NERC Open Research Archive



Article (refereed) - postprint

Newton, Ian; McGrady, Michael J.; Oli, Madan K. 2016. A review of survival estimates for raptors and owls. *Ibis*, 158 (2). 227-248. <u>10.1111/ibi.12355</u>

© 2016 British Ornithologists' Union

This version available http://nora.nerc.ac.uk/513506/

NERC has developed NORA to enable users to access research outputs wholly or partially funded by NERC. Copyright and other rights for material on this site are retained by the rights owners. Users should read the terms and conditions of use of this material at http://nora.nerc.ac.uk/policies.html#access

This document is the author's final manuscript version of the journal article, incorporating any revisions agreed during the peer review process. There may be differences between this and the publisher's version. You are advised to consult the publisher's version if you wish to cite from this article.

The definitive version is available at http://onlinelibrary.wiley.com/

Contact CEH NORA team at <u>noraceh@ceh.ac.uk</u>

The NERC and CEH trademarks and logos ('the Trademarks') are registered trademarks of NERC in the UK and other countries, and may not be used without the prior written consent of the Trademark owner.

Received Date : 28-Apr-2014 Revised Date : 12-Jan-2016 Accepted Date : 02-Feb-2016 Article type : Review Editor : Beatriz Arroyo

Running head: Raptor survival estimates

A review of survival estimates for raptors and owls

IAN NEWTON¹, MICHAEL J. MCGRADY² & MADAN K. OLI^{3*}

¹Centre for Ecology and Hydrology, Benson Lane, Crowmarsh Gifford, Wallingford OX10 8BB, UK

²International Avian Research, Am Rosenhügel, 59 A-3500 Krems, Austria ³Department of Wildlife Ecology and Conservation, University of Florida, Gainesville, FL 32611, USA

*Corresponding author. Email: olim@ufl.edu

This paper reviews the literature on survival estimates for different species of raptors and owls, examines the methods used to obtain the estimates, and draws out some general patterns arising. Estimating survival usually involves the marking of birds so that they can be recognised as individuals on subsequent encounters. Annual survival can then be estimated from: (1) birds ringed at known age (usually as nestlings) and subsequently reported by members of the public (usually as found dead), the ratio of recoveries at different ages being used to calculate annual survival; (2) marked breeding adults, trapped or re-sighted in subsequent years in particular study areas, with the proportion re-trapped (or re-sighted) in each year being taken as the minimum annual survival; (3) the same, but with the application of capture-mark-recapture (or re-sighting) methods to estimate annual survival; (4) a combination of reports of known-age dead birds and re-trapping/re-sighting of live birds, (5) use of radio- or satellite-tracking to follow the fates of individuals; and (6) the integration of these methods with other information, such as change in numbers between years, to derive estimates of survival and other demographic parameters. Studies confined to particular areas This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: 10.1111/ibi.12355 This article is protected by copyright. All rights reserved.

usually give estimates of 'apparent annual survival', because they take no account of birds that leave the area. However, radio- or satellite-tracking makes it possible to estimate true survival, including for pre-breeders that have low natal-site fidelity (this usually requires satellite telemetry). As in other birds, the preferred method for estimating survival has changed over time, as new and more robust methods of estimation have been developed. Methods 1-2 were the first be developed, but without statistical under-pinning, while 3-6 were developed later on the basis of formal statistical models. This difference has to be borne in mind in comparing older with newer estimates for particular species. Published survival estimates were found for three species of Cathartidae, one of Pandionidae, 29 Accipitridae, 12 Falconidae, one Tytonidae and nine Strigidae, almost all from northern hemisphere, temperate species. In most of these species more than one estimate was available, and in some separate estimates for different age or sex groups. The main patterns to emerge included: (1) a significant tendency for annual adult survival to increase with body weight, smaller species having annual survival rates mainly of 60-70%, medium sized species mainly in the range 70-90% and the largest of more than 90%, in the absence of obvious humancaused losses; (2) a lower survival in the first or pre-breeding years of life than in subsequent years; (3) a lack of obvious or consistent differences in survival between the sexes, where these could be distinguished; and (4) in the few species for which enough data were available, a decline in annual survival rates in the later years of life.

Keywords: Birds of prey, bird ringing, capture-mark-recapture, demography, mortality, radio-tracking, survival estimation methods

The study of raptors has contributed substantially to our knowledge of basic and applied ecology, knowledge that can be transferred to many other species of birds and mammals (Newton 1979). As top predators, raptors can perform an important ecological role in limiting some prey populations, and their own numbers are in turn often influenced by the numbers of their prey (Newton 1979, 2013). Additionally, raptors require a variety of habitats for breeding and hunting, and serve as important biological indicators of the state of ecosystems (Sergio *et al.* 2005, 2006, 2008), for example with respect to chemical contamination (Ratcliffe 1970, 1993, Newton 1974, 1986, Newton & Wyllie 1992). Although some of the best examples of successful single species conservation programmes involve raptors (Cade *et al.* 1988, Newton & Wyllie 1992, Bretagnolle *et al.* 2008, Sulawa *et al.* 2010), the ecological processes influencing survival and other demographic rates for most species remain poorly understood.

Compared to many other birds, raptors generally occur at low densities, so sample-size problems are often difficult to avoid (Newton 1979). Furthermore, many species place their nests in sites that are difficult to access, which in turn makes adults hard to catch; consequently, marking of both nestlings and breeding adults in sufficient numbers requires a substantial investment of time, money and effort.

The growth rate of many raptor populations is proportionately more sensitive to adult survival rates than it is to reproductive parameters (Mertz 1971, Stahl & Oli 2006, Sergio et al. 2011). This means that identifying factors and processes that affect age- or life-stage specific survival rates is important for understanding raptor population dynamics. Typically, estimation of survival requires marking animals so that they can be individually identified and their fates monitored over time (Lebreton et al. 1992, Williams et al. 2002). Whatever method is used for marking, assumptions are that the marking method itself does not influence the survival of the individuals concerned, and that these individuals are representative of their population. For raptors, marking methods have included metal or coloured plastic rings, wing-tags, radio-tags and Passive Integrated Transponder (PIT) tags; while analytical methods have included models based on ring recoveries, capture-markrecapture (or re-sight), age-composition (e.g. of museum specimens or samples of dead birds), combinations of live-recaptures and dead-recoveries (Brownie et al. 1985, Pollock et al. 1989, Lebreton et al. 1992, Burnham 1993, Conn et al. 2004, Skalski et al. 2005) and, more recently, integrated population models (Besbeas et al. 2002, Schaub & Abadi 2011). Monitoring and statistical methods to estimate survival from various data types have evolved over time, potentially affecting the accuracy and precision of survival estimates (Clobert & Lebreton 1991, Williams et al. 2002, Craig et al. 2004, Nasution et al. 2004).

We review the literature on survival estimates for different species of raptors and owls, discuss the methods used to obtain the estimates, and identify some general patterns arising. Study species were drawn from the families Cathartidae, Pandionidae, Accipitridae, Falconidae, Tytonidae and Strigidae, and sources searched for information included papers, technical reports, books and book chapters published over seven decades (1946-2015) that gave survival estimates of raptors and owls.

METHODS

Methods used to find relevant literature

To locate relevant publications, we searched Web of Science

(http://apps.webofknowledge.com/) and Google Scholar (https://scholar.google.com/), using a combination of key words (survival, demography, population ecology, population dynamics, owl, raptor), and common and scientific names of different species. We also searched books and theses in the English language with titles suggesting that raptor survival may have been reported. Finally, we examined the references listed in those publications that had not been found in other ways. However, we cannot claim to have found every relevant publication in English, and for various reasons (such as small or obviously biased samples) some of those found are not included in this review.

Methods used to estimate survival

The methods used for survival estimation in birds have changed over time as different approaches have been developed and more sophisticated models have become available. The main methods are listed below, in approximate order of their development, and their strengths and limitations are discussed (Table 1). Methods 1 and 2 were the first to be used and can be classed as informal in that they lack statistical underpinning, whereas the later-developed methods 3-6 are all based on formal statistical models (as are later modifications of method 1).

1. Informal methods based on ring-recovery data from dead birds

Ring recovery data come from birds that have been marked (for raptors, typically as nestlings), released and subsequently found and reported dead, usually by members of the public. Under this sampling method, birds are normally encountered only once after ringing. In some countries, many species of birds, including raptors, have been ringed over many years, and databases of their initial marking and subsequent recovery have been maintained by national organizations (e.g. British Trust for Ornithology [BTO, http://www.bto.org] in the UK, United States Geological Survey [USGS, http://www.usgs.gov] Bird Banding Lab in the USA). Data from ring recovery studies allow estimates of survival probabilities and other parameters among birds of different sex and age classes, regardless of their dispersal distances, and may also provide information on causes of death (Newton 1979).

Estimating survival based on dead recoveries entails comparing, for a given cohort of nestlings, the number recovered dead in each year after ringing to give a 'static life table'. The rate at which recoveries decline with time since ringing reflects mortality. Usually, the samples for particular years are small, so the data for multiple years are pooled, and treated as a single cohort. Any time trends in survival and recovery parameters are thereby masked. Many estimates of survival have been made for raptors and other birds by this 'age ratio method', assuming no significant variation in survival or reporting rates over time or between age and sex groups, and no loss of rings during the lives of the birds (e.g. Haukioja & Haukioja 1970, Anderson *et al.* 1985, Newton & Rothery 1997). However, these assumptions are often rejected when tested, including in studies of raptors (Anderson *et al.* 1985). For these and other reasons, more statistically robust ring recovery models (e.g. Brownie *et al.* 1985) were developed to estimate bird survival, and their use for raptors has increased in recent years (Francis & Saurola 2002, Sulawa *et al.* 2010) (Supporting Online Table S1).

When used either with appropriate models or in conjunction with live-encounter (CMR) data, ring-recovery data offer opportunities to estimate true survival (see below), and to examine trends in survival over time.

2. Informal methods based on live encounters

These methods are based on re-trapping or re-sighting the same marked individuals, usually at nest sites, in the same study area in successive years. Minimum annual survival is estimated from the proportion of marked territorial birds present in one year that were also found to be present in a subsequent year – the 'return rate', or its complement 'turnover' (Newton 1979, Newton *et al.* 1983, Sandercock 2006, Hernández-Matías *et al.* 2011). The method does not distinguish between death and emigration, nor does it adequately account for detection (or recovery) probabilities that are <1 and vary over time during the study (Sandercock 2006). Survival estimates based on this method may, however, be acceptable for species that show high site fidelity and in which all (or almost all) individuals still in the study area can be detected each year. It has been used to estimate survival in Peregrine Falcons (Newton & Mearns 1988, Tordoff & Redig 1997), Eurasian Sparrowhawks *Accipiter nisus* (Newton & Rothery 1997) and various eagle species (Green *et al.* 1996, Carrete *et al.* 2002, Hernández-Matías *et al.* 2011), among others (Table S1).

This method of regular sampling can be challenging for species in which individuals are difficult to capture. Some of these challenges have been partially addressed by: (1) marking nestlings, or trapping and marking adults, in a way that they can be identified later without recatching them (e.g. Craig & Enderson 2004, Cadahía *et al.* 2005, Rosenfield *et al.* 2009); some recent studies have used PIT tags, allowing the 'electronic recapture' of birds at nests (Smith *et al.* 2015); (2) using non-invasive methods such as moulted feather patterns or DNA extracted from feathers found at nests to identify individuals (Newton 1986, Wink *et al.* 1999, Rudnick *et al.* 2005, Kenward & Katzner 2007); or (3) using individual recognition through plumage features (Hernández-Matías *et al.* 2011).

Annual adult survival has also been estimated in other ways from this type of data. In a breeding population (where breeding necessitates territory occupancy) with zero population growth, annual adult mortality is equal to the annual recruitment of new adults into the breeding population (Newton 1979, Newton & Mearns 1988). If all adults in the breeding population in previous years have been marked, so that, in any one year, they can be distinguished from new recruits, the proportion that new recruits form of the total breeders in that year reflects the annual mortality of established breeders. Once a study has continued long enough for all individuals to be of known age, the ratio of each age group to the next can be used to estimate the average annual survival, as in method 1 above.

3. Methods based on Capture-Mark-Recapture (CMR)

Like Method 2, CMR studies involve capturing, marking and releasing individuals over time, and keeping track of marked individuals so that an individual capture history can be constructed. The main difference is that formal statistical models are used to analyse the resulting data. For territorial raptor populations, multistate CMR models provide a flexible means of dealing with the related age-specific phenomena of survival and recruitment to a breeding population (Williams *et al.* 2002). Specifically, birds ringed as nestlings may have no chance of being detected (re-sighted) until they join the territorial breeding population, leading to biased estimates based on standard age-specific CMR models. More generally, individuals of the same age may have both different detection probabilities and different survival rates according to whether or not they have acquired a nesting territory, the latter birds often being termed 'floaters'. An appropriate multistate model structure for birds ringed as nestlings in these situations considers a pre-recruitment state with detection probability fixed to zero, together with age-specific recruitment probabilities reflecting the likelihood

that a pre-recruit will enter the breeding population and thereafter be subject to breeder detection probabilities. In this way, CMR models provide a means of dealing with floaters.

These open population CMR models have been used to estimate survival of raptors, permitting gains from births and immigration, and losses from deaths and permanent emigration to be estimated between sampling occasions (Gould & Fuller 1995, Brown *et al.* 2006, Steenhof *et al.* 2006, Faccio *et al.* 2011, Altwegg *et al.* 2014). Examples are given in Tables 2 and S1.

Like Method 2 above, CMR-based approaches have limited use for estimating survival probabilities of species that show low site fidelity (i.e. when dispersal distances are long relative to the size of the study area), or for dispersing sex and age classes (Sandercock 2006), in which permanent emigration is a major cause of the disappearance of birds from a study area. However, open population CMR models can be used together with radio-tracking data to estimate true survival, or with ring-recovery or other ancillary data to estimate true survival and other population parameters (see below) (Burnham 1993, Williams *et al.* 2002, Kendall *et al.* 2006).

4. Formal methods based on joint live encounters and ring recoveries

Studies involving a combination of live encounters and ring recoveries also involve the capture, marking and release of individuals. Subsequent live-encounters usually derive from the area of initial release, but ring recoveries can come from a much larger geographical region, providing information on permanent emigration (Burnham 1993, Barker 1997, Williams *et al.* 2002). The main advantage of combining live encounter and dead recovery models is that apparent survival can be decomposed into the probabilities of true survival and site fidelity (to the area in which live encounters are made) (Burnham 1993, Barker 1997, Williams *et al.* 2002). Additionally, joint models can be used to test explicitly for the effects of age, sex, and environmental conditions on site fidelity and true survival.

Statistical models for joint analysis of live-encounter and dead-recovery data to estimate survival are well developed (Table 1), but have been used rarely on raptors, except for Peregrine Falcons *Falco peregrinus* (e.g. Kauffman *et al.* 2003, Craig & Enderson 2004, Smith *et al.* 2015) (Table S1).

5. Formal methods based on tracking data

Radio-tracking studies involve capturing birds, fitting them with tags, and monitoring their fates (Kenward 1987). Tracking devices can be 'conventional' (VHF, UHF) or satellitebased (e.g. Argos, GPS or Globalstar) (Fuller *et al.* 2005). Radio-tagging is one of the best methods for monitoring the fates of individuals because, using this method, all tagged birds can be detected with near-certainty if they are present within the search area, and their fates determined unambiguously. Using satellite-received tags, the fates of many individuals can be monitored simultaneously wherever in the world they travel (Bowman *et al.* 1995, Whitfield *et al.* 2004, McIntyre 2005). Technical advances through time have made radio-tracking more flexible and increasingly reliable in application.

Use of radio-tags sometimes allows researchers better to distinguish between losses attributed to death and permanent emigration, and hence to estimate true survival as opposed to apparent survival (the combination of mortality and permanent emigration) (Kenward 1999, Williams *et al.* 2002, Fuller *et al.* 2005), as well as to identify the cause of mortality when dead animals can be retrieved (Heisey & Patterson 2006). Radio-tracking data can also be used in conjunction with other data types (see below) to estimate survival rates from capture-recapture data. Consequently, statistical models have been developed to use both live-encounter and radio-tracking data to estimate true survival (Powell *et al.* 2000, Nasution *et al.* 2004).

Tracking data have provided estimates of survival in large raptors, such as eagles (Ferrer & Calderón 1990, Bowman *et al.* 1995, Harmata *et al.* 1999, Hunt 2002, McIntyre 2005) and hawks (Zelenak *et al.* 1997, Kenward 1999), that are difficult to monitor using other methods (Table S1). Tracking also provides the most reliable way to estimate juvenile (first-year) survival in species with low natal-site fidelity, or age-class survival among pre-breeders of long-lived species with delayed maturity (Ferrer & Calderón 1990, Bowman *et al.* 1995, Kenward 1999, Mannan & Matter 2004, McIntyre 2005, Davies & Restani 2006) (Table S1).

Set against these advantages, tracking studies can be prohibitively costly due to the expense of deploying sufficient numbers of tags and monitoring the individuals concerned. There can also be problems caused by the short life-spans of most transmitters, possible reduction in survival associated with the transmitters (Steenhof *et al.* 2005), and the effects of right-censoring when fate and censoring are not independent events (Bennetts *et al.* 1999, Williams

et al. 2002, Zens & Peart 2003). An additional difficulty arises when mortality cannot be distinguished from transmitter failure, a situation more likely in studies that track wide-ranging species via satellites. Signals that become irregular before stopping altogether are usually taken to signify tag failure, but signal characteristics and onboard sensors of some satellite tags can suggest whether a tag has failed, become detached or its carrier has died. Such tags are therefore especially useful in the study of survival.

6. Integrated Population Models

These models provide a framework that can use multiple types of data (typically, time series of abundance, productivity, and CMR and/or tag recovery data) to estimate demographic parameters and make statistical inferences about these parameters (Besbeas *et al.* 2002, Schaub & Abadi 2011, Kéry & Schaub 2012, Tenan *et al.* 2012). Developing integrated population models involves three steps (Schaub & Abadi 2011, Kéry & Schaub 2012): (1) the development of a model that links multiple data types that pertain to the same demographic process(es); (2) development of likelihoods for individual datasets; and (3) integration of these likelihoods to form a joint likelihood for all data types, permitting estimation of relevant parameters and statistical inferences using either frequentist or Bayesian analytical frameworks.

The advantages of integrated population models include: (1) the combination of information contained in several separate datasets, so that parameters estimated using this approach are generally more precise and statistical inference is stronger than would be possible using independent analyses of individual datasets; (2) the estimation of more demographic parameters than would be possible if each dataset were analysed separately; and (3) the direct estimation of standard errors (or credible intervals) for all estimable parameters (Besbeas *et al.* 2002, Schaub & Abadi 2011, Kéry & Schaub 2012). So far, integrated population models have been used to estimate survival probabilities (and other demographic parameters) among raptors for Eurasian Eagle Owl *Bubo bubo*, California Spotted Owl *Strix o. occidentalis* and Red Kite *Milvus milvus* (Schaub *et al.* 2010, Tenan *et al.* 2012, Tempel *et al.* 2014).

Methods used to examine survival in relation to body mass and sex in different species To estimate survival in relation to body mass for different species, we used only data for adult birds explicitly obtained by the statistically formalised methods 3-6 above, thus excluding estimates from methods 1-2 which were most open to bias. Where separate estimates were available for the sexes, we used gender-specific masses; where estimates were for males and females combined, or where the sexes of the birds in the study were not known, we used the arithmetic mean of the male and female masses. Where separate estimates were available for different years in the same study of a species or when estimates were made for different ages of adult birds, we took the geometric mean of those estimates. Where estimates were estimates for each area or circumstances in the same study, we used separate estimates for each area or circumstance (e. g. low versus high prey abundance, one habitat versus another). This meant that variable numbers of estimates were available for different species, which we allowed for statistically by treating species as a random effect in linear mixed effect models (Zuur *et al.* 2009). All estimates used to calculate relationships between survival and body mass or sex are given in Table 2, and the full data are summarised in Table S1. Table 2 lists 45 published survival estimates for 30 species of diurnal raptors, and 23 for nine species of owls. Table S1 lists survival estimates of 45 species of diurnal raptors from 130 publications, and ten species of owl from 29 sources.

RESULTS

Species studied and constraints in data

We found survival estimates for three species of Cathartidae, one of Pandionidae, 29 of Accipitridae, 12 of Falconidae, one of Tytonidae and nine of Strigidae (Table S1). In most of these species, more than one estimate was available from different areas or time periods. Almost all of the 159 studies were from the temperate region of the northern hemisphere. Most attention has been directed to species that suffered severe population declines or were considered threatened or endangered for other reasons (e.g. Newton 1986, Ratcliffe 1993, Anthony *et al.* 2006, Forsman *et al.* 2011, Ganey *et al.* 2014).

Among these studies, some reported sex-, age-, region- or time-specific variation in survival (Newton & Rothery 1997, Newton *et al.* 1997, Martin *et al.* 2006, Karell *et al.* 2009, Forsman *et al.* 2011, Ganey *et al.* 2014), while others simply provided estimates of overall, or pre-breeder and breeder survival (Table S1). Owing to difficulties of recapturing, re-sighting or tracking birds until they become breeding adults, and the limited use of the relatively new statistical models developed to deal with such difficulties, estimates of juvenile and pre-breeding survival were fewer, more variable and probably less reliable than estimates of adult survival, especially for species with delayed maturity (see below). Furthermore, some of the

studies listed in Table S1 refer to populations in which survival rates were lower than expected because of human behaviour of one form or another: for example, lead poisoning of California Condors *Gymnogyps californianus* or direct persecution of Hen Harriers *Circus cyaneus* (Etheridge *et al.* 1997, Meretsky *et al.* 2000). These were obvious cases, but many other studies, especially in the early years, may have involved populations subject to human killing. Despite some variability and inconsistencies among estimates, some general patterns in the survival rates of raptors and owls were evident.

Body-size and survival

In line with well-established allometric relationships (Calder 1984, Gaillard *et al.* 1989, Charnov 1993, Dobson & Jouventin 2010), adult survival among different raptor species was positively related to log-transformed adult body mass in both diurnal raptors and owls (Fig. 1), although with much additional variation within and between species. Analysis of covariance with species as a random effect in a linear mixed model (implemented using R package *nlme*; Pinheiro *et al.* 2014) provided no evidence of different slopes for diurnal raptors and owls (t = 0.806, P = 0.422). In populations largely free of human-induced mortality, small species, such as Eurasian Sparrowhawk and Common Kestrel *Falco tinnunculus*, typically showed annual adult survival rates of around 60-70%, while medium sized falcons and hawks showed adult survival rates around 80-90%, and large eagles and vultures of more than 90%, although exceptions occurred among the estimates for all three groups (Tables 2, S1).

Age differences in survival

It would be expected that young birds, because of their inexperience and low social status, would survive less well than older individuals. Most studies reporting age-specific estimates indicated that annual survival rates of juveniles or sub-adults were substantially lower than those of adults of the same species (for exceptions see Bowman *et al.* 1995 for Bald Eagle *Haliaeetus leucocephalus*, Sulawa *et al.* 2010 for White-tailed Eagle *H. albicilla*). Furthermore, in studies in which adult age-classes were distinguished, survival of the very oldest age groups among breeders tended to decline, presumably reflecting senescence or relegation to floater status (e. g. Newton *et al.* 1997, Sergio *et al.* 2011, Tenan *et al.* 2012, Ganey *et al.* 2014). In populations of long-lived eagles, in which competition for places in the breeding population is often high, mortality rates amongst late-aged sub-adults may also be high (e. g. Saurola *et al.* 2003), a situation attributed to aggressive interactions between

territory seekers and territory holders (Haller 1996, Sulawa *et al.* 2010). Such competition may result in some adults being killed, but the effect on the overall rate of adult mortality may be less obvious if it disproportionately affects senescent birds. Evans *et al.* (2009) found no increase in mortality in White-tailed Eagles as they approached adulthood in an environment in which potential but unoccupied breeding habitat was abundantly available, although adults survived generally better than non-adults.

The degree of difference in estimated survival between juvenile and adult raptors depends largely on estimation method (Table S1). Because juveniles disperse in greater proportion and over longer distances than adults, any method based on a confined study area that does not distinguish between death and emigration leads to under-estimation of juvenile survival, and enhances the difference in estimated survival between juveniles and adults. In theory, ring recoveries could provide reliable estimates of survival in different age groups, including first-year birds, but it has long been suspected that dead first-year birds are more likely to fall into human hands than dead older birds, which based on Method 1 would overestimate firstyear mortality (Newton 1979, Frances & Saurola 2002). Juveniles are more likely to be shot or trapped, or to venture near human habitation. For these reasons, estimates for juveniles based on radio-tracking are probably the most reliable available, providing that the tags do not affect survival, that right censoring is independent of bird fate, that they function properly for long enough and that attempts are made to follow each individual wherever it goes. We judge that at least four of the radio-tracking studies in Table S1 meet these criteria sufficiently well to provide reliable estimates of first-year survival. These studies gave estimates of adult and first-year survival in the Bald Eagle of 88% and 71% respectively, and in the Common Buzzard Buteo buteo of 88-91% and 66-73% respectively (Bowman et al. 1995, Kenward et al. 2000). In the Snail Kite Rostrhamus sociabilis, estimates of adult survival in three different years were given as 89%, 86% and 82%, and of juvenile survival in the same years as 67%, 45% and 44% respectively (Bennetts et al. 1999). In the Goshawk Accipiter gentilis, with an adult survival of 83% in both sexes, estimated first-year survival was 71% in females and 59% in males, the only study to separate the sexes of first year birds (Kenward et al. 1999). Finally, in a study of the Red Kite that used an integrated population model to provide the estimates, adult and first-year survival were calculated at 96% and 89%, respectively (Tenan et al. 2012). So in these various estimates, first-year survival emerged as 7-48 percentage points lower than adult survival in the same population. Other estimates in Table S1 obtained by radio-tracking refer to adult and first-year survival in different

populations of a species, so are less strictly comparable, while some of the estimates obtained by other methods could be acceptably accurate, but the methodology leaves them open to doubt.

Sex differences in survival

In raptors and owls, females are bigger than males. In most species this difference is small, but in others females can weigh up to twice as much as males (Newton 1979). Particular interest therefore attaches to any sex differences in survival, and on the basis of weight, females would be expected to show higher annual survival rates than males. For 32 comparisons involving seven species of diurnal raptors and four species of owls listed in Table S1, separate survival estimates with standard errors were available for adults of both sexes. In most of these comparisons, the estimates for each sex varied by no more than a few percentage points, and either males or females could show higher survival. However, differences exceeding five percentage points were found in 12 comparisons. In diurnal raptors these large differences were in highly dimorphic species, but they were not consistent within species (see the different estimates for Peregrine Falcon and Eurasian Sparrowhawk in Table S1). Among owls, sex differences greater than five percentage points were found in one study of Tawny Owls Strix aluco (females showing higher survival), and in three studies of Burrowing Owls Athene cunicularia (males showing higher survival in two, females in one). Sufficient information was provided using methods 3-6 to test the statistical significance of apparent sex differences in seven studies of diurnal raptors, and in 23 studies of owls, including 16 of Spotted Owls. Only four were statistically significant (program CONTRAST; Hines & Sauer 1989): Southern Crested Caracara *Caracara plancus*: $\chi^2 =$ 69.230, female 3% higher than male, P < 0.0001, Morrison 2003; Hen Harrier, $\gamma^2 = 5.586$, female 18% higher than male, P = 0.018, Picozzi 1984; two of three studies of Burrowing Owl, $\chi^2 = 4.5$, male 12% higher than female, P = 0.034, Millsap 2002; $\chi^2 = 15.56$, male 23% higher than female, P = 0.0001, Wellicome *et al.* 2014).

Finally, we tested the relationship between the ratios of female:male survival and body mass. With species included as a random effect, no evidence of a relationship emerged, either for diurnal raptors alone (t = -1.536, P = 0.199), owls alone (t = 1.240, P = 0.341) or both groups combined (t = -0.647, P = 0.524). Sex differences in survival of owls and raptors might be expected for reasons other than body size: for example, human persecution (in which females suffer higher mortality because they are more easily killed at the nest), and differential

migration (in which the sexes winter in partly different regions imposing different mortality rates). In view of these findings and possibilities, we conclude that the data provide no convincing or consistent evidence for differential survival between the sexes of adult raptors and owls linked to size dimorphism.

Possible methodological differences in survival estimates

Examination of survival rates of Eurasian Sparrowhawks and Peregrine Falcons studied in different time periods hinted that survival rate estimates may often have been lower in earlier than in later periods. This apparent temporal trend was not statistically significant in either species, and could in any case be influenced by change in estimation methods. Nevertheless, a change in survival over the years could have been expected in response to legal protection and banning of organochlorine pesticides, as the various survival estimates for these species spanned the period before and after these events (e.g. Newton 1986, Cade *et al.* 1988, Wyllie & Newton 1991, Newton & Wyllie 1992).

It is accepted that estimation methods alone can cause substantial variation in survival estimates (Clobert & Lebreton 1991). For example, Francis and Saurola (2002) compared estimates of age-specific survival among Tawny Owls from the same population using formal estimation approaches based on alternative data types: (1) recoveries of birds ringed as nestlings; (2) recoveries of birds tagged as juveniles and adults; (3) recaptures of birds tagged as juveniles; (4) recaptures of birds tagged as juveniles and adults; (5) recoveries and recaptures of birds tagged as juveniles; and (6) recoveries and recaptures of birds tagged as juveniles and adults. Depending on the data type and analytical method, estimated survival of first-year birds ranged from 10.1% to 47.8%, while that for adults was effectively the same at 69.9-72.2%. First-year survival based on recoveries of birds ringed as nestlings was biased high because of violation of the assumption of age-independent recovery rates: juveniles were recovered disproportionately more often than older birds. Differences in survival caused primarily by estimation method or data type have also been reported for the Peregrine Falcon (Gould & Fuller 1995), Eurasian Sparrowhawk (Newton 1986), Bonelli's Eagle Aquila fasciata (Hernández-Matías et al. 2011), Snail Kite (Bennetts et al. 1999) and Red Kite (Tenan et al. 2012).

DISCUSSION

Apparent survival versus true survival

In studies based on specific study areas, most published estimates of survival also include permanent emigration. In effect, they estimate the product of true survival and site fidelity. The degree to which true and apparent survival in adults differs depends on the proportion of adults that leave the area, and in some species, such as Peregrine Falcon and Bald Eagle, this proportion seems so low that apparent survival could closely approximate true survival. Although estimating true survival is desirable, it may be sufficient to estimate apparent survival when comparing rates among different groups of birds of the same species (treatments, management options, habitats, or other factors), if permanent emigration is similar among the groups being compared. For studies directed at changes in abundance on a study area, it may not be necessary to split losses into deaths versus emigration, or gains into local reproduction versus immigration (see Appendix 3 of Franklin et al. 2004). However, true survival estimates are required when the objective is to test life-history theory based on comparative studies, to estimate population growth rates using projection matrix approaches or to develop recovery plans for imperilled species. In all these cases, assessment of population status, or population growth rate based on apparent survival without appropriately accounting for immigration may result in misleading conclusions (e.g. Cooch et al. 2001, Gerber 2005). In particular, it is common practice to assess the conservation status of species by computing asymptotic population growth rates using population projection matrices (e.g. Caswell 2001). When such matrices use apparent survival, then population losses include movement, while the recruitment information used in such matrices typically includes reproduction (clutch size, nest success, etc.), but not immigration. Such matrices are therefore asymmetric with respect to movements, typically including movement in estimates of losses (emigration) but not in estimates of gains (immigration), and thereby leading to underestimates of projected growth rates (e.g. Nichols & Hines 2002). This is a methodological deficiency which is by no means specific to raptors and owls.

Demographic analyses of, for example, Spotted Owls (e. g. Franklin *et al.* 2004, Forsman *et al.* 2011) use CMR methods (e.g. Pradel 1996) at specific study sites to estimate realized (as contrasted with asymptotic) population growth rates directly. These growth rates estimate changes in numbers of birds on specific study sites, with losses including both permanent emigration and death, and gains including recruitment from both local reproduction and

immigration. These growth rates are symmetric with respect to movement and are intended to reflect true changes in numbers in the landscape, rather than theoretical changes that would occur if there were no movement.

The question of whether variation in survival detected in comparative analyses reflects true variation in survival or variation in methodology is also relevant to many other groups of birds. Historically, investigations of waterfowl were among the first to use probabilistic methods that considered the detection process (e.g. Johnson *et al.* 1992). Critiques such as those of Clobert and Lebreton (1991) and Boulinier *et al.* (1997) led to accelerating adoption of CMR methods for survival studies of birds. Some studies of raptors provide examples of cutting-edge research in demography and population dynamics. Examples include studies on the Spotted Owl (e.g., Blakesley *et al.* 2010, Forsman *et al.* 2011, Ganey *et al.* 2014), Red Kite (Tavecchia *et al.* 2012, Tenan *et al.* 2012) and Eagle Owl (Schaub *et al.* 2010). Although raptors are generally rarer than many other birds, study of this group has contributed substantially to our understanding of avian population dynamics, in part because many raptor species are relatively conspicuous and long-lived, and can carry large visual identification marks or transmitters. They are also highly territorial which, together with site fidelity in most species, gives high re-encounter rates.

Territorial breeders and floating non-breeders

Estimates of adult survival in raptors are mostly based on breeding birds, and so do not include floating (non-breeding) adults. This is true of many other birds, including some that move from breeder to floater status in different years in response to factors such as fluctuations in food supply. Non-breeding adults could survive better or worse than breeding adults, depending on the conditions in which they find themselves. For example, in a landscape which is filled to capacity with territorial breeding pairs, floaters may be constrained to occupy less favourable areas where their mortality rates are higher. Alternatively, in landscapes where food is plentiful but breeding sites are limiting, floaters may be able to survive as well or better than breeders, though unable to obtain a nesting territory (e. g. Hunt *et al.* 1998, Newton 1998). In addition, survival of breeding adult raptors may be lower than that of floaters in areas where adults are often shot at the nest, a common occurrence in some species in some areas (e. g. Etheridge *et al.* 1997, Whitfield *et al.* 2004).

Continuing threats to raptors

Although raptor populations have generally recovered from the global declines caused by organochlorine pesticides during the 1950-1960's (e. g. Newton 1998, Banks et al. 2010), there is no shortage of new threats. Inadvertent diclofenac poisoning has caused a catastrophic collapse of populations of several species of south Asian vultures (Oaks et al. 2003, Green et al. 2006), and the toxic effects of lead ingestion have had adverse impacts on the critically endangered California Condor, the rare Steller's Sea Eagle Haliaeetus pelagicus and other raptors (Watson et al. 2009). Organophosphate insecticides have caused mass mortality of Swainson's Hawks Buteo swainsoni and other raptors in South America (Goldstein et al. 1996). Illegal poisoning or other killing affected 40% of 103 Red Kites found dead in Scotland (Smart et al. 2010), and was the most important recent cause of mortality of Red Kites in Spain, suppressing population growth rate by 20% (Tenan et al. 2012); it is also an ongoing problem in parts of Africa greatly reducing vulture numbers (Ogada et al. 2012, 2015). Those raptors that are predators of game birds or racing pigeons have long been subject in Britain and elsewhere to shooting, poisoning and other forms of persecution (e. g. Etheridge et al. 1997, Whitfield et al. 2004, 2008, Newton 2013). Other relatively new threats to raptors include collisions and electrocutions related to power generation and distribution (including wind turbines), and communication towers (Subramanian 2012, Tavecchia et al. 2012, Angelov et al. 2013), and the unsustainable harvest of some species in some regions for falconry (Kovács et al. 2014). As with previous threats, mortality due to these anthropogenic causes is often partly or entirely additive to natural mortality, and frequently leads to population declines. Those raptor species that migrate are thereby exposed to different mortality threats in different regions.

Future studies

In addition to providing data necessary to estimate survival probabilities, radio-tracking studies can provide a wealth of other information (e. g. local movements, dispersal and migration routes, space and habitat use, abundance) unattainable in other ways (Fuller *et al.* 2005). We can therefore expect that radio-tracking will continue to play an important role in raptor research. However, it is expensive, requires high investment of time and effort, and radio-transmitters can affect survival (Steenhof *et al.* 2006). In most analyses of radio-tracking data, individual birds are censored for one reason or another, and a challenge in such analyses is to ensure that censoring is independent of bird fate, as dependence can produce biased survival estimates (Bennetts *et al.* 1999).

In future, we can also expect to see an increase in the number of studies that use multiple data types, including time series of annual counts, radio-tracking, live-recaptures, mark-resighting, and dead-recoveries (from a larger area), all incorporated within a single analytical framework (Burnham 1993, Barker 1997, 1999, Williams et al. 2002, Nasution et al. 2004, Schaub et al. 2010, Tenan et al. 2012). The use of financial incentives can improve rates of ring-reporting by members of the general public in hunted species (Nichols et al. 1991, 1995), as can a shift from standard mail to telephone and web-based reporting (Royle & Garrettson 2005, Boomer et al. 2013). In addition, the use of electronic recaptures using technologies such as Passive Integrated Transponders can potentially improve recapture rates and quality of monitoring data (Barbour et al. 2013). Recent developments in GPS tracking technology that allow the precise location of an animal at frequent intervals, can provide data necessary for accurate and precise estimates of survival and other demographic measures, as well as pin-pointing the time and place of death (McIntyre 2012). Data sharing and collaborative initiatives (e.g. Movebank; Kranstauber et al. 2011) offer opportunities to overcome sample size limitations, extend temporal and spatial scales of inference, harness analytical skills of other researchers and potentially facilitate analyses of older data using new methods. Future research should be driven mostly by conservation concerns, perceived challenges and available funding. However, studies on relatively common species may continue to fill gaps in understanding, while improved survival estimation procedures and advances in technology make population studies of some hitherto 'difficult' species feasible. Whereas species such as the Common Kestrel and Eurasian Sparrowhawk provided initial insight into raptor ecology, it is now possible to effectively study other species that, for example, are larger, longer-lived and have delayed maturity, and also to accumulate information on sex- and agespecific survival. Given the critical roles of raptors in a variety of ecosystems and their roles as indicator and flagship species, investment in raptor research that uses state-of-the-art

methods to estimate mortality may be well justified.

We thank all those researchers who have published estimates of survival in raptors and owls. Data collected by Richard Mearns, George Smith, Chris Rollie and a large number of volunteers in southern Scotland and northern England permitted us to undertake some analyses reported in this manuscript. We are greatly indebted to Dr. Jim Nichols who offered much insightful advice through the preparation of this paper, and also Drs. B. Arroyo, B. Sandercock, G. Tavecchia, G. Hunt and an anonymous reviewer for many helpful comments and feedback at various stages.

- Altwegg, R., Jenkins, A. & Abadi, F. 2014. Nestboxes and immigration drive the growth of an urban Peregrine Falcon *Falco peregrinus* population. *Ibis* 156: 107-115.
- Anderson, D. R., Burnham, K. P. & White, G. C. 1985. Problems in estimating agespecific survival rates from recovery data of birds ringed as young. J. Anim. Ecol. 54: 89-98.
- Angelov, I., Hashim, I. & Oppel, S. 2013. Persistent electrocution mortality of Egyptian
 Vultures *Neophron percnopterus* over 28 years in East Africa. *Bird Conserv. Int.* 23:
 1-6.
- Anthony, R. G., Forsman, E. D., Franklin, A. B., Anderson, D. R., Burnham, K. P.,
 White, G. C., Schwarz, C. J., Nichols, J. D., Hines, J. E., Olson, G. S., Ackers, S.
 H., Andrews, L. S., Biswell, B. L., Carlson, P. C., Diller, L. V., Dugger, K. M.,
 Fehring, K. E., Fleming, T. L., Gerhardt, R. P., Gremel, S. A., Gutierrez, R. J.,
 Happe, P. J., Herter, D. R., Higley, J. M., Horn, R. B., Irwin, L. L., Loschl, P. J.,
 Reid, J. A. & Sovern, S. G. 2006. Status and trends in demography of Northern
 Spotted Owls, 1985-2003. *Wildlife Monogr.* 163: 1-48.
- Banks, A. N., Crick, H. Q. P., Coombes, R., Benn, S., Ratcliffe, D. A. & Humphreys, E.
 M. 2010. The breeding status of Peregrine Falcons *Falco peregrinus* in the UK and Isle of Man in 2002. *Bird Study* 57: 421-436.
- Barbour, A. B., Ponciano, J. M. & Lorenzen, K. 2013. Apparent survival estimation from continuous mark-recapture/resighting data. *Methods Ecol. Evol.* 4: 846-853.
- **Barker, R.** 1997. Joint modeling of live-recapture, tag-resight, and tag-recovery data. *Biometrics* **53**: 666-677.
- **Barker, R.** 1999. Joint analysis of mark-recapture, resighting and ring-recovery data with age-dependence and marking-effect. *Bird Study* **46**: 37-41.

- Bennetts, R. E., Dreitz, V. J., Kitchens, W. M., Hines, J. E. & Nichols, J. D. 1999. Annual survival of Snail Kites in Florida: radiotelemetry versus capture-resighting data. *Auk* 116: 435-447.
 - Besbeas, P., Freeman, S. N., Morgan, B. J. T. & Catchpole, E. A. 2002. Integrating markrecapture–recovery and census data to estimate animal abundance and demographic parameters. *Biometrics* 58: 540-547.
 - Blakesley, J. A., Seamans, M. E., Conner, M. M., Franklin, A. B., White, G. C.,
 Gutierrez, R. J., Hines, J. E., Nichols, J. D., Munton, T. E., Shaw, D. W. H.,
 Keane, J. J., Steger, G. N. & McDonald, T. L. 2010. Population dynamics of
 Spotted Owls in the Sierra Nevada, California. *Wildlife Monogr.* 174: 1-36.
 - Boomer, G. S., Zimmerman, G. S., Zimpfer, N. L., Garrettson, P. R., Koneff, M. D., Sanders, T. A., Magruder, K. D. & Royle, J. A. 2013. Band reporting probabilities for Mallards recovered in the United States and Canada. *J. Wildl. Manage.* 77: 1059-1066.
 - Boulinier, T., Sorci, G., Clobert, J. & Danchin, E. 1997. An experimental study of the costs of reproduction in the Kittiwake *Rissa tridactyla*: Comment. *Ecology* 78: 1284.
 - Bowman, T. D., Schempf, P. F. & Bernatowicz, J. A. 1995. Bald Eagle survival and population dynamics in Alaska after the" Exxon Valdez" oil spill. *J. Wildl. Manage*. 59: 317-324.
 - Bretagnolle, V., Mougeot, F. & Thibault, J. C. 2008. Density dependence in a recovering Osprey population: demographic and behavioural processes. J. Anim. Ecol. 77: 998-1007.
 - Brown, J. L., Collopy, M. W., Gott, E. J., Juergens, P. W., Montoya, A. B. & Hunt, W.G. 2006. Wild-reared Aplomado Falcons survive and recruit at higher rates than hacked falcons in a common environment. *Biol. Conserv.* 131: 453-458.

Brownie, C., Anderson, D. R., Burnham, K. P. & Robson, D. S. 1985. Statistical inference from band recovery data - A handbook, Washington, D.C., USA: U.S. Fish and Wildlife Service, Resource Publication No. 156.

- Burnham, K. P. 1993. A theory for combined analysis of ring recovery and recapture data in marked individuals. In Lebreton, J. D. & North, P. M. (eds.) *Marked individuals in the study of bird populations*:199-213. Basel, Switzerland: Birkhaeuser-Verlag.
- Cadahía, L., Urios, V. & Negro, J. J. 2005. Survival and movements of satellite-tracked Bonelli's Eagles *Hieraaetus fasciatus* during their first winter. *Ibis* 147: 415-419.
- Cade, T. J., Enderson, J. H., Thelander, C. G. & White, C. M. 1988. Peregrine Falcon populations: their management and recovery, Boise, USA: The Peregrine Fund.
- Calder, W. A. 1984. Size, function, and life history. Cambridge, MA: Harvard University Press.
- Carrete, M., Sánchez-Zapata, J. A., Martínez, J. E. & Calvo, J. F. 2002. Predicting the implications of conservation management: a territorial occupancy model of Bonelli's Eagle in Murcia, Spain. *Oryx* 36: 349-356.
- **Caswell, H.** 2001. *Matrix population models: construction, analysis, and interpretation,* Sunderland, Massachusetts, USA: Sinauer Associates.
- **Charnov, E. L.** 1993. *Life history invariants: some explorations of symmetry in evolutionary ecology*, Oxford: Oxford University Press.
- Clobert, J. & Lebreton, J. -D. 1991. Estimation of demographic parameters in bird populations. In Perrins, C. M. Lebreton, J. D. & Hirons, G. J. M. (eds.) *Bird population studies relevance to conservation and management*: 75-104. Oxford: Oxford University Press.
- Conn, P. B., Kendall, W. L. & Samuel, M. D. 2004. A general model for the analysis of mark-resight, mark-recapture, and band-recovery data under tag loss. *Biometrics* 60:

900-909.

- Cooch, E., Rockwell, R. F. & Brault, S. 2001. Retrospective analysis of demographic responses to environmental change : a lesser snow goose example. *Ecol. Monogr.* 71: 377-400.
- Craig, G. R. & Enderson, J. 2004. Peregrine Falcon biology and management in Colorado, 1973-2001, Boulder, Colorado: Colorado Division of Wildlife Technical Publication No. 43.
- Craig, G. R., White, G. C. & Enderson, J. H. 2004. Survival, recruitment, and rate of population change of the Peregrine Falcon population in Colorado. *J. Wild Manage*.
 68: 1032-1038.
- Davies, J. M. & Restani, M. 2006. Survival and movements of juvenile Burrowing Owls during the post-fleding period. *Condor* 108: 282-291.
- Dobson, F. S. & Jouventin, P. 2010. The trade-off of reproduction and survival in slowbreeding seabirds. *Can. J. Zoolog.* 88: 889-899.
- Dunning, J. B. 1992. CRC handbook of avian body masses (1st ed.). Boca Raton, Florida: CRC Press.
- Dunning, J. B. 2007. CRC handbook of avian body masses (2nd ed.). Boca Raton, Florida: CRC Press.
- Etheridge, B., Summers, R. W. & Green, R. E. 1997. The effects of illegal killing and destruction of nests by humans on the population dynamics of the Hen Harrier *Circus cyaneus* in Scotland. *J. Appl. Ecol.* **34**: 1081-1105.
- Evans, R. J., Wilson, J. D., Amar, A., Douse, A., Maclennan, A., Ratcliffe, N. &
 Whitfield, D. P. 2009. Growth and demography of a re-introduced population of
 White-tailed Eagles *Haliaeetus albicilla*. *Ibis* 151: 244-254.

Faccio, S. D., Amaral, M., Martin, C. J., Lloyd, J. D., French, T. W. & Tur, A. 2011.

Movement patterns, natal dispersal, and survival of Peregrine Falcons banded in New England. Technical Report to USFWS. Concord, USA: Vermont Center for Ecostudies.

- Ferrer, M. & Calderón, J. 1990. The Spanish Imperial Eagle Aquila adalberti C.L. Brehm 1861 in Doñana National Park (south west Spain): A study of population dynamics. *Biol. Conserv.* 51: 151-161.
- Forsman, E. D., Anthony, R. G., Dugger, K. M., Glenn, E. M., Franklin, A. B., White,
 G. C., Schwarz, C. J., Burnham, K. P., Anderson, D. R., Nichols, J. D., Hines, J.
 E., Lint, J. B., Davis, R. J., Ackers, S. H., Andrews, L. S., Biswell, B. L., Carlson,
 P. C., Diller, L. V., Gremel, S. A., Herter, D. R., Higley, J. M., Horn, R. B., Reid,
 J. A., Rockweit, J., Schaberel, J., Snetsinger, T. J. & Sovern, S. G. 2011.
 Population demography of Northern Spotted Owls. *Stud. Avian Biol.* 40: 1-106.
- Francis, C. M. & Saurola, P. 2002. Estimating age-specific survival rates of tawny owls recaptures versus recoveries. J. Appl. Stat. 29: 637-647.

Franklin, A. B., Gutiérrez, R. J., Nichols, J. D., Seamans, M. E., White, G. C.,
Zimmerman, G. S., Hines, J. E., Munton, T. E., Lahaye, W. S., Blakesley, J. A.,
Steger, G. N., Noon, B. R., Shaw, D. W. H., Keane, J. J., Mcdonald, T. L. &
Britting, S. 2004. Population dynamics of the California Spotted Owl (*Strix occidentalis occidentalis*): a meta-analysis. *Ornithological Monographs* 54: 1-54.

- Fuller, M. R., Millspaugh, J. J., Church, K. E. & Kenward, R. E. 2005. Wildlife radiotelemetry. In Braun, C.E. (ed) *Techniques for wildlife investigations and management*: 377-417. Bethesda, USA: The Wildlife Society.
- Gaillard, J. M., Pontier, D., Allaine, D., Lebreton, J. D., Trouvilliez, J. & Clobert, J. 1989. An analysis of demographic tactics in birds and mammals. *Oikos* 56: 59-76.

Ganey, J. L., White, G. C., Ward, J. P., Kuyle, S. C., Apprill, D. L., Rawlinson, T. A. &

Jonnes, R. S. 2014. Demography of Mexican Spotted Owls in the Sacramento mountains, New Mexico. *J. Wildl. Manage.* **78**: 42-49.

- Gerber, L. R. 2005. The role of dispersal and demography in determining the efficacy of marine reserves. *Can. J. Zoolog.* 62: 863-871.
- Goldstein, M. I., Woodbridge, B., Zaccagnini, M. E. & Parana, S. 1996. An assessment of mortality of Swainson's Hawks on wintering grounds in Argentina J. Raptor Res. 30: 106-107.
- Gould, W. R. & Fuller, M. R. 1995. Survival and population size estimation in raptor studies a comparison of two methods. J. Raptor Res. 29: 256-264.
- Green, R. E., Pienkowski, M. W. & Love, J. 1996. Long-term viability of the re-introduced population of the White-tailed Eagle *Haliaeetus albicilla* in Scotland. *J. Appl. Ecol.*33: 357-368.
- Green, R. E., Taggart, M. A., Das, D., Pain, D. J., Kumar, C. S., Cunningham, A. A. &
 Cuthbert, R. 2006. Collapse of Asian vulture populations: risk of mortality from residues of the veterinary drug diclofenac in carcasses of treated cattle. *J. Appl. Ecol.*43: 949-956.
- Haller, H. 1996. Der Steinadler in Graubünden: Langfristige Untersuchungen zur
 Populationsökologie von Aquila chrysaetos im Zentrum der Alpen. Der
 Ornithologische Beobachter 9.
- Harmata, A. R., Montopoli, G. J., Oakleaf, B., Harmata, P. J. & Restani, M. 1999. Movements and survival of Bald Eagles banded in the Greater Yellowstone ecosystem. J. Wildl. Manage. 63: 781-793.
- Haukioja, E. & Haukioja, M. 1970. Mortality rates of Finnish and Swedish Goshawks Accipiter gentilis. Finnish Game Research 31: 13-20.

Heisey, D. M. & Patterson, B. R. 2006. A review of methods to estimate cause-specific

mortality in presence of competing risks. J. Wildl. Manage. 70: 1544-1555.

- Hernández-Matías, A., Real, J. & Pradel, R. 2011. Quick methods for evaluating survival of age-characterizable long-lived territorial birds. *J. Wildl. Manage.* **75**: 856-866.
- Hines, J. E. & Sauer, J. R. 1989. Program CONTRAST A general program for the analysis of several survival or recovery rate estimates. Washington, DC: U. S. Fish & Wildlife Service, Fish & Wildlife Technical Report 24.
- Hunt, G. 2002. Raptor floaters at Moffat's equilibrium. Oikos 82: 191-197.
- Hunt, W., Jackman, R., Hunt, T., Driscoll, D. & Culp, L. 1998. A population study of Golden Eagles in the Altamont pass wind resource area: population trend analysis 1994-1997. Report to the National Renewable Energy Laboratory. Santa Cruz, USA: Predatory Bird Research Group.
- Johnson, D. H., Nichols, J. D., & Schwartz, M. D. 1992. Population dynamics of breeding waterfowl. Ecology and management of breeding waterfowl. In Bruce D. J.
 Batt, Afton, A. D., Anderson, M. G., Ankney, C. D., Johnson, D. H., Kadlec, J. A. & Krapu, G. L. (eds.) *Ecology and Management of Breeding Waterfowl:* 446-485.
 Minneapolis: University of Minnesota Press.
- Karell, P., Ahola, K., Karstinen, T., Zolei, A. & Brommer, J. E. 2009. Population
 dynamics in a cyclic environment: consequences of cyclic food abundance on Tawny
 Owl reproduction and survival. *J. Anim. Ecol.* 78: 1050-1062.
- Kauffman, M. J., Frick, W. F. & Linthicum, J. 2003. Estimation of habitat-specific demography and population growth for Peregrine Falcons in California. *Ecol. Appl.* 13: 1802-1816.
- Kenward, R. E. 1987. Wildlife radio tagging equipment, field techniques and data analysis. London: Academic Press.

Kenward, R. E. 1999. Demographic estimates from radio-tagging: models of age-specific

survival and breeding in the Goshawk. J. Anim. Ecol. 68: 1020-1033.

- Kenward, R. & Katzner, T. 2007. Rapid sustainability modeling for raptors by radiotagging and DNA-fingerprinting. J. Wildl. Manage. 71: 238-245.
- Kenward, R. E., Marcstom, V. & Karlbom, M. 1999. Demographic estimates from radio-tagging: models of age-specific survival and breeding in the Goshawk. *J. Anim. Ecol.*68: 1020-1033.
- Kenward, R. E., Walls, S. S., Hodder, K. H., Pahkala, M., Freeman, S. N. & Simpson, V.
 R. 2000. The prevalence of non-breeders in raptor populations: evidence from rings, radio-tags and transect surveys. *Oikos* 91: 271-279.
- Kéry, M. & Schaub, M. 2012. Bayesian Population Analysis using WinBUGS, Oxford: Academic Press.
- Kovács, A., Williams, N. P. & Galbraith, C. A. 2014. Saker Falcon Falco cherrug Global Action Plan (SakerGAP), Abu Dhabi, United Arab Emirates: The Coordinating Unit of the Memorandum of Understanding on the Conservation of Migratory Birds of Prey in Africa and Eurasia (Raptors MOU).
- Kranstauber, B., Cameron, A., Weinzerl, R., Fountain, T., Tilak, S., Wikelski, M. & Kays, R. 2011. The Movebank data model for animal tracking. *Environ. Modell. Softw.* 26: 834-835.
- Lebreton, J. D., Burnham, K. P., Clobert, J. & Anderson, D. R. 1992. Modeling survival and testing biological hypotheses using marked animals: a unified approach with case studies. *Ecol. Monog.* 62: 67-118.
- Mannan, R. W. & Matter, W. J. 2004. Movements and survival of fledgling Cooper's Hawks in an urban environment. J. Raptor Res. 38: 26-34.
- Martin, J., Nichols, J. D., Kitchens, W. M. & Hines, J. E. 2006. Multiscale patterns of movement in fragmented landscapes and consequences on demography of the Snail

Kite in Florida. J. Anim. Ecol. 75: 527-539.

- McIntyre, C. L. 2005. Golden Eagles in Denali National Park and Preserve: Productivity and survival in relation to landscape characteristics of nesting territories. Ph. D. thesis, Oregon State University.
- McIntyre, C. L. 2012. Quantifying sources of mortality and wintering ranges of Golden Eagles from interior Alaska using banding and satellite tracking. *J. Raptor Res.* 46: 129-134.
- Meretsky, V., Snyder, N. F. R., Beissinger, S. R., Clendenen, S. A. & Wiley, J. W. 2000. Demography of the California Condor: implications for reestablishment. *Conserv. Biol.* 14: 957-967.
- Mertz, D. B. 1971. Life history phenomena in increasing and decreasing populations. In Patil, G. P., Pielou, E. C. & Waters, W. E. (eds.) *Statistical Ecology. II. Sampling and modeling biological populations and population dynamics:* 361-399. Pennsylvania University Park: Pennsylvania State University Press.
- Millsap, B. L. 2002. Survival of Florida Burrowing Owls along an urban-development gradient. J. Raptor Res. 36: 3-10.
- Morrison, J. 2003. Age-specific survival of Florida's Crested Caracaras. *J. Field Ornithol.*74: 321-330.
- Nasution, M. D., Brownie, C., Pollock, K. H. & Powell, R. A. 2004. The effect on model identifiability of allowing different relocation rates for live and dead animals in the combined analysis of telemetry and recapture data. *J. Agr. Biol. Envir. St.* **9**: 27-41.
- Newton, I. 1974. Changes attributed to pesticides in the nesting success of the Sparrowhawk in Britain. *J. Appl. Ecol.* 11: 95-101.

Newton, I. 1979. *Population ecology of raptors*. Berkhamsted: T. and A.D. Poyser.Newton, I. 1986. *The Sparrowhawk*. Berkhamsted: T. and A.D. Poyser.

Newton, I. 1998. Population limitation in birds. London : Academic Press.

Newton, I. 2013. Bird populations. London: Collins.

- Newton, I., Marquiss, M. & Rothery, P. 1983. Age structure and survival in a Sparrowhawk population. *J. Anim. Ecol.* **52**: 591-602.
- Newton, I. & Mearns, R. 1988. Population ecology of Peregrines in south Scotland. In Cade, T. J., Enderson, J. H., Thelander C. G. & White, C. M. (eds.) *Peregrine Falcon Populations: Their Management and Recovery:* 651-665. Boise, ID, USA: The Peregrine Fund.
- Newton, I. & Rothery, P. 1997. Senescence and reproductive value in Sparrowhawks. *Ecology* **78**: 1000-1008.
- Newton, I., Rothery, P. & Wyllie, I. 1997. Age-related survival in female Sparrowhawks Accipiter nisus. Ibis 139: 25-30.
- Newton, I. & Wyllie, I. 1992. Recovery of a Sparrowhawk population in relation to declining pesticide contamination. J. Appl. Ecol. 29: 476-484.
- Nichols, J. D., Blohm, R. J., Reynolds, R. E., Trost, R. E., Hines, J. E. & Bladen, J. P. 1991. Band reporting rates for Mallards with reward bands of different dollar values. *J. Wildl. Manage.* 55: 119-126.
- Nichols, J. D. & Hines, J. E. 2002. Approaches for the direct estimation of λ, and demographic contributions to λ, using capture-recapture data. *J. Appl. Stat.* 29: 539-568.
- Nichols, J. D., Hines, J. E., Reynolds, R. E., Blohm, R. J., Trost, R. F. & Bladen, J. P. 1995. Geographic-variation in band reporting rates for Mallards based on reward banding. J. Wildl. Manage. 59: 697-708.
- Oaks, J. L., Gilbert, M., Virani, M. Z., Watson, R. T., Meteyer, C. U., Rideout, B. A., Shivaprasad, H. L., Ahmed, S., Chaudhrt, M. J. I., Arshad, M., Mahmood, S.,

Ali, A. & Khan, A. 2003. Diclofenac residues as the cause of vulture population decline in Pakistan. *Nature* **427**: 630-633.

- Ogada, D., Shaw, P., Beyers, R. L., Buij, R., Murn, C., Thiollay, J. M., Beale, C. M.,
 Holdo, R. M., Pomeroy, D., Baker, N., Krüger, S. C., Botha, A., Virani, M. Z.,
 Monadjem, A. & Sinclair, A. R. E. 2015. Another continental vulture crisis:
 Africa's vultures collapsing toward extinction. *Conservation Letters* In press.
- Ogada, D. L., Keesing, F. & Virani, M. Z. 2012. Dropping dead: causes and consequences of vulture population declines worldwide. *Ann. NY. Acad. Sci.* **1249**: 57-71.
- Picozzi, N. 1984. Sex ratio, survival and territorial behaviour of polygynous Hen Harriers *Circus c. cyaneus* in Orkney. *Ibis* 126: 356-365.
- Pinheiro, J., Bates, D., DebRoy, S., Sarkar, D. & R Core Team (2014). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-117, <URL: http://CRAN.Rproject.org/package=nlme>.
- Pollock, K. H., Winterstein, S. R., Bunck, C. M. & Curtis, P. D. 1989. Survival analysis in telemetry studies: the staggered entry design. J. Wildl. Manage. 53: 7-15.
- Powell, L. A., Conroy, M. J., Hines, J. E., Nichols, J. D. & Krementz, D. G. 2000. Simultaneous use of mark-recapture and radiotelemetry to estimate survival, movement, and capture rates. *J. Wildl. Manage*.64: 302-313.
- Pradel, R. 1996. Utilization of capture-mark-recapture for the study of recruitment and population growth rate. *Biometrics* 52: 703-709.
- **R Core Team** (2014). *R: A language and environment for statistical computing. R Foundation for Statistical Computing.* Vienna, Austria. URL http://www.Rproject.org/.
- Ratcliffe, D. A. 1970. Changes attributable to pesticides in egg breakage frequency and eggshell thickness in some British birds. *J. Appl. Ecol.* **7**: 67-115.

Ratcliffe, D. A. 1993. *The Peregrine Falcon*, 2nd ed. Calton: Poyser.

- Rosenfield, R. N., Bielefeldt, J., Rosenfield, L. J., Booms, T. L. & Bozek, M. A. 2009. Survival rates and lifetime reproduction of breeding male Cooper's Hawks in Wisconsin, 1980 – 2005. Wilson J. Ornithol. 121: 610-617.
- Royle, J. A. & Garrettson, P. R. 2005. The effect of reward band value on mid-continent mallard band reporting rates. *J. Wildl. Manage.* **69**: 800-804.
- Rudnick, J. A., Katzner, T. E., Bragin, E. A., Rhodes, O. E. & DeWoody, J. A. 2005. Using naturally shed feathers for individual identification, genetic parentage analyses, and population monitoring in an endangered Eastern Imperial Eagle (*Aquila heliaca*) poplation from Kazakhstan. *Mol. Ecol.* 14: 2959-67.
- Sandercock, B. K. 2006. Estimation of demographic parameters from live-encounter data: a summary review. J. Wildl. Manage. 70: 1504-1520.
- Saurola, P., Stjernberg, T., Högmander, J., Koivusaari, J., Ekblom, H. & Helander, B.
 2003. Survival of juvenile and sub-adult Finnish White-tailed Sea Eagles in 19911999: a preliminary analysis based on resightings of colour-ringed individuals. In
 Helander, B. (ed.) *Sea Eagle 2000:* 155-167. Stockholm: Swedish Society for Nature
 Conservation.
- Schaub, M. & Abadi, F. 2011. Integrated population models: a novel analysis framework for deeper insights into population dynamics. J. Ornithol. 152 (Suppl. 1): S227-S237.
- Schaub, M., Aebischer, A., Gimenez, O., Berger, S. & Arlettaz, R. 2010. Massive immigration balances high anthropogenic mortality in a stable Eagle Owl population: Lessons for conservation. *Biol. Conserv.* 143: 1911-1918.
- Sergio, F., Newton, I. & Marchesi, L. 2005. Conservation: top predators and biodiversity. *Nature* **436**: 192.

Sergio, F., Newton, I. & Marchesi, L. 2008. Top predators and biodiversity: much debate,

few data. J. Appl. Ecol. 45: 992-999.

- Sergio, F., Newton, I., Marchesi, L. & Pedrini, P. 2006. Ecologically justified charisma: preservation of top predators delivers biodiversity conservation. *J. Appl. Ecol.* 43: 1049-1055.
- Sergio, F., Tavecchia, G., Blas, J., López, L., Tanferna, A. & Hiraldo, F. 2011. Variation in age-structured vital rates of a long-lived raptor: Implications for population growth. *Basic Appl. Ecol.* 12: 107-115.
- Skalski, J. R., Ryding, K. E. & Millspaugh, J. 2005. Wildlife demography: analysis of sex, age, and count data. San Diego: Academic Press.
- Smart, J., Amar, A., Sim, I. M. W., Etheridge, B., Cameron, D., Christie, G. & Wilson,
 J. D. 2010. Illegal killing slows population recovery of a re-introduced raptor of high conservation concern the Red Kite *Milvus milvus*. *Biol. Conserv.* 143: 1278-1286.
- Smith, G. D., Murillo-Garcia, O. E., Hostetler, J. A., Mearns, R., Newton, I., McGrady,
 M. J. & Oli, M. K. 2015. Demography of population recovery: survival and fidelity
 of Peregrine Falcons at various stages of population recovery. *Oecologia* 178: 391-401.
- Stahl, J. T. & Oli, M. K. 2006. Relative importance of avian life-history variables to population growth rate. *Ecol. Model.* 198: 22-39.
- Steenhof, K., Bates, K. K., Fuller, M. R., Kochert, M. N., Mckinley, J. O. & Lukacs, P.
 M. 2006. Effects of radiomarking on prairie falcons: Attachment failures provide insights about survival. *Wildlife Soc. Bull.* 34: 116-126.
- Steenhof, K., Fuller, M., Kochert, M. & Bates, K. 2005. Long-range movements and breeding dispersal of Prairie Falcons from southwest Idaho. *Condor* 107: 481-496.
 Subramanian, M. 2012. An ill wind. *Nature* 486: 310-311.

Sulawa, J., Robert, A., Köppen, U., Hauff, P. & Krone, O. 2010. Recovery dynamics and

viability of the White-tailed Eagle (*Haliaeetus albicilla*) in Germany. *Biodivers*. *Conserv.* **19**: 97-112.

- Tavecchia, G., Adrover, J., Navarro, A. M. & Pradel, R. 2012. Modelling mortality causes in longitudinal data in the presence of tag loss: application to raptor poisoning and electrocution. J. Appl. Ecol. 49: 297-305.
 - Tempel, D. J., Peery, M. Z. & Gutierrez, R. J. 2014. Using integrated population models to improve conservation monitoring: California Spotted Owls as a case study. *Ecol. Model.* 289: 86-95.
 - Tenan, S., Adrover, J., Munoz Navarro, A., Sergio, F. & Tavecchia, G. 2012.
 Demographic consequences of poison-related mortality in a threatened bird of prey. *PloS one* 7: e49187.
 - Tordoff, H. B. & Redig, P. T. 1997. Midwest Peregrine Falcon demography 1982-1995. J. *Raptor Res.* **31**: 339-346.
 - Watson, R. T., Fuller, M., Pokras, M. & Hunt, W. G. 2009. Ingestion of lead from spent ammunition: implications for wildlife and humans, Boise, ID, USA: The Peregrine Fund.
- Wellicome, T.I., Fisher, R.J., Poulin, R.G., Todd, D., Bayne, E.M. Tyler, D.T., Schmutz,
 J.K., De Smet, K. & James, P.C. 2014. Apparent survival of adult Burrowing Owls that breed in Canada is influenced by weather during migration and on their wintering. *Condor* 116: 446-458.
- Whitfield, D. P., Fielding, A. H., McLeod, D. R. A. & Haworth, P. F. 2004. Modelling the effects of persecution on the population dynamics of Golden Eagles in Scotland. *Biol. Conserv.* 119: 319-333.
- Whitfield, D. P., Fielding, A. H., McLeod, D. R. A. & Haworth, P. F. 2008. A Conservation Framework for Golden Eagles: Implications for their Conservation and

Management in Scotland: Scottish Natural Heritage Commissioned Report No.193 (ROAME No. F05AC306).

- Williams, B. K., Nichols, J. D. & Conroy, M. J. 2002. Analysis and management of animal populations, San Diego, USA: Academic Press.
- Wink, M., Staudter, H., Bragin, Y., Pfeffer, R. & Kenward, R. 1999. The use of DNA fingerprinting to estimate annual survival rates in the saker falcon (*Falco cherrug*). J. Ornith. 140: 481-489.
- Wyllie, I. & Newton, I. 1991. Demography of a increasing population of Sparrowhawks. J. Anim. Ecol. 60: 749-766.
- Zelenak, J. R., Rotella, J. J. & Harmata, A. R. 1997. Survival of fledgling Ferruginous Hawks in northern Montana. *Can. J. Zoolog.* 75: 152-156.
- Zens, M. S. & Peart, D. R. 2003. Dealing with death data: individual hazards, mortality and bias. *Trends Ecol. Evol.* 18: 366-373.

Zuur, A. F., Ieno, E. N., Walker, N. J., Saveliev, A. A. & Smith, G. M. 2009. *Mixed effects models and extensions in Ecology with R.* New York: Springer.

SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article **Table S1.** Survival estimates for diurnal raptors and owls.

FIGURE LEGEND

Figure 1. The relationship between body mass and adult survival in diurnal raptors, owls and both diurnal raptors and owls combined. See Table 2 for a detailed description of data.

Table 1. Summary of methods for estimating survival appropriate to different data types, with their pros and cons.

Method (data type)	Analytical	Estimate of	Advantages	Disavantages
	procedures	survival		
1. Ring-	Informal based on	True, usually.	Less expensive than CMR or radio-	Informal models do not allow for
recovery	age ratios among	Apparent, if	tracking studies	variations of recovery and
	recoveries, or formal	recovery occurs		reporting rates.
	using Seber and	only in local areas	Offer a cost-effective way of	Madala and former for a second state
	Brownie		monitoring raptor vital rates over the	Models with rewest assumptions
	parameterizations of		long-term	both adult and young birds. Adults
	ring recovery models			are often difficult to catch or re-
				observe.
				Low recovery and/or reporting
				rates often lead to inadequate
				sample size
2. Observationa	Informal methods:	Bias can be in	Cheapest and easiest method to	Several assumptions are difficult
l data with	territory turnover, age	either direction,	estimate survival	to meet, including: equal
and without	ratio	depending on		detectability for age classes; stable
individual		methodological		age distribution; stationary
identification		details		population

This article is protected by copyright. All rights reserved.

	3.	Capture- Mark- Recapture or Resight (CMR)	Cormack-Jolly-Seber, multistate models	Apparent	Less costly and more efficient sampling than radio-tacking studies. Allows monitoring of a large number of individuals simultaneously	Estimates do not distinguish between permanent emigration and death Does not distinguish between death and emigration Limited use for raptor species that exhibit low site fidelity
60	4.	Joint live encounters – ring recoveries	Burnham and Barker models	True	Distinguish between permanent emigration and death by estimating fidelity and survival rates spearately	Multiple sources of information are unlikely to be available for many species. Sample size often small
Cent	5.	Radio- tracking	Kaplan-Meier, Cox Proportional Hazard or discrete time analogues, CMR models (if detection rates are <1)	Can be true – depends on whether birds leave the area, and become immune to sampling efforts	Distinguishes between death and permanent emigration Allows cause-specific mortality analysis Provides additional ecological	High financial cost and time- consuming sampling Short lifespans of most radio transmitters Possible effects of radio-tags on

				information (e.g., movement, space	survival
				and resource use patterns)	
					Potential problems with non-
					independence of censoring and
					fate
	6. Multiple data	Integrated population	True, usually.	Distinguishes between permanent	Requires multiple data types,
	types (e.g.,	models	Apparent, if data	emigration and death, given	which may not available for many
	time series of		permitting	appropriate data	species
	counts, live		separation of loss		
	encounter,		into emigration and	Permit estimation of important	Bias induced by inappropriate
	ring		death are not	demographic parameters, including	modelling of one parameter can
	recoveries		available	survival and population growth rates	translate into biases in multiple
	recoveries,		available		parameter estimators
	radio-			Estimates generally more precise	
Y	tracking)				Requires strong statistical and
					programming skills

Table 2. Annual adult survival estimates (SE in parentheses when available) and body mass for 30 species of diurnal raptors and nine species of owls used to examine the relationship between body mass and adult survival. When survival was reported for males (M) or females (F), adult body mass is given for that sex. When survival is reported for both sexes (B), average male and female body mass is given for males and females. Estimation methods are described in Table 1, and body masses are from Dunning (1992, 2007). Notes indicate specific circumstances (if any) under which survival rates were estimated.

	Species	Sex	Mass (g.)	Annual adult survival (SE)	Method	Reference	Notes
	A. Diurnal raptors						
	Andean Condor Vultur gryphus	В	11300	0.94	5	Temple & Wallace (1989)	
	Osprey Pandion haliaetus	В	1505.5	0.64	5	Klaassen et al. (2014)	
	Egyptian Vulture Neophron percnopterus	В	2082	0.75 (0.02)	3	Grande et al. (2009)	non-breeding
		В	2082	0.833 (0.022)			breeding
Ð	European Honey-buzzard <i>Pernis apivorus</i>	В	758	0.813 (0.023)*	3	Bijlsma <i>et al.</i> (2012)	multiple adult ages
Ũ	Eurasian Griffon <i>Gyps fulvus</i>	В	7436	0.987 (0.006)	3	Sarrazin <i>et al.</i> (1994)	released birds

Golden Eagle Aquila chrysaetos	В	4263.5	0.896 (0.371)	5	Hunt et al. (1998)	windfarm, breeders
	В	4263.5	0.909 (0.025)		Hunt <i>et al.</i> (2002)	windfarm, breeders
Bonelli's Eagle A. fasciata	В	2000	0.87	3	Hernández-Matías <i>et al</i> . (2011)	
Eurasian Sparrowhawk Accipiter nisus	F	325	0.586 (0.103)*	3	Newton <i>et al.</i> (1997)	multiple adult ages
	F	325	0.567 (0.091)*			
	F	325	0.656 (0.089)*			
	F	325	0.574 (0.081)*			
Northern Goshawk A. gentilis	М	1137	0.83 (0.09)	5	Kenward (1999)	
	F	912	0.83 (0.09)			
Western Marsh Harrier Circus aeruginosus	В	711.5	0.56	5	Klaassen et al. (2014)	
Hen Harrier C. cyaneus	М	430	0.72 (0.07)	3	Picozzi (1984)	
ý	F	430	0.90 (0.03)		× /	
					Etheridge <i>et al</i> .	
	F	430	0.397	3	(1997)	grouse moor
	F	300	0.778			other moor
Montagu's Harrier C. pygargus	В	315.5	0.59	5	Klaassen et al. (2014)	

Red Kite Milvus milvus	В	1080	0.87 (0.12)	3	Smart <i>et al.</i> (2010)	wild-bred birds
(В	1080	0.85 (0.20)			released birds
	В	1080	0.77 (0.03)	3	Tavecchia <i>et al.</i> (2012)	
	В	1080	0.955	6	Tenan <i>et al.</i> (2012)	poison free
Black Kite <i>M. migrans</i>	В	567	0.789 (0.008)	3	Sergio et al. (2011)	
White-tailed Eagle Haliaeetus albicilla	В	4793	0.715 (0.046)	3	Saurola <i>et al.</i> (2003)	
	В	4793	0.966 (0.014)	3	Evans et al. (2009)	wild-bred birds
	В	4793	0.942 (0.022)			released birds
Bald Eagle <i>H. leucocephalus</i>	М	5350	0.86 (0.05)	5	Bowman <i>et al.</i> (1995)	
	F	4130	0.90 (0.04)			
	В	4740	0.766*	5	Harmata <i>et al.</i> (1999)	multiple adult ages
Snail Kite Rostrhamus sociabilis	В	420	0.861 (0.034)	3	Bennetts et al. (1999)	Cormack-Jolly-Seber
	В	420	0.822 (0.034)	3		multi-strata models
	В	420	0.894 (0.029)	5		radio telemetry
Swainson's Hawk Buteo swainsoni	В	958.5	0.843 (0.019)	4	Schmutz <i>et al.</i> (2006)	
Galapagos Hawk	В	1099	0.94	3	Rivera Parra <i>et al.</i> (2012)	before goat eradication
2. 6	В	1099	0.84		()	after goat eradication
	Red Kite Milvus milvusBlack Kite M. migransWhite-tailed Eagle Haliaeetus albicillaBald Eagle H. leucocephalusSnail Kite Rostrhamus sociabilisSwainson's Hawk Buteo swainsoniGalapagos Hawk B. galapagoensis	Red Kite Milvus milvusB B B BBlack Kite M. migransBBlack Kite M. migransBWhite-tailed Eagle Haliaeetus albicillaBBBBald Eagle H. leucocephalusMF BBSnail Kite Rostrhamus sociabilisBSwainson's Hawk Buteo swainsoniBSwainson's Hawk Buteo swainsoniBSqalapagos Hawk B, galapagoensisBBB <tr< td=""><td>Red Kite Milvus milvusB B 1080 B1080 B 1080B1080B1080B1080B1080Black Kite M. migransB300B</td><td>Red Kite Milvus milvusB B 10801080 0.85 (0.20) B0.85 (0.20) 0.85 (0.20) BB10800.955Black Kite M. migransB5670.789 (0.008)White-tailed Eagle Haliaeetus albicillaB47930.715 (0.046) 0.942 (0.022)Bald Eagle H. leucocephalusM53500.86 (0.05)F4130 47400.90 (0.04) B60.0766*Snail Kite Rostrhamus sociabilisB420 4200.822 (0.034) BB420 4200.894 (0.029)Swainson's Hawk Buteo swainsoniB1099 B0.94 BB1099 10990.84</td><td>Red Kite Milvus milvusB B B1080 10800.87 (0.12) 0.85 (0.20) 0.77 (0.03)3B10800.9556Black Kite M. migransB5670.789 (0.008)3White-tailed Eagle AlbicillaB47930.715 (0.046)3B47930.966 (0.014) 0.942 (0.022)3Bald Eagle H. leucocephalusM53500.86 (0.05)5F4130 47400.900 (0.04) 0.766*5Snail Kite Rostrhamus sociabilisB420 4200.822 (0.034) 0.894 (0.029)3Swainson's Hawk Buteo swainsoniB958.50.843 (0.019)4Galapagos Hawk B B10990.943B10990.843</td><td>Red Kite Milvus milvus B 1080 0.87 (0.12) 3 Smart et al. (2010) B 1080 0.77 (0.03) 3 Tavecchia et al. (2012) B 1080 0.955 6 Tenan et al. (2012) B 1080 0.955 6 Tenan et al. (2012) Black Kite M. migrans B 567 0.789 (0.008) 3 Sergio et al. (2011) White-tailed Eagle Haliaeetus B 4793 0.715 (0.046) 3 Saurola et al. (2003) albicilla B 4793 0.966 (0.014) 3 Evans et al. (2009) Bald Eagle M 5350 0.86 (0.05) 5 Bowman et al. (1995) H. leucocephalus F 4130 0.90 (0.04) 3 Bennetts et al. (1999) Snail Kite B 420 0.822 (0.034) 3 Bennetts et al. (1999) Snail Kite sociabilis B 420 0.822 (0.034) 3 Bennetts et al. (1999) Swainson's Hawk B 958.5 0.843 (0.019) 4 Schmutz et al. (2006) Galapagos Hawk B 1099 0.94</td></tr<>	Red Kite Milvus milvusB B 1080 B1080 B 1080 B1080B1080B1080B1080Black Kite M. migransB 300 B	Red Kite Milvus milvusB B 10801080 0.85 (0.20) B0.85 (0.20) 0.85 (0.20) BB10800.955Black Kite M. migransB5670.789 (0.008)White-tailed Eagle Haliaeetus albicillaB47930.715 (0.046) 0.942 (0.022)Bald Eagle H. leucocephalusM53500.86 (0.05)F4130 47400.90 (0.04) B60.0766*Snail Kite Rostrhamus sociabilisB420 4200.822 (0.034) BB420 4200.894 (0.029)Swainson's Hawk Buteo swainsoniB1099 B0.94 BB1099 10990.84	Red Kite Milvus milvusB B B1080 10800.87 (0.12) 0.85 (0.20) 0.77 (0.03)3B10800.9556Black Kite M. migransB5670.789 (0.008)3White-tailed Eagle AlbicillaB47930.715 (0.046)3B47930.966 (0.014) 0.942 (0.022)3Bald Eagle H. leucocephalusM53500.86 (0.05)5F4130 47400.900 (0.04) 0.766*5Snail Kite Rostrhamus sociabilisB420 4200.822 (0.034) 0.894 (0.029)3Swainson's Hawk Buteo swainsoniB958.50.843 (0.019)4Galapagos Hawk B B10990.943B10990.843	Red Kite Milvus milvus B 1080 0.87 (0.12) 3 Smart et al. (2010) B 1080 0.77 (0.03) 3 Tavecchia et al. (2012) B 1080 0.955 6 Tenan et al. (2012) B 1080 0.955 6 Tenan et al. (2012) Black Kite M. migrans B 567 0.789 (0.008) 3 Sergio et al. (2011) White-tailed Eagle Haliaeetus B 4793 0.715 (0.046) 3 Saurola et al. (2003) albicilla B 4793 0.966 (0.014) 3 Evans et al. (2009) Bald Eagle M 5350 0.86 (0.05) 5 Bowman et al. (1995) H. leucocephalus F 4130 0.90 (0.04) 3 Bennetts et al. (1999) Snail Kite B 420 0.822 (0.034) 3 Bennetts et al. (1999) Snail Kite sociabilis B 420 0.822 (0.034) 3 Bennetts et al. (1999) Swainson's Hawk B 958.5 0.843 (0.019) 4 Schmutz et al. (2006) Galapagos Hawk B 1099 0.94

Ferruginous Hawk B. regalis	В	1468.5	0.708 (0.024)	4	Schmutz <i>et al</i> . (2008)	
Eurasian Buzzard <i>B. buteo</i>	В	875	0.88 (0.03)	5	Kenward <i>et al.</i> (2000)	
Northern Crested Caracara	М	1220	0.876 (0.003)	5	Morrison (2003)	
Curucuru cheriway	F	1117	0.906 (0.002)			
Lesser Kestrel Falco naumanni	В	152.5	0.706*	3	Hiraldo et al. (1996)	multiple annual estimates
	В	152.5	0.67 (0.06)	3	Prugnolle <i>et al.</i> (2003)	
	В	152.5	0.72 (0.015)	3	Serrano <i>et al.</i> (2005)	large colonies
	В	152.5	0.653 (0.019)			medium colonies
	В	152.5	0.647 (0.019)			small colonies
	В	152.5	0.718 (0.013)	3	Mihoub et al. (2010)	
Mauritius Kestrel F. punctatus	В	119.5	0.782	3	Nicoll <i>et al.</i> (2003)	
	В	119.5	0.8 (0.04)	3	Nicoll et al. (2004)	hacked birds
	В	119.5	0.8 (0.04)			fostered birds
	В	119.5	0.75 (0.03)			wild-bred birds
American Kestrel F. sparvarius	М	120	0.75 (0.05)	3	Hinnebusch <i>et al.</i> (2010)	
	F	111	0.74 (0.04)			

	Sooty Falcon						
	F. concolor	В	235	0.656 (0.069)	3	McGrady <i>et al.</i> (2015)	
	Aplomado Falcon F. femoralis	В	341.5	0.91	3	Brown et al. (2006)	wild-bred breeder
		В	341.5	0.91			hacked breeder
		В	341.5	0.872			wild-bred non-breeder
		В	341.5	0.303			hacked non-breeder
	Merlin F. columbarius	В	117	0.62 (0.11)	3	Lieske et al. (2000)	
	Prairie Falcon F. mexicanus	F	908	0.87	3	Steenhof et al. (2006)	without transmitters
		F	908	0.49			with transmitters
	Peregrine Falcon F. peregrinus	F	959	0.788 (0.031)	3	Gould & Fuller (1995)	
		В	598	0.725 (0.023)	3	Johnstone (1997)	
1.0		Μ	598	0.765 (0.038)	3		
		F	959	0.701 (0.032)	3		
		В	814.5	0.859 (0.025)	4	Kauffman <i>et al.</i> (2003)	
		В	814.5	0.800 (0.054)	4	Craig <i>et al.</i> (2004)	
		М	598	0.73 (0.02)	3	Franke <i>et al.</i> (2011)	
		F	959	0.73 (0.02)		× /	
		В	824.5	0.81	3	Faccio <i>et al.</i> (2013)	
		В	650	0.852	6	Altwegg <i>et al.</i> (2014)	
		В	697.5	0.810 (0.034)	4	Smith <i>et al.</i> (2015)	

B. Owls

	B. Owls						
	Common Barn Owl	В	403	0.720 (0.044)	4	Altwegg et al. (2003)	
		В	403	0.570 (0.023)	4	Altwegg et al. (2006)	
	Snowy Owl Bubo scandiacus	F	2279	0.923 (0.057)	5	Therrien et al. (2012)	best case
		F	2279	0.852 (0.07)			worst case
	Eagle owl <i>B. bubo</i>	В	2686	0.606	6	Schaub <i>et al.</i> (2010)	
	Tawny Owl Strix aluco	В	475	0.755 (0.026)*	4	Francis & Saurola (2002)	capture-mark-recapture models
	D	В	475	0.760 (0.020)*			merarchical model capture-mark-recapture models.
+		В	475	0.683 (0.035)*			poor vole years hierarchical model, poor vole
		В	475	0.727 (0.039)			years capture-mark-recapture models,
		В	475	0.736 (0.035)*			medium vole years hierarchical model, medium
		В	475	0.780 (0.035)			vole years capture models,
		В	475	0.846 (0.025)*			good vole years hierarchical model, good vole
		В	475	0.780 (0.036)			years

e)						
	В	475	0.739 (0.059)*	3	Karell et al. (2009)	experienced breeders **
	В	475	0.570 (0.070)*			first-time breeders**
	Μ	524	0.79 (0.03)	3	Millon et al. (2010)	
	F	426	0.86 (0.03)*			
	В	475	0.710 (0.029)	3	Pavón-Jordán <i>et al.</i> (2013)	two yrs old and >3 yrs old
Spotted Owl S. occidentalis	F	646	0.814 (0.050)	3	Seamans <i>et al.</i> (1999)	
	F	646	0.832 (0.029)			
	В	0.86	0.86	3	Bond et al. (2002)	
					Zimmerman et al.	
	В	606	0.82 (0.03)	3	(2007)	
	М	566	0.942(0.020)	2	Blakesley <i>et al.</i>	
		500	0.843(0.020)	3	(2010)	
	Г	040 566	0.811(0.021) 0.840(0.017)			
	F	500 646	0.840(0.017) 0.848(0.016)			
	M	0 4 0 566	0.848 (0.016)			
	F	500 646	0.890 (0.016)			
	M	566	0.848 (0.015)			
	F	646	0.848 (0.015)			
	M	566	0.819 (0.013)	3	Forsman <i>et al.</i> (2011)	
	F	646	0.819 (0.013)	-		
	М	566	0.863 (0.008)			
	F	646	0.859 (0.009)			
	М	566	0.851 (0.007)			
	F	646	0.853 (0.007)			
	М	566	0.864 (0.010)			

	F	646	0.865 (0.010)			
	М	566	0.857 (0.013)			
	F	646	0.854 (0.014)			
	Μ	566	0.847 (0.080)			
	F	646	0.848 (0.080)			
	М	566	0.846 (0.009)			
	F	646	0.844 (0.009)			
	Μ	566	0.852 (0.014)			
	F	646	0.828 (0.016)			
	Μ	566	0.844 (0.018)			
	F	646	0.841 (0.019)			
	М	566	0.853 (0.010)			
4	F	646	0.851 (0.010)			
	М	566	0.857 (0.008)			
	F	646	0.856 (0.008)			
	М	566	0.847 (0.040)*	3	Ganey et al. (2014)	annual estimates 2004-2009
	F	646	0.859 (0.036)*			annual estimates 2004-2009
	В	606	0.828	6	Tempel <i>et al.</i> (2014)	
Ural Owl S uralensis	В	785	0.80 (0.016)	3	Pavón-Jordán et al.	
oral own 5. araceisis	Б	705	0.00 (0.010)	5	(2013)	
Little Owl Athene neeting	М	164	0.651 (0.042)	2	Schemb at al. (2006)	
Little Owi Ainene nociua		104	0.031(0.043)	3	Schaub <i>et ut</i> . (2000)	
	Г	104	0.010(0.042)			
	IVI E	104	0.075(0.033)			
	Г Г	104	0.074(0.020)			
	Г М	164	0.087 (0.068)			
	M	164	0.740 (0.057)			

	F	164	0.659 (0.051)			
					Le Gouar et al.	
	В	164	0.753 (0.019)	3	(2011)	
	В	164	0.66	5	Thorup <i>et al.</i> (2013)	
	14	146	0.01 (0.04)	2		
Burrowing Owl A. cunicularia	М	146	0.81 (0.04)	3	Millsap (2002)	highly developed area
	F	156	0.69 (0.04)			highly developed area
	Μ	146	0.62 (0.05)			moderately developed area
	F	156	0.69 (0.04)			moderately developed area
	В	151	0.545	4	Barclay et al. (2011)	
	В	151	0.71			increasing population
	В	151	0.465			decreasing population
	М	146	0.44 (0.05)	3	Wellicome <i>et al.</i> (2014)	
	F	156	0.21 (0.03)		(2014)	
Boreal Owl Aegolius funereus	В	142	0.46	5	Hayward <i>et al.</i> (1993)	

*geometric mean of different estimates (see text) **during periods of low, increasing and decreasing population

