown that the return of echolocation pulses off spinning turbine blades may be erratic, potentially causing navigational problems for bats (Long, Flint & Lepper 2010). Other experimental studies have shown that some species of bat may avoid foraging in noisy environments (Schaub, Ostwald & Siemers 2008) but whether noise generated by turbines (including SWTs) has the same effect has not been tested. In either case, it would be reasonable to assume that the strength of such interference would dissipate relatively rapidly with distance, especially if it involves relatively high-frequency noise (which should attenuate more strongly over shorter distances;Russ 2012), limiting the spatial scale of any adverse effect of the turbine.

This apparently relatively localised effect of SWTs may in turn also explain the lack of evidence for any cumulative effect (i.e. effects were not different between single- or multiple turbine sites). If avoidance behaviour only operates over relatively short (as per our previous findings, <25m) distances, it may be possible for bats to avoid individual turbines rather than whole "clusters" of turbines. Indeed, in our multiple turbine sites, turbines were sited 13 - 122 m apart with this distance being less than 23 m in only one case (mean 49.4m). This suggests that in most cases bats would be able to avoid individual turbines without coming in very close proximity of another, limiting the possibility of any cumulative effects. Further detailed study of individual flight paths would be required to confirm whether this is the case, however. Moreover, it should be noted that although the possibility of cumulative effects is often raised for larger turbines and wind farms (Masden *et al.* 2010; SNH 2012), empirical studies that unequivocally demonstrate such effects are rather rare and have tended to focus on birds rather than bats (Schaub 2012). Although this may be due to the logistical difficulty of study such large-scale effects, it is possible that cumulative effects themselves are rare, except in specific circumstances.

There are a number of important caveats with our findings that should be considered. First of all, these findings and conclusions should not be generalised to species other than *Pipistrellus* sp. Although Pipistrelle bats are by far the commonest species in both our wider study area as well as in the specific habitats we studied (Middleton 2006), it is likely that the collection of sufficient data to draw species-specific conclusions would require targeted surveys. The need to collect data at a wide range of sites rather than in specific habitats (where less widespread species may be more abundant) prevented us from doing this here. However, larger turbines in wind farms are known to have different effects on different species (Cryan & Barclay 2009) and we therefore consider this to be a research priority for SWTs (Tatchley *et al.* submitted). Secondly, the limited number of sites with more than two turbines in our sample means that we cannot exclude the possibility that a larger number of turbines *would* cause cumulative effects; indeed modelling studies in birds (Schaub 2012) show that this may well be the case. Finally, the combination of recording bat activity over a relatively large scale (up to 500m from SWTs) along transects, and a distance-based approach to turbine effects, required that we analysed our data in distance bands. Our choice of relatively 'coarse' 100m distance bands may mean that we have missed more subtle cumulative effects of SWTs. However, our previous study had already established adverse effects on activity seem to approach zero over 25 m (Minderman *et al.* 2012). More importantly, because of the mobility of the transect-based approach it would become very difficult to accurately allocate individual bat passes to more narrow distance bands, i.e. this would potentially introduce false precision. Data on behavioural responses to turbines on a finer scale would require alternative survey methods, e.g. flight path tracking (Holderied & Helversen 2003).

In summary, based on the data presented here it is unlikely that SWTs in the range of 6-25m hub height have adverse effects on bat activity on a landscape (>100 m) scale, or that such effects of cumulative (when considering 2-4 installed turbines). Given the rapid expansion of SWTs both in the UK and worldwide, and the fact that in many cases planning- or siting permission is required, these findings are of key importance to both decision makers on the local level as well as to renewable energy policy makers.

# Acknowledgements

We are extremely grateful to all site- and turbine owners who kindly allowed us access to their property for the survey work; this study would have been impossible without their help. Kathryn Hamilton made substantial contributions to the data collection in 2013, and Sofia Motta Pralon assisted with GIS data entry - thank you both. Many thanks to Cerian Tatchley, Paul Lintott, Nils Bunnefeld, Chris Pendlebury and Claudia Garratt for useful discussions. This study was funded by a University of Stirling Impact Fellowship to JM.

# References

Barton, K. (2015) MuMIn: Multi-Model Inference. R package version 1.13.4.

Bates, D., Maechler, M., Bolker, B. & Walker, S. (2014) Lme4: Linear mixed-effects models using Eigen and S4. R package version 1.1-7.

Burnham, K. & Anderson, D. (1998) *Model Selection and Multimodel Inference: A Practical Information-Theoretic Approach*, 2nd ed. Springer Verlag, New York, USA.

Cryan, P. & Barclay, R. (2009) Causes of bat fatalities at wind turbines: Hypotheses and predictions. *Journal of Mammology*, **90**, 1330–1340.

Drewitt, A. & Langston, R. (2006) Assessing the impacts of wind farms on birds. *Ibis*, **148**, 29–42.

Gelman, A. (2008) Scaling regression inputs by dividing by two standard deviations. *Statistics in Medicine*, **27**, 2865–2873.

Gelman, A. & Hill, J. (2007) *Data Analysis Using Regression and Multilevel/Hierarchical Models*. Cambridge University Press, New York.

Gelman, A. & Su, Y.-S. (2014) Arm: Data Analysis Using Regression and Multilevel/Hierarchical Models. R package version 1.7-07.

Holderied, M.W. & Helversen, O. von. (2003) Echolocation range and wingbeat period match in aerial-hawking bats. *Proceedings of the Royal Society B: Biological Sciences*, **270**, 2293–2299.

Kuvlesky, W.P., Brennan, L.A., Morrison, M.L., Boydston, K.K., Ballard, B.M. & Bryant, F.C. (2007) Wind Energy Development and Wildlife Conservation: Challenges and Opportunities. *Journal of Wildlife Management*, **71**, 2487–2498.

Larsen, J. & Guillemette, M. (2007) Effects of wind turbines on flight behaviour of wintering common eiders: Implications for habitat use and collision risk. *Journal of Applied Ecology*, **44**, 516–522.

Long, C., Flint, J. & Lepper, P. (2010) Wind turbines and bat mortality: Doppler shift profiles and ultrasonic bat-like pulse reflection from moving turbine blades. *JOURNAL OF THE ACOUSTICAL SOCIETY OF AMERICA*, **128**, 2238–2245.

Masden, E.A., Fox, A.D., Furness, R.W., Bullman, R. & Haydon, D.T. (2010) Cumulative impact assessments and bird/wind farm interactions: Developing a conceptual framework. *Environmental Impact Assessment Review*, **30**, 1–7.

Middleton, N. (2006) *An Introduction to the Bats of Scotland*. ECHOES Ecology Ltd., Polmont, UK.

Minderman, J., Fuentes-Montemayor, E., Pearce-Higgins, J.W., Pendlebury, C.J. & Park, K.J. (2014) Estimates and correlates of bird and bat mortality at small wind turbine sites. *Biodiversity and Conservation*, **24**, 467–482.

Minderman, J., Pendlebury, C.J., Pearce-Higgins, J.W. & Park, K.J. (2012) Experimental evidence for the effect of small wind turbine proximity and operation on bird and bat activity. *PLoS ONE*, **7**, e41177.

Nakagawa, S. & Schielzeth, H. (2013) A general and simple method for obtaining R2 from generalized linear mixed-effects models. *Methods in Ecology and Evolution*, **4**, 133–142.

Park, K.J., Turner, A. & Minderman, J. (2013) Integrating applied ecology and planning policy: The case of micro-turbines and wildlife conservation. *Journal of Applied Ecology*, **50**, 199–204.

Pearce-Higgins, J.W., Stephen, L., Langston, R.H.W., Bainbridge, I.P. & Bullman, R. (2009) The distribution of breeding birds around upland wind farms. *Journal of Applied Ecology*, **46**, 1323–1331.

‘Quantum GIS Development Team’. (2014) *Quantum GIS Geographic Information System*. Open Source Geospatial Foundation Project.

‘R Development Core Team’. (2015) *R: A Language and Environment for Statistical Computing*. R Foundation for Statistical Computing, Vienna, Austria.

RenewableUK. (2014) *Small and Medium Wind Strategy: The Current and Future Potential of the Sub-500kW Wind Industry in the UK*. RenewableUK, London, UK.

Russ, J. (2012) *British Bat Calls: A Guide to Species Identification*. Pelagic Pub., Exeter.

Rydell, J., Bach, L., Dubourg-Savage, M., Green, M., Rodrigues, L. & Hedenstrom, A. (2010) Bat mortality at wind turbines in northwestern Europe. *Acta Chiropterologica*, **12**, 261–274.

Schaub, M. (2012) Spatial distribution of wind turbines is crucial for the survival of red kite populations. *Biological Conservation*, **155**, 111–118.

Schaub, A., Ostwald, J. & Siemers, B. (2008) Foraging bats avoid noise. *JOURNAL OF EXPERIMENTAL BIOLOGY*, **211**, 3174–3180.

SNH. (2012) *Assessing the Cumulative Impact of Onshore Wind Energy Developments*. Scottish Natural Heritage, Inverness, UK.

Tuomainen, U. & Candolin, U. (2011) Behavioural responses to human-induced environmental change. *Biological Reviews*, **86**, 640–657.

‘UK Meteorological Office’. (2006) *MIDAS Land Surface Stations Data (1853-Current)*. NCAS British Atmospheric Data Centre, Didcot, UK.

WWEA. (2013) *2013 Small Wind World Report Update*. World Wind Energy Association, Bonn, Germany.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | Mean | Median | Std. dev | Min. | Max. |
| Minimum 24h temperature (°C) | 11.76 | 12.00 | 2.75 | 5.00 | 17.6 |
| Julian day number | 196.59 | 195.00 | 28.85 | 147.00 | 246 |
| Time to midnight (min) | -26.07 | -41.00 | 82.54 | -202.30 | 197.77 |
| Time to midnight2 | 7487.47 | 4669.44 | 8375.13 | 00 | 40925.29 |
| Wind speed (m s-1) | 1.07 | 0.40 | 1.46 | 00 | 9.8 |
| Edge density (m m-2) | 0.04 | 0.03 | 0.02 | 00 | 0.13 |
| Proportion woodland | 0.05 | 0.00 | 0.13 | 00 | 0.92 |

**Table 1.** Descriptive statistics of predictor variables in the Generalised Linear Mixed Model (GLMM) for bat activity (Table 2), on their original (unstandardised) scale.

**Table 2.** Candidate set of Generalised Linear Mixed Models for bat activity (probability of bat activity ha-1, binomial errors, complementary log-log link) within 4 AICc points of the “top” model. Coefficients are parameter estimates based on models fit using standardised (centered and scaled to 2 SD) inputs. In the case of factorial predictors, their inclusion is indicated by a +. Empty cells indicate the predictor was not included in a given model. K is the number of parameters in the models. All models included site and transect within site as a nested random effect structure.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Model no. | Intercept | No. of turbines | Distance band | Day number | Edge density | Min. 24h temperature | Proportion woodland | Time to midnight | Time to midnight2 | Wind speed | No. of turbines \* Distance band | k | Log-likelihood | AICc | Δ AICc | Model weight | *R*2 (marginal) | *R*2 (conditional) |
| 1 | -1.319 |  | + | 0.491 | 0.49 |  | 0.35 | -0.20 | -0.561 | -0.694 |  | 13 | -797.3 | 1620.9 | 0 | 0.244 | 0.204 | 0.391 |
| 2 | -1.048 |  |  | 0.504 | 0.507 |  | 0.416 | -0.193 | -0.581 | -0.706 |  | 9 | -801.8 | 1621.8 | 0.905 | 0.155 | 0.201 | 0.389 |
| 3 | -1.320 | 0.182 | + | 0.503 | 0.493 |  | 0.355 | -0.190 | -0.549 | -0.689 |  | 14 | -796.9 | 1622.1 | 1.217 | 0.133 | 0.213 | 0.394 |
| 4 | -1.335 |  | + | 0.506 | 0.493 | -0.080 | 0.349 | -0.201 | -0.559 | -0.702 |  | 14 | -797.1 | 1622.5 | 1.573 | 0.111 | 0.209 | 0.391 |
| 5 | -1.037 | 0.174 |  | 0.512 | 0.505 |  | 0.421 | -0.183 | -0.568 | -0.697 |  | 10 | -801.4 | 1623.1 | 2.203 | 0.081 | 0.207 | 0.391 |
| 6 | -1.305 | -0.075 | + | 0.506 | 0.544 |  | 0.333 | -0.19 | -0.565 | -0.683 | + | 18 | -793.4 | 1623.3 | 2.456 | 0.072 | 0.222 | 0.402 |
| 7 | -1.053 |  |  | 0.517 | 0.505 | -0.078 | 0.415 | -0.193 | -0.577 | -0.709 |  | 10 | -801.6 | 1623.4 | 2.495 | 0.07 | 0.204 | 0.387 |
| 8 | -1.325 | 0.189 | + | 0.516 | 0.491 | -0.086 | 0.354 | -0.189 | -0.545 | -0.692 |  | 15 | -796.6 | 1623.6 | 2.722 | 0.063 | 0.216 | 0.392 |
| 9 | -1.042 | 0.180 |  | 0.526 | 0.503 | -0.084 | 0.420 | -0.182 | -0.564 | -0.700 |  | 11 | -801.2 | 1624.6 | 3.727 | 0.038 | 0.210 | 0.388 |
| 10 | -1.310 | -0.069 | + | 0.520 | 0.542 | -0.084 | 0.332 | -0.188 | -0.561 | -0.686 | + | 19 | -793.1 | 1624.9 | 3.996 | 0.033 | 0.225 | 0.400 |
|  | -1.121 |  |  |  |  |  |  |  |  |  |  | 3 | -853.0 | 1712.1 | 91.183 | <0.001 | <0.001 | 0.274 |

**Table 3.** Model-averaged (using the zero-method, Burnham & Anderson 1998) parameter estimates (*b*) and their standard errors based on the candidate set of Generalised Linear Mixed Effects Models (GLMMs) for bat activity (probability of a bat pass ha-1) shown in Table 2. Coefficients are parameter estimates based on model fits using standardised (centered and scaled to 2 SD) inputs. The reference categories for distance band is the first band (0-100m from turbines) and the standardised inputs for no. of turbines (two-way factor; therefore only centered – see main text and Gelman (2008)) were -0.538 and 0.462 for single and multiple turbines respectively. All models included site and transect within site as a nested random effect structure, and all models had binomial errors and were fitted with a complimentary log-log link.

|  |  |  |
| --- | --- | --- |
| Parameter | *b* | S.E. |
| Intercept | -1.226 | 0.180 |
| Distance band 2 (100-200m) | 0.171 | 0.161 |
| Distance band 3 (200-300m) | 0.277 | 0.225 |
| Distance band 4 (300-400m) | 0.215 | 0.187 |
| Distance band 5 (400-500m) | 0.258 | 0.215 |
| Julian day no. | 0.505 | 0.129 |
| Edge density | 0.502 | 0.096 |
| Proportion woodland | 0.372 | 0.093 |
| Time to midnight | -0.193 | 0.109 |
| Time to midnight2 | -0.563 | 0.200 |
| Wind speed | -0.696 | 0.124 |
| No. turbines | 0.049 | 0.174 |
| Min. 24h temperature | -0.026 | 0.076 |
| Distance band 2 (100-200m) \* No. turbines | 0.049 | 0.177 |
| Distance band 3 (200-300m) \* No. turbines | 0.046 | 0.169 |
| Distance band 4 (300-400m) \* No. turbines | -0.016 | 0.113 |
| Distance band 5 (400-500m) \* No. turbines | 0.049 | 0.178 |

C:\Jeroen\docs\000_Turbines\000_Txt\Bats_LargeScale_MS\FIG1.tiff

**Figure 1.** Observed (bars) and predicted (points and lines) bat activity (probability of a bat pass ha-1) based on Generalised Linear Mixed Effects Models (GLMMs, Tables 2 and 3), in each of five distance bands around small wind turbines. White bars and grey circles are for single-turbine sites, and grey bars and black circles are for multiple-turbine sites. The error bars (lines) represent the predictive uncertainty, calculated as the 95% quantiles of predicted distributions based on N = 1000 samples from the estimated parameter distributions (Gelman & Hill 2007).

# *Supplementary material:*

## Appendix 1: Preliminary habitat data analysis

A total of 10 habitat variables were quantified in each transect section in all sites (Table S1), using 1:1250 UK Ordnance Survey MasterMap Topography digital map (OS) data and QGIS v. 1.8. These variables were chosen on the basis of their known potential effect on bat activity. The five proportion land cover variables (proportion buildings, road or tracks, rough grassland, roadside, and woodland) were calculated from the OS polygon area data and were calculated as the sum of each of these five land cover types in each transect section, divided by the size of the transect section. These were defined as the total area size of all polygons listed as (1) "Buildings" in the *Theme* field; (2) "Road Tracks and Paths"" in the *Theme* field; (3) "Rough grassland" (including all lower classifications) in the *descrTerm* field; (4) "Roadside" in the *descrGroup* field and (5) "Coniferous or Nonconiferous trees"" in the *descrGroup* field. The four distance variables (distance to buildings, linear features, trees and water) were calculated by overlaying the OS data with a 1x1m 'raster' of grid cells, calculating the distance (m) between each raster cell and the nearest raster cell containing each of the four land cover types (respectively, buildings as defined above, any linear feature as contained in the OS "line" data, any woodland as defined above and any water polygon), and averaging these distances for all cells within the transect sections. Finally, edge density (m m-2) was expressed as the total length of all line features contained in the OS Linear data per transect section, divided by the transect section area.

To identify a subset of habitat variables that were most descriptive of bat activity (see main text), we ran an initial model selection (Burnham & Anderson 1998) procedure on a Generalised Linear Mixed Effects (GLMM) (Gelman & Hill 2007) model for the probability of a bat pass per hectare as a function of all 10 habitat variables, transect within site as a random effect, a binomial error distribution with a log-log link function and transect section size (ha) as an offset. This model structure was identical to the model presented in the main text. To avoid problems with colinearity, we limited the full model set to include only one of the five 'proportion' habitat variables (because these represent proportion cover, lower cover of one automatically implies greater cover of another, leading to high colinearity). We further excluded all models including (1) both proportion of woodland and distance to trees, (2) both proportion buildings and distance to buildings, as well as models including edge density and either (3) distance to linear features or (4) proportion of roads or tracks. Again, these latter exclusions were to avoid high colinearity: a greater proportion of woodland in a given transect section naturally implies shorter distances to trees, edge density is comprised of linear features including roads and tracks, etc. Thus, these variables effectively measured very similar things but in slightly different ways. Although this model selection procedure was done to identify which of these competing variables best describe variation in bat activity, they should not be included in the same models being compared. All model fitting procedures and analyses followed the methodology described in the main text.

The resulting full model set comprised of a total of 112 models (Table S2). The candidate set (AICc<4) contained 4 models. The 'top' model retained three of the ten habitat variables: distance to water, edge density and the proportion of woodland. Distance to water was only included in two out of the four models in the top set, however. By contrast, edge density and the proportion of woodland were retained in all four models in the candidate set. Thus, we chose to use edge density and the proportion of woodland as the key two habitat variables explaining bat activity.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | Mean | Median | SD | Min. | Max. |
| Distance to buildings (m) | 200.47 | 184.14 | 128.82 | 9.20 | 623.77 |
| Distance to linear features (m) | 22.74 | 18.33 | 19.24 | 5.37 | 190.97 |
| Distance to trees (m) | 180.72 | 154.98 | 137.68 | 0.19 | 646.47 |
| Distance to water (m) | 201.16 | 168.36 | 154.68 | 13.20 | 689.27 |
| Edge density (m m-2) | 0.04 | 0.03 | 0.02 | 0.00 | 0.13 |
| Proportion buildings | 0.01 | 0.00 | 0.03 | 0.00 | 0.29 |
| Proportion roads or tracks | 0.02 | 0.01 | 0.03 | 0.00 | 0.26 |
| Proportion rough grassland | 0.04 | 0.00 | 0.13 | 0.00 | 1.00 |
| Proportion roadsides | 0.02 | 0.00 | 0.03 | 0.00 | 0.26 |
| Proportion woodland | 0.05 | 0.00 | 0.13 | 0.00 | 0.92 |

**Table S1.** Descriptive statistics of all ten original habitat variables as predictors for bat activity on their original (unstandardised) scale.

**Table S2**. Candidate set of Generalised Linear Mixed Models for bat activity (probability of bat activity ha-1, binomial errors, complementary log-log link) within 4 AICc points of the “top” model, ***as a function of ten candidate habitat predictors only***. Coefficients are parameter estimates based on models fit using standardised (centered and scaled to 2 SD) inputs. K is the number of parameters in the models. All models included site and transect within site as a nested random effect structure.

|  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Model | Intercept | Distance to buildings | Distance to water | Edge density | Proportion woodland | K | Log-likelihood | AICc | Δ AICc | Model weight |
| 1 | -1.146 |  | -0.305 | 0.523 | 0.41 | 6 | -824.8 | 1661.7 | 0.000 | 0.541 |
| 2 | -1.146 | 0.028 | -0.301 | 0.532 | 0.408 | 7 | -824.8 | 1663.7 | 1.962 | 0.203 |
| 3 | -1.139 |  |  | 0.549 | 0.445 | 5 | -826.9 | 1663.9 | 2.184 | 0.181 |
| 4 | -1.138 | 0.059 |  | 0.568 | 0.438 | 6 | -826.8 | 1665.7 | 3.957 | 0.075 |