STATE-SPACE MODELLING OF DATA ON MARKED 1 INDIVIDUALS 2 Olivier GIMENEZ^{1,2,6}, Vivien ROSSI^{3,4,5}, Rémi CHOQUET¹, Camille DEHAIS¹, 3 Blaise DORIS^{1,3}, Hubert VARELLA^{1,3}, Jean-Pierre VILA³ and Roger PRADEL¹ 4 ¹Centre d'Ecologie Fonctionnelle et Evolutive/CNRS - UMR 5175 5 1919 Route de Mende 6 34293 Montpellier - FRANCE ²Centre for Research into Ecological and Environmental Modelling 8 University of St Andrews, St Andrews The Observatory, Buchanan Gardens, KY16 9LZ - SCOTLAND 10 ³Laboratoire d'Analyse des Systèmes et Biométrie - UMR 729 11 INRA/ENSAM 12 2 Place Pierre Viala, 34060 Montpellier - FRANCE 13 ⁴Institut de Modélisation Mathématiques de Montpellier - UMR 5149 14 Universite Montpellier 2 15 CC051, Place Eugène Bataillon, 34095 Montpellier Cedex 5 - FRANCE 16 ⁵ CIRAD - Unité de dynamique des forêts naturelles 17 TA C-37 / D 18 Campus International de Baillarguet, 3434398 Montpellier Cedex 5 - FRANCE 19 ⁶E-mail: olivier.gimenez@cefe.cnrs.fr 20 21 Abstract State-space models have recently been proposed as a convenient and flexible 22 framework for specifying stochastic models for the dynamics of wild animal pop-23 ulations. Here we focus on the modelling of data on marked individuals which 24

is frequently used in order to estimate demographic parameters while accounting
for imperfect detectability. We show how usual models to deal with capturerecapture and ring-recovery data can be fruitfully written as state-space models.
An illustration is given using real data and a Bayesian approach using MCMC
methods is implemented to estimate the parameters. Eventually, we discuss future developments that may be facilitated by the SSM formulation.

$_{7}$ 1 Introduction

The estimation of animal survival is essential in population biology to investigate pop-8 ulation dynamics, with important applications in the understanding of ecological, evo-9 lutionary, conservation and management issues for wild populations (Pollock, 1991; 10 Williams et al., 2002). While the time to event is known in medical, social or engi-11 neering sciences (death, marriage and failure respectively), models for estimating wild 12 animal survival must incorporate nuisance parameters to account for incomplete de-13 tectability in monitoring individuals (Schwarz et Seber, 1999). Typically, individuals 14 are captured, marked and can be resigned or recaptured (encountered thereafter) to 15 construct encounter histories which consist of sequences of 1's and 0's according to 16 whether a detection occurs or not. The likelihood for such data arises from products 17 of multinomial distributions whose cell probabilities are complex functions of survival 18 probabilities - parameters of primary interest - and encounter probabilities - nuisance 19 parameters (Cormack, 1964; Jolly, 1965; Seber, 1965 - CJS thereafter). 20

In this note, we show how the population process can be fruitfully disentangled, by distinguishing the underlying demographic process, i.e. the survival (as well as transitions between sites/states if needed), from its observation, i.e. the detectability. This leads us to consider a natural formulation for capture-recapture models using statespace models (SSMs). Our contribution is in line with a recent paper by Buckland et
al. (2004) who have proposed to adopt SSMs as a convenient and flexible framework
for specifying stochastic models for the dynamics of wild animal populations.

Thus far, SSMs have been mainly used for time series of animal counts (de Valpine, 2004; Millar and Meyer, 2000) or animal locations (Anderson-Sprecher and Ledolter, 1991) to allow true but unobservable states (the population size or trajectory) to be r inferred from observed but noisy data (see Clark et al., 2005 and Wang, 2006 for reviews). The novelty of our approach lies in the use of SSMs to fit capture-recapture models to encounter histories.

In Section 2, we discuss how to express the CJS model under the form of a SSM. The 10 implementation details are provided, and real data are presented to compare parameter 11 estimates as obtained using the standard product-multinomial and the SSM approaches. 12 In Section 3, the flexibility of the state-space modeling approach is demonstrated by 13 considering two widely used alternative schemes for collecting data on marked animals. 14 Finally, Section 4 discusses important developments of capture-recapture models facil-15 itated by the SSM formulation. We emphasize that this general framework has a great 16 potential in population ecology modelling. 17

¹⁸ 2 State-space modelling of capture-recapture data

¹⁹ We focus here on the CJS model for estimating animal survival based on capture-²⁰ recapture data, as this model is widely used in the ecological and evolutionary litterature ²¹ (e.g. Lebreton et al., 1992).

¹ 2.1 Likelihood

We first define the observations and then the states of the system. We assume that 2 n individuals are involved in the study with T encounter occasions. Let $X_{i,t}$ be the 3 binary random variable taking values 1 if individual i is alive at time t and 0 if it is 4 dead at time t. Let $Y_{i,t}$ be the binary random variable taking values 1 if individual i is 5 encountered at time t and 0 otherwise. Note that we consider the encounter event as 6 being physically captured or barely observed. The parameters involved in the likelihood 7 are $\phi_{i,t}$, the probability that an animal *i* survives to time t + 1 given that it is alive 8 at time t (t = 1, ..., T - 1), and $p_{i,t}$ the probability of detecting individual i at time t9 $(t = 2, \ldots, T)$. Let finally e_i be the occasion where individual i is encountered for the 10 first time. A general state-space formulation of the CJS model is therefore given by: 11

12

$$Y_{i,t}|X_{i,t} \sim Bernoulli(X_{i,t}p_{i,t}), \tag{1}$$

$$X_{i,t+1}|X_{i,t} \sim Bernoulli(X_{i,t}\phi_{i,t}), \tag{2}$$

for $t \ge e_i$, with $p_{i,e_i} = 1$ and where Equation (1) and Equation (2) are the observation 14 and the state equations respectively. This formulation naturally separates the nuisance 15 parameters (the encounter probabilities) from the parameters of actual interest i.e. the 16 survival probabilities, the latter being involved exclusively in the state Equation (2). 17 Such a clear distinction between a demographic process and its observation makes the 18 description of a biological dynamic system much simpler and allows complex models 19 to be fitted (Pradel, 2005; Clark et al., 2005). We will refer to this formulation as 20 the individual state-space CJS model (individual SSM CJS hereafter). The rationale 21 behind the above formulation is as follows. We give the full details for the observation 22 Equation (1) only, as a similar reasoning easily leads to Equation (2). If individual i is 23

alive at time t, then it has probability $p_{i,t}$ of being encountered and probability $1 - p_{i,t}$ otherwise, which translates into $Y_{i,t}$ is distributed as $Bernoulli(p_{i,t})$ given $X_{i,t} = 1$. Now if individual i is dead at time t, then it cannot be encountered, which translates into $Y_{i,t}$ is distributed as Bernoulli(0) given $X_{i,t} = 0$. Putting together those two pieces of reasoning, the distribution of the observation $Y_{i,t}$ conditional on the state $X_{i,t}$ is given by Equation (1).

Statistical inference then requires the likelihood of the state-space model specified above. Assuming independence of individuals, the likelihood is given by the product of all individual likelihood components. The likelihood component for individual i is the probability of the vector of observations $\mathbf{Y}_i^T = (Y_{i,e_i}, \ldots, Y_{i,T})$ which gathers the information set up to time T for this particular individual. Conditional on the first detection, the likelihood component corresponding to individual i is therefore given by (e.g. Harvey, 1989)

$$\int_{X_{i,e_i}} \dots \int_{X_{i,T}} [X_{i,e_i}] \left\{ \prod_{t=e_i+1}^T [Y_{i,t}|X_{i,t}][X_{i,t}|X_{i,t-1}] \right\} dX_{i,e_i} \dots dX_{i,T}$$
(3)

14

where [X] denotes the distribution of X and X_{i,e_i} the initial state of individual *i* which is assumed to be alive. Because we deal with binary random vectors, we used the counting measure instead of the Lebesgue measure.

In its original formulation, the CJS makes important assumptions regarding individuals. First, all individuals share the same parameters, which means that the survival and detection probabilities depend on the time index only. In mathematical notation, we have $\phi_{i,t} = \phi_t$ and $p_{i,t} = p_t$ for all i = 1, ..., n, so Equation (1) and Equation (2) become $X_{i,t+1}|X_{i,t} \sim Bernoulli(X_{i,t}\phi_t)$ and $Y_{i,t}|X_{i,t} \sim Bernoulli(X_{i,t}p_t)$ respectively. Second, the CJS model also assumes independence between individuals. By using simple relationships between Bernoulli and Binomial distributions, one can finally fruitfully
 formulates the original CJS model under the following state-space model:

$$Y_t | X_t \sim Bin(X_t - u_t, p_t) \tag{4}$$

3

$$X_{t+1}|X_t \sim Bin(X_t, \phi_t) + u_{t+1} \tag{5}$$

where X_t is the number of survivors from time t plus the number of newly marked 5 individuals at time t, u_t , and Y_t is the total number of previously marked individuals 6 encountered at time t. We will refer to this formulation as the population state-space 7 CJS model (population SSM CJS hereafter). Interestingly, specifying the system un-8 der a state-space formulation now requires much less equations than the individual 9 SSM CJS model, which may avoid the computational burden. Nevertheless, while the 10 individual SSM CJS involves parameters for every single individual and sampling occa-11 sion, the population SSM CJS model makes the strong assumptions that all individuals 12 behave the same as well as independently, which may be of little relevance from the 13 biological point of view. To cope with this issue, in-between modelling can be achieved 14 by considering age effects or groups classes in the population SSM model (Lebreton et 15 al., 1992). Finally, covariates can be incorporated in order to assess the effect of envi-16 ronment such as climate change, most conveniently by writing the relationship between 17 the target probabilities and the predictors on the logit scale (Pollock, 2002). 18

¹⁹ 2.2 Implementation

Fitting SSMs is complicated due to the high-dimensional integral involved in the individual likelihood Equation (3). To circumvent this issue, several techniques have been proposed including Kalman filtering, Monte-Carlo particle filtering (such as sequential importance sampling) and MCMC (see Clark et al., 2005 and Wang, 2006 for reviews).
Our objective here is not to discuss these different methods. For our purpose, we adopt
the MCMC technique which is now widely used in biology (Ellison, 2004; Clark, 2005),
in particular for estimating animal survival (Seber et Schwarz, 1999; Brooks et al.,
2000). Besides, this is to our knowledge the only methodology which comes with an
efficient and flexible program to implement it, which, in our case, will allow biologists
to easily and rapidly adopt our approach.

In addition to the difficulty of estimating model parameters, the use of SSMs raises
several important issues regarding identifiability, model selection and goodness-of-fit
(Buckland et al. 2004) that were not discussed here. Noteworthy, Bayesian modelling
using MCMC methods offer possible solutions reviewed in Gimenez et al. (submitted).

12 2.3 Illustration

We consider capture-recapture data on the European dipper (*Cinclus cinclus*) that were 13 collected between 1981 and 1987 (Lebreton et al., 1992). The data consists of marking 14 and recaptures of 294 birds ringed as adults in eastern France. We applied standard 15 maximum-likelihood estimation (Lebreton et al. 1992) and MCMC techniques (Brooks 16 et al. 2000) using the product-multinomial likelihood and the state-space likelihood of 17 Equation (3) in combination with Equation (1) and Equation (2). We ran two overdis-18 persed parallel MCMC chains to check whether convergence was reached (Gelman, 19 1996). We used 10,000 iterations with 5,000 burned iterations for posterior summariza-20 tion. We used uniform flat priors for both survival and detection probabilities. The 21 simulations were performed using WinBUGS (Spiegelhalter et al., 2003). The R (Ihaka 22 and Gentleman, 1996) package R2WinBUGS (Sturtz et al., 2005) was used to call Win-23 BUGS and export results in R. This was especially helpful when converting the raw 24

encounter histories into the appropriate format, generating initial values and calculate
posterior modes. The programs are available in Appendix A. Posterior summaries for
encounter and survival probabilities are given in Table 1, along with their posterior
probability distributions that are displayed in Figure 1.

5

[Table 1 about here.]

6

[Figure 1 about here.]

Survival estimates were uniformally similar whatever the method used (Table 1). In
particular, there is a clear decrease in survival 1982-1983 and 1983-1984, corresponding
to a major flood during the breeding season in 1983 (Lebreton et al., 1992).

In contrast, posterior medians of detection probabilities using the CJS SSM approach are quite different from the classical maximum likelihood estimates, but more similar to the posterior medians obtained with the product-multinomial likelihood approach (Table 1). These discrepancies are no longer present when posterior modes are examined, as expected (recall that we use non-informative uniform distributions as priors for all parameters).

The last survival probability as well as the last detection probability are estimated with high variability (Table 1 and Figure 1). The fact that these two parameters cannot be separately estimated is not surprising since the CJS model is known to be parameterredundant (Catchpole and Morgan, 1997). Also, the first survival probability and the first detection probability are poorly estimated, due to the fact that very few individuals were marked at the first sampling occasion (approximately 7% of the full data set).

In terms of time computation, the MCMC approach using a product-multinomial likelihood took 30 seconds to run and a few second for the classical approach, while the MCMC approach using the SSM likelihood took 4 minutes (512Mo RAM, 2.6GHz
2 CPU).

³ 3 Further state-space modelling

4 3.1 Multistate capture-recapture models

Multistate capture-recapture models (Arnason, 1973; Schwarz et al., 1993; AS hereafter) 5 are a natural generalization of the CJS model in that individuals can move between 6 states, according to probabilities of transition between those states. States can be either 7 geographical sites or states of categorical variables like reproductive status or size class 8 (Lebreton and Pradel, 2002). We provide here a state-space modelling formulation 9 of the AS model (Dupuis, 1995; Newman, 1998; Clark et al., 2005). Without loss of 10 generality, we consider 2 states. Let $X_{i,t}$ be the random state vector taking values 11 (1,0,0), (0,1,0) and (0,0,1) if, at time t, individual i is alive in state 1, 2 or dead 12 respectively. Let $Y_{i,t}$ be the random observation vector taking values (1,0,0), (0,1,0)13 and (0,0,1) if, at time t, individual i is encountered in state 1, 2 or not encountered. 14 Parameters involved in the modelling include $\phi_{i,t}^{rs}$, the probability that an animal i 15 survives to time t + 1 given that it is alive at time t (t = 1, ..., T - 1) and makes the 16 transition between state r and state s over the same interval (r, s = 1, 2), as well as $p_{i,t}^r$ 17 the probability of detecting individual i at time t in state r (t = 2, ..., T, r = 1, 2). A 18 state-space formulation for the multistate AS model is given by: 19

$$Y_{i,t}|X_{i,t} \sim Multinomial \left(1, X_{i,t} \begin{bmatrix} p_{i,t}^1 & 0 & 1 - p_{i,t}^1 \\ 0 & p_{i,t}^2 & 1 - p_{i,t}^2 \\ 0 & 0 & 1 \end{bmatrix} \right)$$
(6)

20

$$X_{i,t+1}|X_{i,t} \sim Multinomial \left(1, X_{i,t} \begin{bmatrix} \phi_{i,t}^{11} & \phi_{i,t}^{12} & 1 - \phi_{i,t}^{11} - \phi_{i,t}^{12} \\ \phi_{i,t}^{21} & \phi_{i,t}^{22} & 1 - \phi_{i,t}^{21} - \phi_{i,t}^{22} \\ 0 & 0 & 1 \end{bmatrix} \right)$$
(7)

where Equation (6) and Equation (7) are the observation and the state equations 2 respectively. This formulation has similarities with that of Pradel (2005) who used 3 hidden-Markov models to extend multistate models to cope with uncertainty in state 4 assignment. Again, it should be noted that the state-space formulation allows the de-5 mographic parameters to be distinguished from the nuisance parameters. A similar 6 reasoning to that adopted for the CJS model leads to Equations (6) and (7). As ex-7 pected, Equation (6) and Equation (7) reduce to Equation (1) and Equation (2) if one 8 considers a single state. Making similar assumptions as in the CJS model leads to the 9 population AS SSM. 10

¹¹ 3.2 Ring-recovery models

1

The capture-recapture models presented above deals with apparent survival, which 12 turns out to be true survival if emigration is negligeable. When marks of individuals 13 (or individuals themselves) are actually recovered, true survival probabilities can be 14 estimated using ring-recovery models (Brownie et al., 1985; RR models hereafter). Let 15 $X_{i,t}$ be the binary random variable taking values 1 if individual i is alive at time t and 16 0 if it is dead at time t. Let $Y_{i,t}$ be the binary random variable taking values 1 if mark 17 of individual i is recovered at time t and 0 otherwise. The parameters involved in the 18 likelihood are $\phi_{i,t}$, the probability that an animal *i* survives to time t + 1 given that 19 it is alive at time t (t = 1, ..., T - 1), and $\lambda_{i,t}$ the probability of recovering the mark 20 of individual i at time t (t = 2, ..., T). A general state-space formulation of the RR 21

¹ model is therefore given by:

2

3

$$Y_{i,t}|X_{i,t}, X_{i,t-1} \sim Bernoulli\left((X_{i,t-1} - X_{i,t})\lambda_{i,t}\right)$$
(8)

$$X_{i,t+1}|X_{i,t} \sim Bernoulli(X_{i,t}\phi_{i,t}) \tag{9}$$

where Equation (8) and Equation (9) are the observation and the state equations re-4 spectively. While the state Equation (9) is the same as that in the individual SSM 5 CJS, the observation Equation (8) deserves further explanation. If individual *i*, alive 6 at time t, does not survive to time t + 1, then its mark has probability $\lambda_{i,t}$ of being 7 recovered and probability $1 - \lambda_{i,t}$ otherwise, which translates into $Y_{i,t}$ is distributed as 8 $Bernoulli(\lambda_{i,t})$ given $X_{i,t-1} = 1$ and $X_{i,t} = 0$ i.e. $X_{i,t-1} - X_{i,t} = 1$. Now if individual i 9 is in a given state (dead or alive) at time t and remains in this state till time t+1, then 10 its mark cannot be recovered, which translates into $Y_{i,t}$ is distributed as Bernoulli(0)11 given $X_{i,t-1} = 0$ and $X_{i,t} = 0$ or $X_{i,t-1} = 1$ and $X_{i,t} = 1$ i.e. $X_{i,t-1} - X_{i,t} = 0$. The dis-12 tribution of the observation $Y_{i,t}$ conditional on the combination of states $X_{i,t-1} - X_{i,t}$ is 13 thus given by Equation (8). Similar comments to that of previous sections can be made 14 here as well. Finally, we note that because the probability distribution of the current 15 observation does not only depend on the current state variable, the model defined by 16 Equation (8) and Equation (9) does not exactly matches the definition of a state-space 17 model but can be rewritten as such (see Appendix B). 18

¹⁹ 4 Discussion

We have shown that, by separating the demographic process from its observation, CR models for estimating survival can be expressed as SSMs. In particular, the SSM formulation of the CJS model competes well with the standard method when applied to a real data set. Bearing this in mind, we see at least two further promising developments
to our approach.

First, by scaling down from the population to the individual level while modelling 3 the survival probabilities, individual random effects can readily be incorporated to cope 4 with heterogeneity in the detection probability (Huggins, 2001) and deal with a frailty in 5 the survival probability (Vaupel and Yashin, 1985). Second, the combination of various 6 sources of information which has recently received a growing interest, (e.g. recovery 7 and recapture data, Catchpole et al., 1998; recovery and census data, Besbeas et al., 8 2002; Besbeas et al., 2003) can now be operated/conducted in a unique SSM framework 9 and hence benefits from the corpus of related methods. Of particular importance, we 10 are currently investigating the robust detection of density-dependence by accounting 11 for error in the measurement of population size using the combination of census data 12 and data on marked individuals. 13

Because most often, data collected in population dynamics studies give only a noisy output of the demographic process under investigation, the SSM framework provides a flexible and integrated framework for fitting a wide range of models which, with widespread adoption, has the potential to advance significantly ecological statistics (Buckland et al., 2004; Thomas et al., 2005).

5 Acknowledgments

O. Gimenez's research was supported by a Marie-Curie Intra-European Fellowship
within the Sixth European Community Framework Programme. This project was
funded by the Action Incitative Régionale BioSTIC-LR 'Modélisation intégrée en dynamique des populations : applications à la gestion et à la conservation'. The authors
would like to warmly thank J.-D. Lebreton for his support during this project. We also
thank a referee whose comments help improving a previous draft of the paper.

6 Literature Cited

Anderson-Sprecher, R., Ledolter, J., 1991. State-space analysis of wildlife telemetry
data. Journal of the American Statistical Association. 86, 596–602.

Arnason, A.N., 1973. The estimation of population size, migration rates and survival
in a stratified population. Research in Population Ecology. 15, 1–8.

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.

- Besbeas, P., Lebreton, J.D., Morgan, B.J.T., 2003. The efficient integration of abundance and demographic data. Journal of the Royal Statistical Society Series C Applied
 Statistics. 52, 95–102.
- ¹⁹ Brooks, S. P., Catchpole, E. A., Morgan, B. J. T., 2000. Bayesian animal survival
 ²⁰ estimation. Statistical Science. 15, 357–376.
- ²¹ Brownie, C., Anderson, D.R., Burnham, K.P., Robson, D.S., 1985. Statistical infer²² ence from band recovery data a handbook. United States Fish and Wildlife Service,
- ²³ Washington.

- ¹ Buckland, S.T., Newman, K.B., Thomas, L., Koesters, N.B., 2004. State-space models
- ² for the dynamics of wild animal populations. Ecological Modelling. 171, 157–175.
- ³ Catchpole, E.A., Freeman, S.N., Morgan, B.J.T., Harris, M.P., 1998. Integrated recov-
- ⁴ ery/recapture data analysis. Biometrics. 54, 33–46.
- ⁵ Choquet, R., Reboulet, A.-M., Pradel, R., Gimenez, O., Lebreton, J.-D., 2004. M⁶ SURGE: new software specifically designed for multistate capture-recapture models.
- 7 Animal Biodiversity and Conservation. 27, 207–215.
- ⁸ Clark, J.S., 2005. Why environmental scientists are becoming Bayesians. Ecology
 ⁹ Letters. 8, 2–14.
- ¹⁰ Clark, J.S., Ferraz, G.A., Oguge, N., Hays, H., DiCostanzo, J., 2005. Hierarchical Bayes
- ¹¹ for structured, variable populations: From recapture data to life-history prediction.
 ¹² Ecology. 86, 2232–2244.
- ¹³ Cormack, R. M., 1964. Estimates of survival from the sighting of marked animals.
 ¹⁴ Biometrika. 51, 429–438.
- ¹⁵ De Valpine, P., 2004. Monte Carlo state-space likelihoods by weighted posterior kernel
- ¹⁶ density estimation. Journal of the American Statistical Association. 99, 523–536.
- ¹⁷ Dupuis, J., 1995. Bayesian estimation of movement and survival probabilites from
 ¹⁸ capture-recapture data. Biometrika. 82, 761–772.
- ¹⁹ Ellison, A.M., 2004. Bayesian inference in ecology. Ecology Letters. 7, 509–520.
- ²⁰ Gimenez, O., Bonner, S., King, R., Parker, R. A., Brooks, S., Jamieson, L. E., Grosbois,
- ²¹ V., Morgan, B. J. T., Thomas, L., submitted. WinBUGS for population ecologists:
- ²² Bayesian modeling using Markov Chain Monte Carlo methods. Environmental and
- 23 Ecological Statistics.
- 24 Gelman, A., 1996. Inference and monitoring convergence. In Gilks, W. R., Richard-
- ²⁵ son, S. Spiegelhalter, D. J. (eds), Markov chain Monte Carlo in practice, pp 131–143.

- ¹ Chapman and Hall, London.
- ² Harvey, A.C., 1989. Forecasting, structural time series models and the Kalman Filter.
- ³ Cambridge University Press, Cambridge.
- ⁴ Huggins, R., 2001. A note on the difficulties associated with the analysis of capture-
- ⁵ recapture experiments with heterogeneous capture probabilities. Statistics and Proba-
- ⁶ bility Letters. 54, 147–152.
- ⁷ Ihaka, R., Gentleman R., 1996. R: a language for data analysis and graphics. Journal
- ⁸ of Computational and Graphical Statistics. 5, 299–314.
- Jolly, G. M., 1965. Explicit estimates from capture-recapture data with both death and
 immigration-stochastic model. Biometrika. 52, 225–247.
- ¹¹ 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. Ecological Monograph. 62, 67–118.
- Lebreton, J.D., Almeras, T., Pradel, R., 1999. Competing events, mixtures of information and multistrata recapture models. Bird Study. 46, 39–46.
- Lebreton, J.D., Pradel, R., 2002. Multistate recapture models: modelling incomplete
 individual histories. Journal of Applied Statistics. 29, 353–369.
- Millar, R.B., Meyer, R., 2000. Non-linear state space modelling of fisheries biomass
 dynamics by using Metropolis-Hastings within-Gibbs sampling. Journal of the Royal
- ²⁰ Statistical Society Series C Applied Statistics. 49, 327–342.
- Newman, K. B. 1998. State-space modeling of animal movement and mortality with
 application to salmon. Biometrics. 54, 1290–1314.
- ²³ Pollock, K.H., 1991. Modeling capture, recapture, and removal statistics for estimation
- ²⁴ of demographic parameters for fish and wildlife populations past, present, and future.
- ²⁵ Journal of the American Statistical Association. 86, 225–238.

- Pollock, K. H., 2002. The use of auxiliary variables in capture-recapture modelling: an
 overview. Journal of Applied Statistics. 29, 85–102.
- ³ Pradel, R., 2005. Multievent: An extension of multistate capture-recapture models to
- ⁴ uncertain states. Biometrics. 61, 442–447.
- ⁵ Schwarz, C.J., Schweigert, J.F., Arnason, A.N., 1993. Estimating migration rates using
- ⁶ tag-recovery data. Biometrics. 49, 177–193.
- ⁷ Schwarz, C. J., Seber, G. A. F., 1999. Estimating animal abundance: review III.
 ⁸ Statistical Science. 14, 427–56.
- Seber, G. A. F., 1965. A note on the multiple-recapture census. Biometrika. 52,
 249-259.
- ¹¹ Spiegelhalter, D. J., Thomas, A., Best, N. G., Lunn, D., 2003. WinBUGS user manual
 ¹² Version 1.4. MRC Biostatistics Unit, Cambridge, UK.
- ¹³ Sturtz, S., Ligges, U., Gelman, A., 2005. R2WinBUGS: a package for running Win-
- ¹⁴ BUGS from R. Journal of Statistical Software. 12, 1–16.
- ¹⁵ Thomas, L., Buckland, S.T., Newman, K.B., Harwood, J., 2005. A unified framework
- ¹⁶ for modelling wildlife population dynamics. Australian and New Zealand Journal of
 ¹⁷ Statistics. 47, 19–34.
- ¹⁸ Vaupel, J.W., Yashin, A.I., 1985. Heterogeneity ruses some surprising effects of
 ¹⁹ selection on population dynamics. American Statistician. 39, 176–185.
- ²⁰ Wang, G., 2006. On the latent state estimation of nonlinear population dynamics using
- ²¹ Bayesian and non-Bayesian state-space models. Ecological Modelling. doi: 10.1016/
- 22 j.ecolmodel.2006.09.004.
- ²³ Williams, B. K., Nichols, J. D., Conroy, M. J., 2002. Analysis and management of
 ²⁴ animal populations. Academic Press, San Diego.

¹ Appendix A: WinBUGS code for fitting the CJS ² model using the SSM formulation

```
3
  # MODEL
                                                      #
  # State-space formulation of the Cormack-Jolly-Seber model
                                                      #
5
  # observations = 0 (non-encountered) and 1 (encountered)
                                                      #
6
  # states = 0 (dead) and 1 (alive)
                                                      #
7
  model
a
  {
10
  # Define the priors for survival phi and detectability p
11
12 p[1] <- 1
13 phi[1] <- 1
14 for (j in 2:K)
  {
15
 phi[j] ~ dbeta(1,1)
16
 p[j] ~ dbeta(1,1)
17
  }
18
  # Define the SYSTEM PROCESS
19
  for (i in 1:n)
20
    {
21
    # if first capture
22
   PrX[i,e[i]] <- 1 # Pr(alive | first capture) = 1</pre>
23
    X[i,e[i]+1] ~ dbern(PrX[i,e[i]]) # alive (a 1 is generated with certainty)
24
```

```
PrO[i,e[i]] <- X[i,e[i]+1] # detection probability at initial detection is 100%</pre>
1
    # otherwise
2
        for (j in (e[i]+1):K)
3
        {
4
        PrX[i,j] <- phi[j] * X[i,j]</pre>
5
        X[i,j+1] ~ dbern(PrX[i,j])
6
        PrO[i,j] <- p[j] * X[i,j+1]</pre>
7
        }
8
    # fullfil the remaining cells with zeros
9
        for (j in 1:(e[i]-1))
10
        {
11
       PrX[i,j] <- 0
12
       X[i,j] <- 1
13
       PrO[i,j] <- 0
14
        }
15
    }
16
  # Define the OBSERVATION PROCESS
17
  for (h in 1:nx)
18
    {
19
    data[h,3] ~ dbern(PrO[data[h,1],data[h,2]])
20
    }
21
  }
22
  23
 # DATA
24
  # 'K' is the number of encounter occasions
25
```

1 # 'n' is the number of individuals

2 # 'nx' is 'K' times 'n'

3 # 'e' is the vector of first encounters ('n' components)

4 # 'data' is a matrix with dimensions 'nx' times 3 where

5 # the first column gives the current individual (1,...,'n'),

 $_{6}$ # the second column gives the current encounter occasion (1,...,'K'),

7 # the third column gives the observation (= 1 if detection, = 0 otherwise)

8 # corresponding to the current individual and current encounter occasion

¹⁰ Appendix B

Let $Z_{i,t} = [X_{i,t-1}, X_{i,t}]$ be a bivariate random vector where its two components are denoted $Z_{i,t}^1$ and $Z_{i,t}^2$. Equation (8) becomes

$$Y_{i,t}|Z_{i,t} \sim Bernoulli\left((Z_{i,t}^1 - Z_{i,t}^2)\lambda_{i,t}\right)$$
(10)

¹⁴ and Equation (9) becomes

$$Z_{i,t+1}|Z_{i,t} = \begin{cases} \left(Z_{i,t+1}^{1}|Z_{i,t}\right) = Z_{i,t}^{2} \\ Z_{i,t+1}^{2}|Z_{i,t} \sim Bernoulli(Z_{i,t}^{2}\phi_{i,t}). \end{cases}$$
(11)

15

13

¹⁶ The system defined by Equation (10) and Equation (11) is a state-space model and it ¹⁷ is equivalent to the model defined by Equation (8) and Equation (9).

Note that an alternative state-space formulation can be adopted using a multistate
formulation of the RR model (Lebreton et al., 1999) and Section 3.1.



Figure 1: Posterior distributions for the survival and detection probabilities using the CJS model applied to the Dipper data set as estimated by the state-space model and MCMC methods.

Table 1: Estimated survival and detection probabilities for the Dipper data using the CJS model and three different methods, the state-space model (SSM) using a Monte Carlo Markov Chain (MCMC) method, the product-multinomial model (PMM) using a MCMC method and the PMM using a maximum-likelihood (ML) method. The two first methods were implemented using program WinBUGS (Spiegelhalter et al., 2003), while program M-SURGE (Choquet et al., 2004) was used to implement the last one.

	SSM using MCMC	PMM using MCMC	PMM using ML
Parameter	Posterior median/mode (SD)	Posterior median/mode (SD)	MLE (SE)
ϕ_1	0.721/0.722 (0.132)	0.723/0.693 (0.132)	0.718(0.156)
ϕ_2	$0.448/0.456\ (0.071)$	$0.448/0.460\ (0.071)$	$0.435\ (0.069)$
ϕ_3	$0.480/0.493\ (0.060)$	$0.480/0.476\ (0.061)$	$0.478\ (0.060)$
ϕ_4	0.628/0.624 (0.061)	$0.627/0.616\ (0.060)$	$0.626\ (0.059)$
ϕ_5	$0.602/0.601 \ (0.057)$	$0.602/0.607 \ (0.057)$	$0.599\ (0.056)$
ϕ_6	0.713/0.640 (0.142)	0.720/0.628 (0.143)	- (-)*
p_2	$0.671/0.658\ (0.134)$	0.670/0.691 (0.134)	$0.696\ (0.166)$
p_3	$0.883/0.918\ (0.083)$	0.883/0.904 (0.083)	$0.923\ (0.073)$
p_4	0.888/0.914 (0.063)	0.889/0.912 (0.063)	$0.913 \ (0.058)$
p_5	0.882/0.885 (0.057)	0.883/0.904 (0.057)	$0.901 \ (0.054)$
p_6	0.913/0.920 (0.052)	0.912/0.935 (0.051)	0.932(0.046)
p_7	$0.735/0.724\ (0.142)$	$0.727/0.648\ (0.143)$	- (-)*
* NT · 1 · · ·			

* Non-identifiability detected.