Dear authors,

based on the reports of three external reviewers, my own reading of your paper, and the help of Camille Jolivel for an extra careful reading of your manuscript, I would be happy to recommend your paper for PCI Ecology, pending some modifications and corrections.

The three reviewers, myself and Camille Jolivel have appreciated a lot your paper. The topic discussed is very interesting because CMR-based methods consider heavy assumptions, which could have large consequences on the final results. It is therefore interesting to see to what extent these assumptions are justifiable. The particular impact of mark-loss on estimation biases is scarce in the literature and is worth being studied further as you show here, especially because mark losses can have counter-intuitive effects.

Dr. Sylvain Billiard for PCI Ecology

General evaluation and major comments

1. Better define and give a precise formulation of your model, more in line with how such models are presented in the literature. I strongly suggest to follow reviewer’s 1 advice to dedicate a whole section on the model, its formulation, hypothesis, link with previous literature, etc.

2. All reviewers, myself and Camille Jolivel felt that a better use of supp. information should be made. Actually, they are too extensive, some are certainly worth to include in the main text, a suggestion by reviewer 3 to provide a tentative of synthesis of some supp. mat. by focusing on a specific case and show estimates in real parameter space rather than metrics summarising the biases (for a better understanding of the effect and direction of the biases due to mark loss). Below, I also suggest another option to include some of your results in the main text.

3. I felt a bit frustrated that a very nice work of data compilation has been done, but which is maybe underutilised. It might be worth describing and discussing a bit some general patterns observed in Fig. S1 supp. information 1.

4. As several reviewers pointed out, I found interesting the conclusion and proposition that double mark should be used. I wonder if it could be possible to have a quantitative assessment of that point with the methodological framework developed by the authors: compare biases with a single vs. two marks and evaluate the quantitative gain of estimation.
Additional comments (with the help of Camille Jolivel)

- Abstract: Most of the information given are about the simulation studies, especially sections 2., 3. and 4. The data analysis on bats is drown out in the abstract. We suggest to compile all information on bats in a 5th section.
- Keywords: We suggest to remove from the keywords list the words already in the title of the paper.
- Introduction: Italicise all parameters and variables, do not italicise numbers in quotes; Homogenise the font for quotes.
- Introduction: Suggestion of reorganisation of the first paragraph: Capture-mark-recapture (CMR) methods have [...] is the loss of marks (see Supported Information 1, Table S2). Mark loss has two consequences [...] which is called “recycling”. Local survival is the product of true survival etc.
- L78: Explain the two parts of the equation
- L100-101: Homogenise the use of brackets for citations "(Nichols & Kendall, 1995 ; see also Cam 2009 ..."
- L91 vs. L105: Incoherence between ”has not been explored” vs. ”is scarce”
- Material and method: You might improve the accuracy and the information content of the title of the sections. For instance: change “real data analysis” with “Application on a bats dataset”; “Computation details” is not sufficiently different to “data simulation”. A title containing what is within this section could be “Estimation procedures and assessments”.
- L130: Consider explaining more precisely what you call “realistic”. You might rather mean “representative”? 
- L133: It is a bit odd to cite Fig. 1.b before Fig. 1.a
- L140: remove comma after “Reptilia”
- L145: It is not quite clear how you translated your data survey in these four scenario. Consider explaining more accurately. In addition, it might be worth making explicit the features you chose to neglect. Also discuss whether or not, by splitting the range of observation into four scenarios, your analysis is still adequate for any kind of species considered.
- L169: add notation for the “mark loss probabilities” noted “ptl” in Fig 1.a
- L171: Not sure why 50 datasets here, and only ten in FigS1 supp Inf 2?
- L177: It would be more appropriate to call $R_{i,t}$ a probability rather than a rate because rate refers to a speed, you could consider calling $R_{i,t}$ the retention event probability.
- L177: Not clear whether the “mark loss probabilities” = 1 - “retention probability“?
- L177: Not easy to understand that $j$ in... means over the different situation. The model should be better formalised.
- L182
- L186: “recycled individuals” is defined here, whereas the term appears several times before. Consider defining explicitly what “recycled” means as soon as necessary.
- L200: remove comma after “both”
- L219: delete extra dot
- L224: delete extra dot
- L234: Add missing dot
• L237: Consider adding some general and basic literature about JAGS, Gibbs, etc. for people that would not be familiar with these methods. e.g. Kruschke, J. (2014). Doing Bayesian data analysis: A tutorial with R, JAGS, and Stan.

• L238: Give the meaning of the acronym MCMC

• L242: "Chain convergence was assessed with the Gelman-Rubin statistic denoted R-hat”

• L243: “converge” → “convergence”

• L247: “expected”, rather say “assumed”? 

• L257: Only figure S43 is mentioned here, a series of figure is expected I guess.

• Results: Homogenise notations of states. L277 “D” vs. L282 A and C.

• L296: I feel a little frustrated that results in fig. S4-S11 are given in supp. mat. only. I think it would be worth moving these figures in the main text by providing a synthetic figure as a matrix as Fig 2 and 3.

• Discussion

• Double check the reference in the literature section, there are some mistakes and typos here and there (examples:)


• Figures: Consider splitting Figure 1 into two figures. Fig1 1a and 1b show information different enough to justify a split. In addition, it would save room for more explanation of each figure within the caption. Since these figures are at the core of the paper, it might be worth to make them as informative as possible.

• If you chose not to split the figure into two, then move down “(b)” to the correct position.

• Consider adding the meaning of notations “pc”, “pr”, “pd” in the main text

• Table 1: “Adultes” → “Adults” ; I suspect an inconsistency in notation: $p_A, p_B, ...$ might be $p_{r_A}, p_{r_B}, ...$ as the probability of re-sighting is $p_r$ in Fig 1a ; To be checked but I think “long-live” and “short-live” should be replaced by “long-lived species” and “short-lived species” ; The parameters for the Uniform and Normal distribution should be defined ($N(a,b)$ is the normal distribution with mean $a$ and variance $b$ ; Not sure what does mean the $N(−0.3$ between brackets? A range? A subtraction?

• Fig.1a: I guess the state at time $t + 1$ should be $j$ and not $i$ since it has changed with probability $\psi$ ; Move down the “(b)” ; Fig 1b: I would replace “Scenario 2” by its meaning “Long-lived species” ; Give explicit reasons in the caption why there are fluctuations across years in detection and survival probability in sub-figures.

Reviews

Please find hereafter the reports of three external reviewers.
Reviewer’s 1 report.
Review of Mark loss can strongly bias demographic rates in multi-state models: a case study with simulated and empirical datasets

May 2, 2022

Dear authors, I enjoyed reading your paper very much. Your work is original in that no one has ever assessed the bias in transition parameters due to tag-loss in capture-recapture models. I have a few comments that I hope you’ll find useful. Regards.

Main comments

• You have done an impressive work in gathering survival estimates and the magnitude of tag loss for hundreds of species. Congratulations on that. Tables S1 and S2 are awesome!

• I feel like lines 71-83 are too technical for an introduction. I would move this paragraph to Material and methods, and explain the problem with words here.

• Line 265: I find the use of ROPE disturbing because from the beginning of the paper we expect an assessment of bias. Why not simply consider posterior means/medians and compute bias in the frequentist way? I’m not asking to drop the ROPE metric, but to add a more traditional measure of bias (hopefully the simulations were saved and it will not take long).

• You’re missing several important references. First, multistate models should not be called multistate CJS models. R. Cormack, G. Jolly and G. Seber developed models with a single alive state. Multistate models were developed by Neil Arnason and Carl Schwarz in a series of papers, and multistate models are sometimes referred to as the Arnason-Schwarz model. As of terminology, I guess multistate models or Arnason-Schwarz models do the job. The references that need to be cited are:


Second, about tag-loss, there are two papers that need to be included in your paper:


• Line 235: I’ve had a hard time identifying the model with tag-loss you’re using. If I understand correctly, it is specified with Eqn 1. This equation should come with the relevant literature, unless you’ve developed this model yourself. I recommend having a specific section describing both models with and without tag loss, which are common to both the simulations and case study, so that the reader can easily go back to it. In this new section, you should make it clear how the model with tag-loss you’re using differs from existing models with double-marking. This new section could also include the literature review you do earlier in the paper, but if not, you should refer to it.

Minor comments

• Lines 31-33: I find the statement “little attention has been paid to the effect of common violations of the CMR model assumptions” a bit unfair. I understand that the abstract has to be punchy and attractive, but there are a lot of papers out there on the issues of heterogeneity, incorrect state assignment to individuals, and more broadly goodness-of-fit testing for multistate models. I would tone down this statement a little, and simply write that the tag-loss issue is tackled.

• I noticed that you use both multi-state and multistate, I would homogenise throughout the paper.

• Line 99: Bugs is not a software program, you mean WinBUGS, JAGS, OpenBUGS, STAN, NIMBLE, etc. I would also keep MARK and E-SURGE for user-friendly software here. Please, add the relevant references too, folks have put a lot of efforts in developing these pieces of software, and their dedication has served the community well, the least we can do is to acknowledge their work.

• Sometimes you have initials for first names in the cited references. I guess we do not need them in the text, only in the list of references.

• Lines 104-108: You do a great job at reviewing the literature on tag loss in CJS models, and the applications of multistate models, but here we’re missing what you expect to be new and/or different from the existing studies in terms of bias in demographic parameters. Actually, you have these predictions diluted in the paragraph lines 116-125, which is more about a specific case study. The general predictions should come with the simulation study.

• Why going for a Bayesian approach? You don’t have random effects in the simulations, nor do you have prior information to incorporate. Also, bias is kind of a frequentist notion, it would have been more natural to go for maximum likelihood estimation. Plus, this would have allowed for more than 50 simulated datasets (for each scenario or combination of parameters), which I guess you had to keep under a limit due to the computational burden of MCMC methods. Estimation of bias requires more simulations in general. I’m not asking to go for a frequentist approach, just to provide the reader with some justifications.

• Line 132: What about the dead state? It is absorbing too ;-)

• Lines 125-126: Not sure there is a formal definition of bias in posterior distribution, isn’t it?

• Line 193: What does ‘To illustrate the simulations’ mean? Some readers would be perfectly happy with simulations only. The case study brings less generality, because you don’t know the truth (at least only 66% of it). I like the case study, it just needs to be better introduced: What does it add to the simulations?

• For the bat case study, you need to perform goodness-of-fit tests and cite the relevant literature. Bias in parameter estimates could be caused by transience or trap-dependence issues.

• Line 224: Bias on which parameter(s)?

• Lines 243-248: It is difficult to follow you blindly here without knowing for which parameters convergence was not reached. Actually, with only 50 simulations, we might expect some ‘bias’ in the results due only to lack of convergence, not tag loss. I encourage you to be more specific.
• Line 237: That’s an old version of R ;-) I recommend updating JAGS, JagsUI and R.

• Line 274: Do we care about detection being biased?

• Lines 298-299: I am not sure I follow, it’s one thing to quantify bias on survival, it is another to demonstrate bias on the relationship between a covariate and survival. You might have a bias in survival, but if this bias is the same for all individuals or time intervals, then I guess there is no problem to assess the effect of individual or temporal covariate. Am I missing something?

• Lines 311-328: The results are discussed with no reference whatsoever to the existing literature. What was found in other papers? Mobilizing the existing literature and comparing your results to previous findings will help you to emphasize what’s new and original in your work.

• Lines 331-332: I’ve probably missed it, how can you be sure that you quantify bias in the case study with only two third of the individuals for which you know they’ve lost their tag?

• Line 345: Dispersal, not dispersion (I think).

• Lines 402-404: I applaud the authors for their efforts of making data and code available. I would deposit the code and data on GitHub/GitLab for versioning and also to make it easy to re-use your code (copying and pasting from a PDF to R can produce funky behaviors sometimes).

• Legend of Table 1: You write ‘U=Univariate distribution)’, you mean ‘Uniform distribution’ I guess. I also find it unclear the meaning of the minus something you have in the short-life species column. A Uniform(a,b) - 0.5 is Uniform(a-0.5,b-0.5), and Normal(mu,sigma)-0.3 is Normal(mu-0.3,sigma). Did you simulate directly from these ‘translated’ distributions, or did you simulate from Uniform(a,b) and Normal(mu,sigma) then substract something?

• Figure 1: Nice work!

• Figures 2 and 3: These figures cannot be read independently of the main text. I would remind the reader in the legend what the scenarios are, and what the numbers and letters on the axes refer to. Also I’d define ROPE.

• Line 266: I find ‘for most parameters in most situations’ to be a vague statement. I encourage you to provide some quantitative information so that the reader can make a judgment by herself/himself.

• Figure S2: I am not sure the relationship will remain significant once you account for uncertainty in both adult and juvenile survival probabilities. You’re doing statistics on statistics here.
Reviewer 2’s report.

The manuscript presents a simulation and analysis of real data, assessing the bias due to tag loss in parameter estimates from multi-state CMR models. This is a timely contribution, as the effects of tag loss in CMR models have received some attention in the literature, but not in the case of multi-state models. I do not have the technical expertise to evaluate the correctness of the code presented (I trust other reviewers will do that), but I found the text clear and generally easy to understand what was done. The introduction is possibly the part of the manuscript that could do with a bit more work. Some sections, such as between lines 62-67 could use more references. The results and figures in the manuscript are clear, although I found that the supporting information is quite extensive and difficult to follow. I agree with the interpretation of the results, that tag loss is relevant and should be taken into account in CMR studies. To me it was also surprising that it biased the transition probabilities more strongly than survival estimates. The need for double marking is also an important message that the authors properly emphasize.
Reviewer’s 3 report

The authors created an extensive simulation experiment to study the effect of mark-loss on bias of parameter estimates in multistate capture-recapture models. Overall the author did a good job illustrating the fact that mark-loss can induce substantial parameter bias in multistate models. Moreover, this bias can present itself in an unintuitive fashion due to the complexity of the model and the interactions of the parameters in the likelihood.

Although it is strictly a personal preference, the article might appeal to the ecological community that uses these models if the authors illustrated some of these bias effects on the real scale of the parameters in the main portion of the paper (not in an appendix). I.e., for perhaps with just the bat analysis, the authors could create a figure with, say survival or transition probabilities under each model. That way users can see the effects in real terms, rather than EMD or ROPE metrics that don’t have a meaningful interpretation in real parameter space.

In addition to my overall comments, I have attached an annotated pdf with more specific comments and questions.
Titles: Mark loss can strongly bias demographic rates in multi-state models: a case study with simulated and empirical datasets

Authors

Frédéric Touzalin$^{1,2}$, Eric J. Petit$^3$, Emmanuelle Cam$^4$, Claire Stagier$^1$, Emma C. Teeling$^1$, Sébastien J. Puechmaille$^{5,6,7}$

Affiliations

$^1$School of Biology and Environmental Science, Science Centre West, University College Dublin, Dublin, Ireland.
$^2$Bretagne Vivante-SEPNB, Brest, France.
$^3$DECOD (Ecosystem Dynamics and Sustainability), INRAE, Institut Agro, Ifremer, Rennes, France.
$^4$Université de Bretagne occidentale, Brest, LEMAR, CNRS, IRD, Ifremer, F-29280 Plouzane, France.
$^5$Zoological Institute and Museum, University of Greifswald, Greifswald, Germany.
$^6$ISEM, University of Montpellier, CNRS, IRD, Montpellier, France.
$^7$Institut Universitaire de France, Paris, France.

Correspondence

Frédéric Touzalin
Email: fredtouzalin@gmail.com
Abstract

1. The development of methods for individual identification in wild species and the refinement of Capture-Mark-Recapture (CMR) models during the past decades has greatly improved the assessment of population demographic rates to answer ecological and conservation questions. In particular, multistate models, with their flexibility for the analysis of complex study systems, have become popular in the ecological community. However, despite the extensive use of these models, little attention has been paid to the effect of common violations of the CMR model assumptions, such as mark loss and the often-associated recycling of remarked individuals.

2. To explore this knowledge gap we used a wide range of simulation scenarios reflecting frequently encountered real case studies inspired from the survival rates of 700 vertebrates’ species. We estimated the effects of mark loss and recycled individuals on parameter estimates using a multi-state Cormack-Jolly-Seber (MSCJS) framework. We explored parameter bias through simulations of a metapopulation system with different capture and survival rates. We also illustrated how mark loss can be easily estimated and accounted for using an empirical long-term (10 years) CMR dataset of bats, individually identified using both PIT tag technology as marks that can be lost, and multi-locus genotypes as ‘permanent marks’.

3. The results from our simulated scenarios demonstrated that the occurrence of bias and the parameters concerned were highly dependent on the study system, and no general rules on parameter behaviour can be established a priori. The model structure and the interdependency among parameters make it challenging to predict how bias could affect estimates.

4. Our results highlight the need to assess the effect of mark loss when using MSCJS models. Ignoring such violations of model assumptions can have important implications for ecological inferences and conservation policies. In general, the use of permanent marks (e.g. genotype), should always be preferred when modelling of population dynamics and if not possible, combining types of temporary marks (e.g. PIT tags, bands) should be considered.
KEYWORDS

1 Introduction

Capture-mark-recapture (CMR) methods have become a standard approach to estimate demographic rates of wild species thanks to the development of a range of different models. Accurately quantifying population dynamic parameters is critical to assess the state of populations, understand their dynamics, and ultimately make efficient management and conservation decisions. However, all CMR models make a number of assumptions of homogeneity (Johnson et al., 1986; Williams et al., 2002) that are prone to generate biases if violated. A widespread violation of CMR model assumptions, originally identified four decades ago (Nelson et al., 1980) is the loss of marks (see Supported Information 1, Table S2). Local survival is the product of true survival and mark retention rate for individual staying in the study area. If the retention rate starts to drop below one without being considered in the model, while true survival remains constant, then local survival is underestimated and becomes confused with the probability of presence of the mark, even more if true survival is high. Mark loss has two consequences: (1) when mark shedding occurs it induces non-identifiability of individuals, and individuals that have lost their mark may be considered either dead or out of the study area, despite being alive and present; (2) if such individuals are captured again, they will not be recognized and will be remarked as newly recruited individuals, which is called “recycling”.

Consider “1011”, a simple four occasions encounter history, where “1” implies the individual was caught and “0” not. If we note $\phi_t$ the survival rate between occasion $t$ and $t+1$ and $p_t$ the capture probability at occasion $t$ (with $q_t = 1 - p_t$), this encounter history occurs with probability $\phi_1 q_2 \phi_2 p_3 \phi_3 p_4$. Now consider the individual can make a transition between 2 states, and its history
becomes “1022”, with individual in state “1” at $t_1$ and in state “2” at $t_3$ and $t_4$. If we define $\psi^{i,j}$ the
transition probability from state $i$ to state $j$ (i and j in \{1,2\}), conditional on survival, the new
encounter history occurs with probability $\phi^{(1)}_1 \psi^{(1,1)} q^{(1)}_2 \phi^{(1,2)} p^{(2)}_3 \phi^{(2,2)} p^{(2)}_4 +$
$\phi^{(1)}_1 \psi^{(1,2)} q^{(2)}_2 \phi^{(2,2)} p^{(2)}_3 \phi^{(2,2)} p^{(2)}_4$, with indices for state-specific parameters noted in parentheses.

If this individual loses its mark after its first capture and is remarked when recaptured at $t_3$, its
encounter history becomes ‘two’ different histories from ‘two’ different individuals: one becoming
“1000”, the second “0022”. In this case, survival and mark loss patterns are different, not only is
survival underestimated (at least for the “first” history) but so are the transition probabilities,
because there is no longer a change of state (the second history starts directly at state 2).

Many CMR studies, despite using different mark types, are affected by mark loss, the rate of which
varies according to a myriad of factors, such as species (see Supported Information 1, Table S2),
mark type (Smout et al., 2011a), sex (Conn et al., 2004), mass (Schwarz et al., 2012), size (Acolas
et al., 2007), mark location (Kaemingk et al., 2011) or physiological stage (Besnard et al., 2007).

Mark loss has previously been shown to produce negative bias in survival estimates and detection
(Nichols et al., 1992; Nichols & Hines, 1993). Many statistical tools have been developed to
remove the confounding effect of mark loss in single state models, (Robson & Regier, 1966; G. A.
F. Seber & Felton, 1981). However, the effect of these violations has not been explored in the multi-
state modelling framework.

Multi-state models were developed to deal with situations where the “state” (e.g. location,
behaviour, physiology, reproductive or social status) of an individual may affect its survival or
detection probability, and where the individual can change “state” during life (reviewed in Lebreton
et al., 2009). These models became popular and widely used because of their flexibility to address a
large range of study systems and biological questions, and because they can be used in a relatively
straightforward manner by biologists given the development of user-friendly software (e.g. Mark,
E-surge, Bugs). They are used to address many ecological and evolutionary hypotheses based on
variations in life history traits (state transitions) throughout an individual’s life (Nichols & Kendall (1995) see also Cam (2009) for an extensive discussion on the subject), or density dependence effect (Schofield & Barker, 2008), co-evolution (Benkman et al., 2005), dispersal probability among subpopulations or living areas (Hestbeck et al., 1991; Spendelow et al., 1995), disease prevalence in wild populations (Jennelle et al., 2007). However, the literature dealing with the effect of mark loss on multi-state model behaviour is scarce and much remains to be explored.

To address this gap in knowledge, here we used simulation-based multi-state Cormack-Jolly-Seber (MSCJS) approaches to investigate the impact of mark loss on estimates of model parameters within a Bayesian framework. Given the increasing use of MSCJS models, our goal is to assess the potential bias in the marginal posterior distributions of demographic parameter estimates using a metapopulation context, based on biologically realistic scenarios and if possible, provide comprehensive guidelines for both, fieldwork and data analysis. The MSCJS model share assumptions with the CJS model (Cormack, 1964; Jolly, 1965; G. Seber, 1965), particularly in relation to mark loss. In addition, states are assumed to be recorded without error and survival, transition and observation probability must be the same for all individuals at a given occasion and state.

To illustrate our approach with an empirical example, we also used our decade-long mark recapture data set of PIT-tagged and genotyped Greater mouse-eared bat (Myotis myotis), a taxonomic group particularly susceptible to PIT-tags loss (Freeland & Fry, 1995). We used ‘genotype’ as an individual permanent mark to estimate bias between models accounting/not accounting for mark loss and recycling, and suggested recommendations for future studies. Similar to single state CJS models, we predicted that the MSCJS model would potentially be subject to underestimation of survival and overestimation of transitions in case of tag loss and recycling (Nichols & Hines, 1993). As transition probability is conditional on survival and detection on state in our MSCJS model, we expect errors in estimates to propagate to model parameters in different ways according to state.
transition rate. We then hypothesised that posterior distributions of parameters are likely to be biased even in case of low survival and recapture rate.

2 Material and method

2.1 Data simulation

To quantify the potential bias induced by mark loss on parameter estimates in the MSCJS framework, we defined several scenarios corresponding to realistic situations inspired from the literature (Supporting Information 1). We considered a study period of 10 years, during which an individual can change state among 4 states (A, B, C, D), state 'D' being an absorbing state (without possibility to change when it is reached, e.g. permanent emigration), see Fig. 1.b. We considered two age classes: (1) individuals less than 1-year-old (hereafter ‘juveniles’); (2) individuals older than 1 year (hereafter ‘adults’). Sampling took place every year and state assignment can only be made one year after birth at the earliest. Consequently, all individuals were adults when detected the second time. When individuals are in state D they cannot be captured (nor marked), only resighted. We allowed stochastic annual variations in survival rate for each age class.

To set the most common values for survival parameters, we surveyed 700 species in six vertebrate classes- Actinopterygii, Chondrichthyes, Amphibia, Aves, Reptilia, and Mammalia. We used the existing database for tetrapods (Conde et al., 2019) and also compiled data from the literature for fish and bat species (Supporting Information 1, Table S1). The distribution of survival rates encompasses a large range of values (Supporting Information 1, Fig. S1). We observed a positive correlation between adult and juvenile survival for a subset of 143 species in which both rates were available (Supporting Information 1, Fig. S2). Based on these surveys, we defined 4 scenarios of study: (1) long-lived species with high detection; (2) long-lived species with low detection; (3) short-lived species with low detection; (4) short-lived species with high detection. Values of parameter simulated are reported in Table 1. and Fig 1.b and the complete simulation steps of the
MSCJS model between two-time intervals are displayed in Fig. 1.a. We considered two common ways of re-encountering individuals after marking: (1) recapture when individuals are physically caught again; (2) resighting, which is a passive way to detect individuals without handling them (e.g. using colour or alphanumeric rings or PIT-tags). This approach is motivated by the fact that in most studies, the probability of resighting is typically higher than the probability of recapture, therefore they can be differently affected by estimation biases. To investigate the effect of mark loss on the estimates of state transition probabilities, we simulated data using different transition probabilities between states (Fig. 1.b). Irrespective of their age, females were allowed to transition at each occasion between state A, B and C, at a permanent rate depending on the state they were before transition, but not in state D. Only juvenile males were able to transition to the D state, in different proportions depending on their initial state, while adult male never changed their state (Fig. 1.b). This sexual dissymmetry in transition is common in mammals (and other taxa) and allows us for example to distinguish permanent emigration (absorbing state) from other movements.

At the first occasion, individuals in state A, B and C were composed of 40 juveniles (sex ratio 1:1) and 60 adults (80% females, 20% males); there was no individual in state D. On each subsequent occasion, 40 juveniles and 5 adults were marked in each state (A, B,C) except in state D, where it is not possible to recapture individuals but only to observe them.

As mark shedding is often dependant on time since marking and occurs most frequently shortly after marking in many species (in our case study; Fabrizio et al., 1999; Fokidis et al., 2006), for each scenario, we investigated a range of mark loss rates commonly reported in the literature (Supported Information 1, Table S2). We simulated three mark loss probabilities (0.05, 0.25, 0.4) during the first year after marking and a constant 0.05 rate thereafter, which generated a diversity of cases of mark loss and recycled individuals (Supported Information 2, Fig. S3). We simulated fifty datasets for each of the 12 combination of parameters, and ran two MSCJS models: one with recycled individuals, which mimicked the situation where it was not possible to identify individuals.
which have lost their mark; the second, accounting for mark loss, which mimicked situations where
a permanent mark was used to identify individuals (i.e. no recycling possible). In this last model we
only added to the previous model retention of mark as a Bernoulli process:

\[
R_{i,t} \sim \begin{cases} 
  \text{Bernoulli}(p_{r1}), & \text{if } i \text{ was juvenile and marked at } t - 1, \\
  \text{Bernoulli}(p_{r2}), & \text{if } i \text{ was adult and marked at } t - 1, \\
  \text{Bernoulli}(p_{r3}), & \text{if } i \text{ was adult and the mark was already present at } t - 1, \\
  0, & \text{if } i \text{ had no mark at } t - 1 \text{ and was not marked at } t - 1,
\end{cases}
\] (eqn 1)

with \(R_{i,t}\) the retention rate of individual \(i\) at time \(t\) and \(p_{rj}\) the probability of retention \(j\) (\(j\) in \{1,2,3\})
that depend both of the age of the individual \(i\) and the status of the mark at time \(t-1\). In total, 1,200
simulations were performed (Supported Information 2, Fig. S1). The computational codes for a
fully reproducible example dataset are provided in the Supporting Information 2.

2.2 Data analysis

In the 600 simulated dataset, taking into account mark loss (without recycling), we identified
individuals who lost their mark prior to running the model, and fitted a Bayesian MSCJS model,
with priors and codes detailed in Supporting Information 2 (ModelA.jags). In the 600 simulated
dataset, not accounting for tag loss, as individuals who lost their mark cannot be identified (i.e.
recycled), we simply cut life histories each time mark loss occurred and added a new line into the
dataset as a recycled individual. We then fitted a similar Bayesian MSCJS model, but removed the
part concerning mark loss estimation (Supporting Information 2, ModelW.jags). The model codes
and the initial values used for parameter estimation are provided in Supporting Information 2. All
prior distributions used for the model without recycling were the same as those used for the model
with recycled individuals.

2.3 Real data analyses

To illustrate the simulations, we analysed an empirical dataset from a 10-year study of CMR data on
the Greater-mouse eared bat (Myotis myotis) in Brittany (France; 2010-2019). A total of 2,561
Individuals were marked in 5 roosts: La Roche Bernard (47°31’N, 2°18’W), Férel (47°28’N, 2°20’W), Noyal-Muzillac (47°35’N, 2°27’W), Béganne (47°35’N, 2°14’W) and Limerzel (47°38’N, 2°21’W). The bats were individually tagged using PIT tags, ID-100C (Troven®), with a unique 10-digit code. These small passive tags (2.12x11mm, 0.1gr) allowed identification from passive readers. All individuals caught in roosts without PIT-tags were systematically tagged, which included both, individuals who lost their tag and those which were never tagged before, and genotyped as a second and permanent marking method. Genotypes were established from DNA extracted from wing biopsies from all individuals tagged and from all males untagged caught during swarming surveys (n=4,258 samples; details in Supporting Information 3, Fig. S1), as male capture probability at roosts after birth was extremely low. A total of 16 microsatellite markers optimised for *Myotis myotis* were used to establish individual genotypes (Foley et al., 2020). To minimize genotyping errors, we analysed, when available, 2 different samples per individual and all samples were genotyped and scored twice by two different people. We also performed genetic profile comparisons to detect errors (Frantz et al., 2003; Puechmaille & Petit, 2007). We hypothesised that the error rate of genotypes was low enough to be negligible and did not include this source of uncertainty in the models. We also checked each winter for the presence of lost tags on the floor of the roosts, which allowed us to identify, at least in part (66% compared to the genotype), the individuals that lost their tag, with the rest of losses having occurred outside of the roosts. Most of these tags were lost during the first year (Supporting Information 3, Fig. S2), which is confirmed by the absence of their records from passive reading detectors. In total, 252 individuals out of 2,561 (~10%) were identified as having lost their tag at least once. From those individuals, 94 individuals were recaptured and retagged a second time and three retagged a third time. As retagging occurred during the last capture occasion for 13 individuals, ignoring tag loss led to the recycling of 81 individuals in total among the 94. To analyse these data, we fitted a multisite model (a special case of multi-state models when states are sites and transition are movement between sites.), similar to...
the MSCJS model used for simulated data (Supporting Information 3). Emigration from the five studied subpopulations was assessed using capture and resighting data obtained between capture occasions in swarming and wintering sites. We accounted for individual heterogeneity, including a random effect in the estimation of survival probability, a widespread phenomenon in free populations and a known source of bias in CMR models. (Gimenez et al., 2018).

We defined eight possible detection states which allowed us to estimate separately, capture and resighting probabilities (see Supporting Information 3, Table S1). As empirical data suggested the possible movement of individuals between all subpopulations and outside, we did not constrain transition between subpopulations except for movement of juveniles from D toward A, B, C which was not possible and therefore set to 0. In this study, we also tested the effect of surgical adhesive (Vetbond®) after PIT-tag injection, in order to assess tag-loss reduction in comparison with self-healing (Lebl & Ruf, 2010; van Harten et al., 2020). In this model, tag retention probabilities were modelled like in the simulation study (eqn. 1), by accounting for time since marking (two discrete classes: first year or subsequent years), individual age class (juvenile or adult), and the use of surgical adhesive (yes or no) As per the simulated datasets, two models were run, a first one taking into consideration tag loss and recycling, and a second model ignoring this information.

### 2.4 Computation details

We used JAGS (Plummer, 2003) with the jagsUI package (Kellner, 2016) from R 3.6.0 (R Core Team, 2019) for analyses of simulated and empirical datasets. We used four MCMC chains with 150,000 iterations each and drew samples from posterior distributions after discarding the first 50,000 iterations (burn-in). We retained every twentieth iteration generating 20,000 samples from the posterior distribution for each parameter. Chain convergence was assessed with the Gelman-Rubin statistic (Brooks & Gelman, 1998). Among 1,200 simulations, some showed R-hat values > 1.1, indicating a convergence failure. Converge problems concerned few parameters and the corresponding R-hat values were still low. To avoid prohibitively long computing time, we did not
attempt to increase the number of iterations to achieve the complete convergence of MCMC chains for these parameters in the simulations concerned. Our results are based on 50 simulated datasets per scenario, and it is expected that lack of convergence for these few parameters has no substantial influence on our results.

For assessing bias in parameter estimates when mark loss or recycling is not accounted for, we computed the Earth Mover Distance (EDM), using the EMD-\(L_1\) algorithm (Ling & Okada, 2007), a distance function that quantifies the difference between two distributions. This metric measures the minimum cost of turning, point by point, a distribution into another. We also estimated a ROPE (Region Of Practical Equivalence, Kruschke, 2018) for each scenario to assess the degree of difference between distribution represented by the EDM metric. To define this ROPE for each scenario, we randomly built 1,000 pairs of models from the 50 simulations and calculated the associated 1,000 EDMs from the posterior distributions of the estimated parameters (Supporting Information 2, Fig. S43). The obtained distributions of EDM represented variations expected for inferences obtained from simulations initiated with the same parameter values. The ROPE was then defined between 0 and the upper value of the 80% highest posterior density interval (hdi) from the distribution of these EDM. Finally, the proportion of EDM for each simulated case outside the ROPE was computed, giving a direct indication of bias, the higher this proportion, the higher the bias. Comparisons of EDM between the models that did and did not account for tag loss and recycling to their respective ROPE is a way to illustrate cases in which not accounting for tag loss leads to estimates that substantially differ from estimates obtained when accounting for tag loss. However, it is not a way to estimate absolute bias. This is not a problem when the model that accounts for tag loss correctly recovers simulated parameter values, which was the case for most parameters in most situations.
3 Results

3.1 Simulation results

Mark loss and recycling increase in scenarios with higher survival, recapture and mark loss rate (Supported Information 2, Fig. S3). However, the estimates from the model accounting for mark loss and recycling did not show major departures from the simulated values in any scenario (Supporting Information 2, Fig. S3-S42). Irrespective of scenarios, little to no estimation bias appeared when tag loss was set to 0.05, only resighting probability showed some underestimations.

The number of parameters whose estimates were biased increased with increasing mark loss rate, but the extent was scenario-dependent (Fig. 2 & 3). In particular, Adult survival showed substantial underestimation (Fig. 2.b) in scenario 1 for all state but “D”, but bias decreased with time (Supporting Information 2, Fig. S7-S14). For scenario 4, moderate underestimates in adult survival were also observed during the first study year. Juvenile survival and capture probabilities were less biased, nevertheless moderate underestimations occurred for high mark loss rate (Supporting Information 2, Fig. S11-S14 and S23-S26).

Resighting probability showed substantial bias, with underestimates mainly in state A and C, for all scenarios and mark loss rates (Fig. 3.a). A large percentage of the transition probabilities estimates were also biased when mark loss and recycling were ignored (Fig. 3.b-e), with in general an underestimates of the probability to remain in the same state and, as a corollary, an overestimates of the probability of changing of state (Supporting Information 2, Fig. S27-S37). Overestimations occurred in juvenile males except from state C, where transition rate was the highest (0.9). For females, the same pattern was observed in state A and B for which transition rates were low. On the contrary, for state transition from state C to B (set high = 0.4), we observed an underestimation of the transition probability and an overestimation of the probability to remain in state C with scenarios set with low detection rates (scenario 2 and 3, Supporting Information 2, Fig. S36-S37).
3.2 Bat metapopulation

Most of the estimated parameters (survival, capture, resighting and state transition probabilities) show both negative and positive biases, with no clear pattern, but no substantial bias appeared in the model not accounting for tag loss and recycling compared to the model taking them into account (Supporting Information 3, Fig. S4-S11). Larger biases appeared in survival estimates, with the estimated median survival difference between the two models reaching values greater than 0.2 (Supplementary Information 3, Fig. S4-S5), which could profoundly affect the results of studies interested in evaluating the effect of annual covariates (e.g., climate variables) on survival probability. The estimated tag loss probability was high for juveniles but substantially reduced by the use of surgical glue (Fig. 4), decreasing by one third from 0.28 (90%hdi [0.23,0.33]) to 0.19 (90%hdi [0.16,0.22]). However, in adults the use of surgical glue did not affect tag loss rate, with an overlap of 69% of the probability distributions. The adult tag loss rate was around 0.1 which is two times less than in juveniles when surgical glue was used. Considering the period following one-year post-tagging, the probability of tag loss when surgical glue was used is higher (median 0.03, 90% hdi [0.02, 0.04]) versus not used (median 0.02, 90%hdi [0.01, 0.02]). This difference is in fact an artefact due to a lack of search for lost tags on the ground of the colonies in the first year of the study (Supplementary Information 3, part 3 and Fig. S12). Other parameter estimates can be found in Supporting Information 3, part 2.6.

4 Discussion

We used simulations to assess the effect of mark loss and recycled individuals on parameter estimates in the MSCJS model, where capture and survival probabilities vary over time, survival also varies among age classes and mark loss processes depend on time since marking. We showed that not only survival, but capture, resighting and state transition probabilities estimates can be substantially affected by this violation of CMR model assumptions. Survival is mainly
underestimated in cases where capture and detection is high, this trend is accentuated when survival is also high, which moderates previous studies suggesting that biases mainly occur in species with both high survival rates, catchability and mark loss (Diefenbach & Alt, 1998). Our simulation results confirmed that inaccuracy of model estimates is positively linked to the rate of mark loss, but can also occur when tag loss rate is low (5%) and can be independent of survival and capture rates. Indeed, in datasets with few recycled individuals, i.e. with low survival and capture rates, transition and resighting probabilities can be severely biased if mark loss is high. This suggests that results from studies where survival and capture are low should also be interpreted with caution if mark loss is suspected but not taken into consideration. Severity of bias can also vary over time, with bias in survival decreasing with time as in our simulated datasets. This is partly due to the mark loss pattern we chose and stresses the fact that, even in studies conducted over short periods, parameters can be substantially biased. In cases where mark loss increases with time since marking, we would expect an increase of bias with time.

The simulation and the empirical study results together highlight how the complexity and interdependence of parameters can compound or counteract estimation biases in the absence of mark loss modelling. While the simulations showed some global rules on the direction of the biases, the real example showed instead how unexpected the bias pattern can be. Although we tried to cover in our simulation demographic variations classically observed across vertebrates, they have not been carried out under the immense range of parameters combinations that can be encountered. Most study systems and monitoring methods have their own specificities and a priori prediction of bias without simulating them can be misleading. The propagation of uncertainty in parameter estimates due to mark loss is still challenging to predict and increases with system complexity. Therefore, prior to planning a CMR study, we advocate that researchers/managers run simulations to evaluate in which conditions (i.e. under which parameters combination) their study would provide reliable estimates of the parameters of interest (e.g. demographic, state transition).
Preliminary studies with multiple marks could also be considered when possible (Smout et al., 2011b). This would allow CMR study design to be optimised prior to the study being conducted and hence limit biases from the onset.

MSCJS models have been partly developed to estimate movement between sites, recruitment, dispersion, temporary or permanent emigration (Lebreton et al., 2003, 2009; Schaub et al., 2004). Our simulation results suggest that state transition probabilities are sensitive to mark loss, even at low rates, e.g. the probability to stay in the same state (philopatry) or to change state showed both under and overestimations. These parameters are often of central interest in many studies to answer ecological and demographic questions and are used for management and conservation purposes (Cam et al., 2004; Horton et al., 2011). Although the loss of marks is regularly reported for a wide variety of tags and taxa, it is only marginally considered in studies interested in estimates of population dynamics parameters (Nelson et al., 1980; Ostrand et al., 2012; Smout et al., 2011b; Tavecchia et al., 2012). Most model developments to account for mark loss have focussed on the Jolly-Seber model (Jolly, 1965; G. Seber, 1965) for abundance estimates, where mark loss and recycling are prone to generate serious biases (Malcolm-White et al., 2020). The loss of the mark remains largely unconsidered despite an increasing use of MSCJS models in ecology, demography, management and conservation (Huntsman et al., 2020; Melnychuk et al., 2017). In light of our study, we recommend the use of permanent or double temporary marks, where dependence in loss is taken into account (Laake et al., 2014; McMahon & White, 2009), as any analysis of CMR data is potentially affected by this violation of model assumption (Riecke et al., 2019).

Despite the fact that PIT tags are suitable in an increasing number of studies and allow the collection of data without physically re-capturing individuals, our case study highlights the importance of a second marking method to estimate potential bias in demographic rates. Tag loss has long been known in small mammal species, in particular those that fly or glide (Freeland & Fry,
1995). Here we confirmed that PIT-tag shedding can be reduced in the short term by the use of surgical adhesive (Lebl & Ruf, 2010; van Harten et al., 2020).

As illustrated, surgical adhesive is not sufficient to reduce tag loss to zero, and the use of a permanent mark (e.g. genotype) is required, for all or part of the population studied (Laake et al., 2014). Similar situations, where permanent marks should be considered, arise when marks deteriorate and become unreadable, equivalent to an increase of mark loss with time, like neck collars or ear tags (Conn et al., 2004; Diefenbach & Alt, 1998). In this case, we expect a decrease in accuracy of model parameter estimates for the duration of the study, and further supports the use of permanent marks for CMR studies.

Mark loss is typically not considered from ecological and management perspectives except when researchers are interested in understanding factors influencing mark failures or in improving their marking methods. Our results highlight the need to assess the effect of mark loss each time mark failure is suspected, to avoid spurious conclusions about the dynamics of their studied species. In our experience and according to the literature, PIT-tags are prone to being shed regardless of the studied taxa, often in the short but sometimes in the long term. Mark loss should therefore be carefully considered in all CMR analyses and possibly also in other studies using similar datasets, and should be explicitly modelled when necessary for more accurate estimations of population dynamics.

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Author contribution

F.T. designed the project with the other co-authors. E.C.T., F.T. and BV acquired the funding. E.C.T., E.J.P., S.J.P and E.C. supervised the project. E.C.T., F.T., E.J.P., S.J.P and BV supervised the fieldwork. E.C.T., E.J.P., F.T. and S.J.P. collected samples. S.J.P. & F.T. supervised the microsatellite genotyping. C.S. generated the genotypes. F.T. and E.C. developed the R scripts for simulating and analysing data with help of E.J.P. and S.J.P. F.T. led the writing of the manuscript and all authors contributed to manuscript revisions and gave final approval for publication.

Data accessibility

R scripts for simulating the data, and analysing the data with JAGS, will be archived at the Dryad Digital Repository.

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Table 1: Parameter values used to simulate the 4 scenarios. For random values generated, the corresponding distribution is indicated in brackets (N=Normal distribution; U=Univariate distribution). For transition values between states see Fig. 1.b.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
<th>Value</th>
<th></th>
<th>Value</th>
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</thead>
<tbody>
<tr>
<td>$\phi_{ad.}$</td>
<td>Adults survival in state A, B, C</td>
<td>$\sim$0.92 [logit(N(2.5,0.3))]</td>
<td>$\sim$0.62 [logit(N(2.5,0.3)-0.3))]</td>
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<td>$\phi_{ad.}$</td>
<td>Adults survival in state D</td>
<td>$\sim$0.81 [logit(N(1.5,0.3))]</td>
<td>$\sim$0.51 [logit(N(1.5,0.3)-0.3))]</td>
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<tr>
<td>$\phi_{JRV.}$</td>
<td>Juveniles survival in state A, B, C</td>
<td>$\sim$0.55 [logit(N(0.2,0.3))]</td>
<td>$\sim$0.35 [logit(N(0.2,0.3)-0.2))]</td>
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</tr>
<tr>
<td>$p_{CA}$</td>
<td>Capture probability in state A</td>
<td>$\sim$0.65 [U(0.6-0.7)]</td>
<td>$\sim$0.15 [U(0.6-0.7)-0.5]</td>
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<tr>
<td>$p_{CB}$</td>
<td>Capture probability in state B</td>
<td>$\sim$0.75 [U(0.7-0.8)]</td>
<td>$\sim$0.25 [U(0.7-0.8)-0.5]</td>
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<tr>
<td>$p_{CC}$</td>
<td>Capture probability in state C</td>
<td>$\sim$0.7 [U(0.65-0.75)]</td>
<td>$\sim$0.2 [U(0.65-0.75)-0.5]</td>
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<tr>
<td>$p_{CD}$</td>
<td>Capture probability in state D</td>
<td>0</td>
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<tr>
<td>$p_{A}$</td>
<td>Resighting probability in state A</td>
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<td>0.35</td>
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<tr>
<td>$p_{B}$</td>
<td>Resighting probability in state B</td>
<td>0.95</td>
<td>0.45</td>
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<tr>
<td>$p_{C}$</td>
<td>Resighting probability in state C</td>
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<td>0.4</td>
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<tr>
<td>$p_{D}$</td>
<td>Resighting probability in state D</td>
<td>0.7</td>
<td>0.2</td>
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</table>
Figure 1: Multistate capture-mark-recapture model. Panel (a) illustrate the fate of individuals captured and realised at t or dead at t. From t to t+1 dead individuals stay dead and cannot be detected (absorbing state), while released individuals can change state. Arrows indicate the possible paths, first in the state-transition, second in the mark retention process and third in the observation steps. Indices on arrows indicate corresponding probabilities. $\phi$ is the survival probability (time- and state-dependent), $\psi$ is the state transition indicated in panel (b) where plain arrows correspond to females’ possible transitions and dotted arrows to males. $ptl$ is the probability to lose the mark (dependent on age and time from marking). $p_c$ is the capture probability (time- and state-dependent), $p_r$ is the resighting probability (state-dependent), and the detection probability equal $(1-p_c)(1-p_r)$. Only juvenile males can reach state D and can be re-observed but not recaptured. On panel (b), simulated survival (Surv.) and detection (Det.) rates were displayed for states A, B and C according to scenario 2 (long-lived species with low detection). Blue lines correspond to adult survival, orange line to juvenile survival, grey line to resighting probability and green line to capture probability.
Figure 2: Tile-plots of the proportion of the distribution of the Earth Mover Distance (across 50 simulated datasets) out of the Region of Practical Equivalence (ROPE), between the model accounting for tag loss and recycling and the model ignoring them. Each tile represents annual (right axis) juvenile survival (a), adult survival (b) and capture probability (c) for each scenario and tag loss probabilities.
Figure 3: Tile-plots of the proportion of the distribution of the Earth Mover Distance (across 50 simulated datasets) out of the Region of Practical Equivalence (ROPE), between model accounting for tag loss and recycling and the model ignoring them for each simulated cases. Each tile represents resighting probability (a) and movement probabilities between subpopulations (direction, “from-to”, are indicated above each tile-plot) of juvenile female (b), adult female (c), juvenile male (d) and adult male (e) for each scenario and tag loss probabilities.
Figure 4: Posterior distribution of the tag loss probabilities according to age classes and time after marking in the *Myotis myotis* dataset. Left panel correspond to tag shedding rate during the year following the tag injection and the right panel for the following years (constant in time). In blue, distribution if surgical adhesive was used after tag injection and in red, without surgical adhesive.