Survival and Population Dynamics of the Marabou Stork in an Isolated Population, Swaziland

Ara Monadjem1*, Adam Kane2, Andre Botha3, Desire Dalton4,5, Antoinette Kotze4,5

1 All Out Africa Research Unit, Department of Biological Sciences, University of Swaziland, Kwaluseni, Swaziland, 2 Department of Zoology, School of Natural Sciences, Trinity College Dublin, Dublin, Ireland, 3 Birds of Prey Programme, Endangered Wildlife Trust, Modderfontein, South Africa, 4 National Zoological Gardens of South Africa, Pretoria, South Africa, 5 Genetics Department, University of the Free State, Bloemfontein, South Africa

Abstract

Investigating the ecology of long lived birds is particularly challenging owing to the time scales involved. Here an analysis is presented of a long term study of the survival and population dynamics of the marabou stork (Leptoptilos crumeniferus), a wide ranging scavenging bird from Sub-Saharan Africa. Using resightings data of tagged nestlings and free flying birds we show that the stork population can be divided into three general life stages with unique survival probabilities and fecundities. Fecundity of the storks is inversely related to rainfall during their breeding season. Corroborative evidence for a metapopulation structure is discussed highlighting the impact of the Swaziland birds on the ecology of the species in the broader region. The importance of tag loss or illegibility over time is highlighted. Clearly, any attempt at conserving a species will require a detailed understanding of its population structure, of the sort examined here.

Citation: Monadjem A, Kane A, Botha A, Dalton D, Kotze A (2012) Survival and Population Dynamics of the Marabou Stork in an Isolated Population, Swaziland. PLoS ONE 7(9): e46434. doi:10.1371/journal.pone.0046434

Editor: Matt Hayward, Australian Wildlife Conservancy, Australia

Received June 29, 2012; Accepted August 29, 2012; Published September 28, 2012

Copyright: © 2012 Monadjem et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: The authors have no funding or support to report.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: ara@uniswaz.sz

Introduction

Survival estimates are critical parameters for various ecological models, particularly population dynamics, which in turn inform conservation [1]. This is particularly important for long-lived vertebrates which may take several years to mature into breeding adults [2], [3]. However, survival estimates are often not available for long-lived birds such as storks; the only species for which robust estimates of both subadults and adults are available is the white stork Ciconia ciconia [4], [5], [6]. Survival in the marabou stork Leptoptilos crumeniferus has been crudely estimated by comparing the proportion of immature to adult birds [7], however robust estimates of survival must be based on following marked individuals through time [8].

The marabou stork is a widespread scavenging bird occurring in savanna habitats throughout sub-Saharan Africa [9]. Despite its extensive distributional range, this species breeds at a rather limited number of localities [10]. For example, despite having been recorded across most of eastern, central and northern South Africa, it has not yet successfully bred in that country [11]. In fact, in all of southern Africa, this species breeds at just one site in Swaziland [12] (Figure 1), one site in Botswana [13] and less than a dozen scattered sites in Zimbabwe [14]. Hence this species has a metapopulation structure [15] with breeding birds concentrated in discrete colonies (Figure 2). This raises the possibility that the breeding populations act as sources for the South African sink. Dispersing birds can cover extensive distances of up to 1500 km thus allowing for dynamics of this sort to occur [10].

One way to infer a metapopulation structure for this species would be through a population dynamics model. Such models require a number of parameters for them to have any predictive power, notably probability of survival and fecundity rates [16]. Moreover, in most species these parameters will change as they age; a first year marabou stork, for instance, is not going to be involved in reproduction [17]. This is the shortcoming of exponential or geometric decay models of population growth which treat all life stages equally, resulting in a serious oversimplification [18]. An extensive knowledge of the breeding ecology and demographics of marabou storks is therefore necessary before such population dynamics modelling can be conducted.

The aim of this paper is to elucidate the population ecology of the marabou stork in southern Africa. It is hypothesized that survival is age dependent and that this species has a positive population growth rate. The objectives of this paper are to:

1) Relate fecundity of marabou storks breeding in Swaziland over a nine year period with climatic conditions,
2) Estimate survival rates based on resightings of tagged nestlings and free flying marabou storks in southern Africa,
3) Use the breeding data and survival estimates of the breeding population in Swaziland to model the population dynamics of this species.

Materials and Methods

Study area

The collection of breeding data and the tagging of nestlings was conducted in the 16 000 ha Hlane National Park (31°53’S, 26°18’E), Swaziland. The vegetation is dominated by knobthorn Acacia nigrescens woodland interspersed with riverine forest. The
climate is subtropical [see [12], [19] for more details on topography and climate]. The marabou stork breeding season covers the austral winter: 2–3 eggs are laid in May or June; eggs hatch after approximately 30 d incubation, usually in June or July, which is the coldest time of year; and the chicks fledge in October [20], [10], [19], [21].

Free-flying marabou storks were captured and tagged at the Moholoholo Wildlife Rehabilitation Centre (24°30’S; 30°54’E, 600 m above sea level), north-eastern South Africa. The site is situated in low-lying savanna, to the west of the Kruger National Park.

There is no Ethics Committee at the University of Swaziland to oversee the compliance of biological studies. However, the current study was conducted under a permit from the Swaziland National Trust Commission to Ara Monadjem. As outlined in our methods, the only contact we had with the study animals was tagging 210 birds based on a standard technique applicable throughout the World. Furthermore, we did not harm or compromise the health of any species during this study.

Patagial tagging was implemented as the preferred method of colour-marking for large raptors, vultures and storks by the Birds of Prey Programme of the Endangered Wildlife Trust in 2006. An extensive review and assessment process of a range of methods was conducted over a period of 18 months before the members and associates of the Programme agreed on and approved the tagging protocol for the use of this method at its Annual Conference [22]. The review included a clinical assessment of the condition of captive and recaptured free-flying tagged birds by independent veterinarians after tagging and no signs of infection or other potentially negative symptoms resulting from the tagging have to date been found [22]. Based on the results of the survival and movements of tagged birds, the method also does not seem to have a detrimental effect on the mobility and foraging ability of other large birds e.g. vultures [23].

Data collection
Breeding marabou storks were monitored regularly between 2003 and 2011 at Hlane National Park. Originally nests were located on foot, but from 2008 we used a microlight to search for nests from the air once per year in July. Active nests (a nest on which eggs were found or adult activity was observed) were visited twice weekly until the chicks hatched or the nest failed. Fledging date was taken as the date of the last visit on which the chick was still on the nest. The mass and wing length was measured during each visit.

Free-flying birds were captured in a specially designed walk-in trap for vultures [24], [25] that was erected alongside the vulture restaurant at Moholoholo Wildlife Rehabilitation Centre. Captured birds were aged and had their mass and wing length measured. Ageing was based on plumage characteristics [26] and birds were assigned to one of three age classes: juvenile (1st year bird), subadult (2–4 years old) and five years old or older. Each bird was fitted with a metal ring issued by AFRING (Animal Demography Unit, University of Cape Town) and a patagial tag. Patagial tags were fitted according to the standard protocol adopted for this practice in southern Africa [22]. It involved the use of a double set of standard cattle tags engraved with a unique number which was fitted to the patagial area on each wing of each bird using a tag applicator. This method was extensively assessed
prior to this study and found not to be detrimental to the birds’ health or inhibiting their ability to forage [22]. All tagged marabou storks were released unharmed and within 120 min of capture.

A dedicated resightings programme was established using radio and television broadcasts, newspaper and magazine articles, and posters in Kruger National Park rest camps. A significant proportion of resightings was submitted by the staff at Moholoholo who kept a daily watch at the vulture restaurant, at which the marabou storks were frequently observed. Resightings were also reported inter alia by managers of other vulture restaurants, game ranchers, farmers and tourists.

Sexing

Only nestlings were sexed. DNA extraction was conducted using the QiagenDNAeasy® Blood and Tissue Kit. The extraction protocol as outlined in the manufacturer protocol was followed. CHD1 gene amplification was conducted using the 2550F/2718R [27] primer set at the National Zoological Gardens of South Africa. Amplification was carried out using 25 μl reaction volume and polymerase chain reaction was conducted with PromegaGo-Taq® Flexi DNA polymerase (Promega Corporation) which has a 1× buffer containing 10 milli molar (mM) Tris®-HCl (pH 9.0), 50 mM potassium chloride (KCl) and 0.1% Triton® X-100. The final reaction conditions were as follows: 1 X PCR buffer, 1.5 mM MgCl2, 200 micro molar (μM) of each 2′-deoxynucleotide triphosphate (dNTP), 5 picomol (pmol) of each of the forward and reverse primer, 0.25 unit (U) Taq DNA polymerase and 10–20 nano gram (ng) genomic DNA template. A no template control as well as positive controls for a male and female bird of known sex was included. The conditions for PCR amplification were as follows: 2 min at 95°C initial denaturation, 30 cycles for 30 s at 95°C, 30 s at 50°C and 2 min at 72°C, followed by extension at 72°C for 10 min. The PCR reaction was carried out in the BOECO TC-PRO Thermal Cycler. PCR products mixed with tracking dye were separated by electrophoresis in a 2% agarose gel for 45 min at 100 V in 1× Tris-borate-EDTA buffer.

Data analysis

Fecundity is a measure of reproductive success and was defined as the number of fledglings successfully raised per pair per annum [28]. A Pearson’s correlation was used to test whether fecundity was related to rainfall in the preceding summer (October to March, inclusive) and rainfall during the winter breeding season (May to September), as these variables had previously been shown to be important for marabou stork breeding [19].

The program MARK was used to estimate survival and recapture of marabou storks using the standard Cormack-Jolly-Seber model [29], [30]. A variety of models that included time dependence, sex and age were developed. Models were ranked using Akaike’s Information Criterion corrected for small sample size (AICc) [31]. The model with the lowest AICc was deemed the best model; where ΔAICc, for any two (or more) models was <2.0, they were both deemed to be equally good.

Survival was estimated separately for the birds tagged as nestlings from those tagged as free-flying. A subset (n = 100, tagged between 2008 and 2011) of those tagged as nestlings were sexed, and were used to test for the role of sex in survival of marabou storks. Models which included sex performed poorly compared with those that did not (ΔAICc >2.9) and hence sex was removed as a factor. Subsequently, the data for the sexes of the nestlings were pooled and only age and time dependence were included.

To test for violations of the assumption of homogeneity of survival and recaptures, GOF (goodness of fit) tests were conducted in the program Release [32]. Test 2 tests the assumption of equal catchability (in our case, resightings) of marked individuals. Test 3 tests the assumption that all individuals have equal probability of survival independent of when they were marked.

Model Building

Leslie Matrices are used to chart the development of a population over time by separating the given group into distinct age classes, with matrix elements representing probability of survival and fecundity of these classes [33]. A variation of a Leslie Matrix was developed to investigate the population dynamics of a marabou stork population. The model used information from females only as they are responsible for reproduction in the population [34] and assumes that the above parameters are constant for each age class. The number of females in each class is given by a column vector \( \mathbf{N}_t \), such that the number determined at time \( t+1 \) is given by \( \mathbf{N}_{t+1} = \mathbf{L}\mathbf{N}_t \), where \( \mathbf{L} \) represents the Leslie Matrix. This process is repeated for each time step. In this case \( t \) represents one year. This gives the following general equation:

\[
\begin{bmatrix}
  n_1(t+1) \\
  n_2(t+1) \\
  \vdots \\
  n_s(t+1)
\end{bmatrix} =
\begin{bmatrix}
  f_1 & f_2 & \cdots & f_s \\
  s_1 & 0 & \cdots & 0 \\
  \vdots & 0 & \ddots & \vdots \\
  0 & 0 & \cdots & s_{s-1}
\end{bmatrix}
\begin{bmatrix}
  n_1(t) \\
  n_2(t) \\
  \vdots \\
  n_s(t)
\end{bmatrix}
\]

where “\( f_i \)” is fecundity and “\( s_i \)” is the probability of survival from one age class to the next. The dominant eigenvalue of the matrix is the \( \lambda \) value and represents the growth rate of the population. If \( \lambda > 1 \), the population is growing; 0 < \( \lambda < 1 \), it is decreasing and \( \lambda = 1 \), the population is stable. The right eigenvector of a given matrix represents the stable age distribution of the group i.e. the proportion of birds in each stage. This is a measure of the contribution the life stages have to overall population growth [35]. So although juveniles do not reproduce they still make a contribution here.

A variation of the Leslie Matrix known as the Lefkovich Matrix was used in order to separate the storks into distinct life stages [16]: juveniles, subadults and adults,

\[
\begin{bmatrix}
  P_1 & F_2 & F_3 & F_4 \\
  G_1 & P_2 & 0 & 0 \\
  0 & G_2 & P_3 & 0 \\
  0 & 0 & G_3 & P_4 \\
  0 & 0 & 0 & G_4
\end{bmatrix}
\]

\( G_x \) (on the subdiagonal) is the probability of surviving for a year and moving into the next stage; \( P_x \) (on the diagonal) is the probability of surviving for a year and remaining in the same stage; \( F_x \) represents fecundity. The model did not take into account density dependent effects. The matrix was created in MS Excel using a variation of a template developed by Spangenberg and Jungck [33] (see results section for the values used to parameterize the model). Seeing as subadult and adult birds can remain in their respective stages for more than a single year this was corrected for accordingly with the following equations,

\[
p_i = \frac{1 - (s_i)^{y_i-1}}{1 - (s_i)^y_i} \cdot s_i
\]
The probability of remaining in an age class at next year is given by $p_i$ and $q_i$ is the probability of moving up an age class at next year. $s_i$ is the survivor rate for $i^{th}$ year and $d_i$ is the length of time spent in this $i^{th}$ stage (taken from [16]).

The below matrix was developed using the values from our results which corrected the original parameters for stage duration and fecundity,

$$q_i = \left(\frac{(s_i)^{d_i} (1-s_i)}{1-(s_i)^{d_i}}\right)$$

This matrix has the three defined age classes i.e. the columns (there is a small non-zero probability that the birds will live past the third stage and fall into a fourth category but any individuals here have zero probability of surviving to the next time step). Subadult birds spend years two to four in their stage and adults spend from age five to 25 in their stage. This assumes a life expectancy of 25 years for wild marabou storks [36], [11]. The fecundity value shown here was taken as an average from 2003–2011 (see Figure 3). The model was run with a varying starting population of three to 10 adult females; the lower number represents a minimum estimate of the nesting population in Swaziland during the early 1960s [37]. However, seeing that nesting sites can easily be overlooked by observers on the ground we also ran the model with a starting population of up to 10 females.

### Results

Between 19 and 31 pairs of marabou storks bred annually at Hlane National Park that fledged between 11 and 43 chicks. The fecundity of marabou storks breeding at Hlane National Park differed greatly between years (Figure 3), with a mean of 1.05 fledged offspring per pair per annum. There was no correlation between fecundity and rainfall in the previous summer ($r = 0.401$, $df = 7$, $P = 0.284$), however, there was a significant inverse correlation with rainfall during the breeding season ($r = -0.785$, $df = 9$, $P = 0.012$).

A total of 193 nestlings and 17 free-flying marabou storks have been tagged since 2005 and which have been resighted 811 and 834 times, respectively. On fledging, male marabou storks had significantly larger mean ($\pm$ SE) wing lengths ($663.1 \pm 4.63$ mm vs $596.1 \pm 5.85$ mm; $t = 8.98$, $P < 0.0001$, $DF = 98$) and heavier mean ($\pm$ SE) masses ($7437 \pm 110$ g vs $6155 \pm 114$ g; $t = 8.09$, $P < 0.0001$, $DF = 98$) than females. Free-flying birds had mean ($\pm$ SE) wing length of $700.7 \pm 7.88$ mm ($n = 17$), and mean ($\pm$ SE) mass of $5038 \pm 291$ mm ($n = 13$).

In the nestling analysis covering the seven years between 2005 and 2011, survival of nestlings post-fledging was age-dependent. The best model for survival had three age classes, 1st year birds, 2nd–4th year birds and $5^{th}$ year birds (Table 1). The next model had $\Delta AICc < 2$ compared with the best model, and had survival separated into five age classes. For both of these models recapture rates were age and time independent (Table 1). The next three

![Figure 3. The relationship between fecundity of marabou storks and rainfall between 2003 and 2011 in Swaziland.](https://doi.org/10.1371/journal.pone.0046434.g003)
These models were separated into three age classes. In the top three models, recapture rates were age dependent, whereas in the fourth best model recaptures were independent of both age and time (Table 2). Again, the test 2 and test 3 results were statistically not significant (chi-square test, P > 0.05), showing that the assumptions tested had not been violated.

The survival rates of marabou storks varied considerably between age classes and the datasets analysed (Table 3). Typically, first year survival was lower than that of older birds. Based on the analysis of birds tagged as nestlings, survival to the end of the first year was 0.6440. However, based on the analysis of free-flying birds, the survival of 1st year birds was only 0.2500. Survival of older age classes (subadults and adults) was generally high ranging from 0.7917 to 0.8727 (Table 4). The only exception was age 5th year birds tagged as nestlings where survival dropped to 0.2193, suggesting loss or fading of tags.

The resulting λ value was 1.0212 indicating a population increase over time (see Figure 4). The population in this model developed from exclusively adults to an age structure which stabilizes at 43.8% adults, 33.4% subadults and 22.5% juveniles (Table 5). The reproductive value increases with age in marabou storks with the adult stage being the highest (see Table 5).

The final population in the model is clearly dependent on the initial number of founding individuals and the fecundity of the storks. Starting with three females a total population of 26 birds is developed from exclusively adults to an age structure which stabilizes at 43.8% adults, 33.4% subadults and 22.5% juveniles (Table 5). The reproductive value increases with age in marabou storks with the adult stage being the highest (see Table 5).

The number of parameters is indicated by ‘n’. The models are arranged from best (top of table) to worst (bottom).

doip:10.1371/journal.pone.0046434.t001

doip:10.1371/journal.pone.0046434.t002

doip:10.1371/journal.pone.0046434.t003

doip:10.1371/journal.pone.0046434.t004

doip:10.1371/journal.pone.0046434.t005

Table 1. The candidate models used to estimate survival in nestling marabou storks tagged in Swaziland between 2005 and 2011, and resighted across southern Africa.

| Model | AICc | Delta AICc | AICc Weights | N |
|-------|------|------------|--------------|---|
| phi(age1, ≥2) p(t) | 52.0191 | 0.29804 | 3 |
| phi(age1, 2, ≥3) p(age1, ≥2) | 52.554 | 0.5349 | 4 |
| phi(age1, ≥2) p(t) | 52.5604 | 0.5413 | 4 |
| phi(age1, ≥2) p(t) | 53.2647 | 1.2456 | 4 |
| phi(age1, ≥2) p(t) | 59.7627 | 7.4736 | 6 |
| phi(age1, ≥2) p(t) | 66.2284 | 14.093 | 8 |
| phi(age1, ≥2) p(t) | 79.696 | 27.6769 | 11 |

Estimates of survival (phi) and recapture (p) were modelled with time (t), and/or age class of the birds (age). Age1 refers to age classes of 1st year birds, age2 to subadults (2nd to 4th year birds) and age3 to adults (≥5th year birds). The inclusion or exclusion of interactions in the models is symbolized by (∗) or (+), respectively. The number of parameters is indicated by ‘n’. The models are arranged from best (top of table) to worst (bottom).

doip:10.1371/journal.pone.0046434.t002

Table 2. The candidate models used to estimate survival in free-flying marabou storks tagged in South Africa between 2007 and 2011, and resighted across southern Africa.

| Model | AICc | Delta AICc | AICc Weights | n |
|-------|------|------------|--------------|---|
| phi(age1, 2–4, ≥5) p(t) | 487.751 | 0.13917 | 4 |
| phi(age1, 2, 3, 4, ≥5) p(t) | 490.1439 | 0.11435 | 5 |
| phi(age1, 2, 3, 4, ≥5) p(age1, ≥2) | 492.1082 | 2.3572 | 6 |
| phi(age1, 2, 3, 4, ≥5) p(t) | 494.1198 | 4.3688 | 6 |
| phi(age1, ≥2) p(.) | 494.3246 | 4.5736 | 10 |
| phi(t) p(.) | 495.4601 | 5.7091 | 7 |
| phi(age1, ≥2) p(.) | 496.0157 | 6.2647 | 4 |
| phi(t) p(.) | 496.0794 | 6.3284 | 10 |
| phi(age1, ≥2) p(age1, ≥2) | 496.2424 | 6.4914 | 4 |
| phi(age1, 2, 3, 4, ≥5) p(.) | 497.811 | 8.0600 | 5 |
| phi(age1, ≥2+t) p(.) | 500.4836 | 10.7326 | 11 |
| phi(age1–2, ≥3+t) p(.) | 502.8836 | 13.1353 | 11 |
| phi(age1–2, ≥3+t) p(.) | 505.8063 | 16.0553 | 15 |

Estimates of survival (phi) and recapture (p) were modelled with time (t), and/or age class of the birds (age). Age1–5 refers to age classes of 1st year birds through to 5th year birds. The number of parameters is indicated by ‘n’. The models are arranged from best (top of table) to worst (bottom).

doip:10.1371/journal.pone.0046434.t003

Table 3. Survival and recapture (resightings) rates of marabou storks tagged as nestlings and free-flying adults in South Africa between 2005 and 2011.

| Analysis | Estimated parameter | rate | SD |
|----------|---------------------|------|----|
| Nestlings Survival of 1st year birds | 0.6440 | 0.0765 |
| Nestlings Survival of 2nd–4th year birds | 0.7917 | 0.0597 |
| Nestlings Survival of ≥5th year birds | 0.2193 | 0.1459 |
| Nestlings Recapture rate | 0.4226 | 0.0523 |
| Free-flying Survival of 1st year birds | 0.2500 | 0.2165 |
| Free-flying Survival of ≥ subadult birds | 0.8727 | 0.2483 |
| Free-flying Recapture of 1st year birds | 0.9999 | 0.0004 |
| Free-flying Recapture of ≥ subadult birds | 0.2545 | 0.1559 |

The analysis refers to the specific dataset used for the estimates: nestlings=all nestlings tagged and resighted between 2005 and 2011; free-flying=all free-flying birds tagged and resighted between 2007 and 2011. See Methods for further details.

doip:10.1371/journal.pone.0046434.t004

Table 4. Parameters used in matrix.

| Stage | Fx | Px | Gx |
|-------|----|----|----|
| Juveniles | 0 | 0 | 0.644 |
| Subadults | 0 | 0.587 | 0.205 |
| Adults | 0.525 | 0.865 | 0.008 |

Fx = Fecundity; Px = Probability of remaining in age class at next year; Gx = Probability of moving up an age class at next year.
Discussion

This study presents the first estimates of survival for the marabou stork based on resightings of tagged individuals that also accounts for recapture (resightings) probability. A previous study estimated the survival of marabou storks based on the proportion of immature birds to adult birds, yielding survival rates of 28% for first year birds, 72% for subadults and 92% for adults [7], [38]. Our estimates are lower than these for adults but higher for juvenile and subadult birds. Our estimate for sub-adult survival mentioned above also matches those for the wood stork, *Mycteria americana* [39]. From the tagging of nestlings, our estimate of 1st year mean survival is 64%. This is higher than survival rates for 1st year white storks estimated at 47% [40], 48% [4] and 33–42% [41], and for 1st year wood storks at 44% [42]. This is not surprising since the marabou stork is larger than these two species and survival rates are directly related to size [38]. However, the estimate of 1st year survival based on the tagging of free-flying birds was exceptionally low (25%), lower than that reported for the white or wood storks. Undoubtedly, this estimate suffers from small sample sizes as only four free-flying 1st year birds were captured and tagged compared with 193 nestlings.

Adult survival was higher, but the specific value depended on the dataset that was used. Based on the free-flying birds the survival rate was 87% and apparently did not differ between subadults and adults. Subadult (2nd–4th year birds) mean survival rates based on resightings of tagged nestlings was 79%, with adult (≥5th year birds) survival rate dropping to 22%. This clearly is not an accurate reflection of adult survival, where captive marabou storks may live to 31 years [11], but may be related to tag loss (see below). The annual survival of adult white storks in Europe exhibited considerable inter-annual variation over a 19-year period ranging from a low of about 70% to almost 100% [41]. By contrast, mean annual survival rates in declining populations of white storks in Europe ranged between 58% and 75% [4].

Survival rates of the marabou stork were not time-dependent. In contrast, the variation in survival rates of the white stork has been linked to rainfall in the Sahel where this species migrates to in the northern winter [40]. The marabou stork is not a migrant [11] which may explain the lack of time-dependence in this species. The lack of time-dependent survival, however, is still surprising in the marabou stork, especially since reproductive success of this species is strongly related to rainfall [19]. The current study corroborated the findings of a previous study [19] showing fecundity to be negatively related to rainfall during the breeding season. The reasons for this have previously been argued to be related to food availability and foraging efficiency of the parents [19] rather than due to the inability of the chicks to thermoreg-

![Graph showing the growth of the total population and each component stage over a 50 year period commencing in 1962.](image)

**Figure 4.** Graph showing the growth of the total population and each component stage over a 50 year period commencing in 1962. The initial population was set to 10 females in this case. $\lambda = 1.0212$. doi:10.1371/journal.pone.0046434.g004

| Stage     | Stable Age Distribution | Reproductive Value |
|-----------|-------------------------|--------------------|
| Juveniles | 0.2251                  | 1.00               |
| Subadults | 0.3337                  | 1.59               |
| Adults    | 0.4378                  | 3.36               |

**Table 5.** Values for the stable age distribution and the reproductive value of the life stages as given by the right and left eigenvectors of the matrix respectively.

doi:10.1371/journal.pone.0046434.t005
ulate in cold and cloudy conditions [43]. Marabou storks are scavengers feeding on a wide range of food resources including carrion from large mammal carcasses, aquatic vertebrates and human waste [7], [17], [44]. The exact nature of the relationship between rainfall, food and marabou stork fecundity remains unclear, but we suggest the following. Since rainfall is necessarily linked with cloud cover, increased rainfall during the breeding season represents increased cloud cover during this period of food stress when parents need to fend for themselves as well as growing chicks. However, marabou storks require thermals for foraging [17], which are only available on sunny days, and hence increased cloud cover may result in less soaring time and less efficient foraging.

There are a number of hypotheses that have been put forward to explain the relationship between rainfall and breeding success. One suggestion is that high rainfall impedes the formation of thermals on which the Marabous soar while they forage [19]. Another is that rainfall impacts on breeding success indirectly by influencing food availability for the storks. During the breeding season freshwater fishes make up a significant portion of the storks’ diet [44]. Indeed the nestlings were observed to regurgitate fish on which they had recently been fed [19]. Low rainfall would grant the birds easier access to fish stocks as they wade through rivers and streams with lower water levels [19].

There was no apparent difference in the survival of male and female marabou storks. A similar result has been reported for the white stork [4]. However, female wood stork fledglings had survival rates of up to five times higher than in the larger males [41]. In line with a previous study [21], male marabou storks had larger masses and wing lengths than female storks. Larger size may be correlated with larger mortality, especially under difficult environmental conditions [45]. This, however, was not the case in this study.

Tag loss or fading is a serious violation of the assumptions of capture-mark-recapture studies [46], [47]. In our study, tag loss seems to have been a factor in the older age classes (≥5th year birds). A similar resightings study of African white-backed vultures fitted with the same tags showed that they were fading and become illegible from about 5–6 years of age and older [23]. In this study, we report a sudden and significant drop in survival rate from the age of 5 years and above, suggesting that the same fate befell these birds. A similar resightings study of African white-backed vultures (Gyps africanus) had reported a sudden and significant drop in survival rate from the age of 5 years and above, suggesting that the same fate befell these birds. A similar resightings study of African white-backed vultures (Gyps africanus) had reported [23].

Marabou storks are capable of dispersing up to 1500 km after fledging, easily allowing them to cover the relatively short distance (<100 km) between the study site in Swaziland and Kruger National Park in South Africa [10] (see Figure 2).

The high reproductive value of adult marabou storks is worth highlighting. This value represents the contribution that birds at any stage make to population growth. Our analysis showed that adults contributed twice as much as subadults and over three times as much as juveniles to the growth of the population. The population growth of wood storks was similarly shown to be highly sensitive to adult survivorship [39]. Clearly, this is an important point to recognize for conservation efforts targeting long-lived species [52]. The inverse relationship between survival and reproductive value means that the emphasis should be on ensuring the survival of adults.

Acknowledgments

We thank Mickey Reilly for permission to study marabou storks at Hlane National Park and for logistical support. Mduzzi Ngwenya, Martha Surridge, Elaine Franklin, Marie Dahl Gydesen and various All Out Africa volunteers assisted with the finding of nests and processing of the chicks. Dave Ducasse has flown various assistants annually over the study area in a microlight in search of marabou stork nests since 2000.

Author Contributions

Conceived and designed the experiments: AM AB. Performed the experiments: AM AB. Analyzed the data: AM A. Kane. Contributed reagents/materials/analysis tools: AM AB DD A. Kotze. Wrote the paper: AM A. Kane.

References

1. Tucker AD, Gibbons JW, Greene JL (2001) Estimates of adult survival and migration for diamondbacked turtles: conservation insight from local extirpation within a metapopulation. Can J Zool 79: 2199–2209.
2. Heppell SS (1998) Application of life-history theory and population model analysis to turtle conservation. Copeia 1998: 367–375.
3. Piper SE, Boshof AF, Scott HA (1999) Modelling survival rates in the Cape Griffon Gyps coprotheres, with emphasis on the effects of supplementary feeding. Bird Study 46: S230–238.
4. Kanyamibwa S, Bairlein E, Schierrer A (1993) Comparison of survival rates between populations of the white stork Ciconia ciconia in Central Europe. Ornis Scandinavica 24: 297–302.
5. Sarther B-E, Gretan V, Tryjanowski P, Barbraud C, Engen S, et al. (2006) Climate and spatio-temporal variation in the population dynamics of long distance migrant, the white stork. J Anim Ecol 75: 80–90.

6. Neuvoux M, Barbraud J-M, Barbraud C (2008) Nonlinear impact of climate on survival in a migratory white stork population. J Anim Ecol 77: 1143–1152.

7. Pomeroy DE (1977) The biology of marabout storks in Uganda. 1. Some characteristics of the species, and the population structure. Aardea 65: 1–24.

8. Lebreton J-D, Burnham KP, Clobert J, Anderson DR (1992) Modelling survival and testing biological hypotheses using marked animals: a unified approach with case studies. Ecological Monographs 62: 67–118.

9. Brown LH, Urban EK, Newman K (1982) The Birds of Africa, Vol. 1. Academic Press, London.

10. Monadjem A, Bamford AJ, Rasmussen M (2000) Dispersal of juvenile Marabou Storks Leptoptilos crumeniferus as determined by sightings. Ostrich 79: 187–189.

11. Anderson MD (2005). Marabou Stork Leptoptilos crumeniferus. Pp 626–627. In: Hockey, PAR, Dean, WRJ Ryan, PG (eds). Roberts – Birds of Southern Africa. VII th ed. The Trustees of the John Voelcker Bird Book Fund, Cape Town.

12. Monadjem A (2005) Breeding biology of the Marabou Stork (Leptoptilos crumeniferus) in Swaziland. Ostrich 76: 90–94.

13. Tarboton W (2001) A guide to the nests and eggs of southern African birds. Struik, Cape Town.

14. Mundy PJ (2003) Breeding of Marabou Storks. Honeyguide 49: 148–150.

15. Hanski I (1998) Metapopulation dynamics. Nature 396: 41–49.

16. Crouse DT, Crowder LB, Caswell H (1987) A Stage-Based Population Model for Loegerhead Sea Turtles and Implications for Conservation. Ecology 68: 1412–1423.

17. Pomeroy DE (1976) The biology of marabout storks in Uganda, 2. Breeding biology and general review. Adrea 66: 1–23.

18. Pastor J (2008) Mathematical ecology of populations and ecosystems: Wiley-Blackwell. 65–77 p.

19. Monadjem A, Bamford AJ (2009) Influence of rainfall on timing and success of reproduction in Marabou Storks Leptoptilos crumeniferus. Ibis 151: 344–351.

20. Monadjem A, Garcelon DK (2005) Nesting distribution of vultures in relation to land use in Swaziland. Biodiversity and Conservation 14: 2079–2093.

21. Monadjem A, Bamford AJ, Hardy ICW, Earnshaw JK, Franklin E, et al. (2010) Temporal and sex-specific variation in growth rates of Marabou Stork Leptoptilos crumeniferus chicks. Ostrich 81: 83–91.

22. Botha A (2007) A review of colour-marking techniques used on vultures in southern Africa. Vulture News 36:52–63.

23. Monadjem A, Botha A, Morn C (2012) Survival of the African white-backed vulture Gyps africanus in north-eastern South Africa. Afr J Ecol. doi: 10.1111/ aje.12009

24. Diekmann M, Scott A, Scott M, Diekmann J (2004) Capture and fitting of satellite- and radiotelemetry equipment onto Cape griffon Gyps coprotheres, African white-backed Gyps africanus and lappet-faced Torgos tracheliotos vultures in the Waterberg area, Namibia, in 2004.Vulture News 51: 34–45.

25. Bamford AJ, Diekmann M, Monadjem A, Hardy ICW (2009) Development of non-explosive based methods for mass capture of vultures. S Afr J Wildlife Res 39, 202–208.

26. Mundy PJ, Butchart D, Ledger JA, Piper SE (1992) The vultures of Africa. Acorn Books & Russel Friedman Books, Randburg & Halfway House, South Africa.

27. Fridolfsson AK, Elleghen H (1999) A simple and universal method for molecular sexing of non-rafting birds. J Avian Biol. 20: 116–121.

28. Postupalsky S (1973) Raptor reproductive success: some problems with methods, criteria, and terminology. In: Hamerstrom FN, Harrell BE, and Olenhoff RR (eds) Management of Raptors. Proceedings of the Conference on Raptor Conservation Techniques. pp 21–31. Fort Collins, Colorado.

29. White GC, Burnham KP (1999) Program MARK: survival estimation from populations of marked animals. Bird Study 46: S120–S139.

30. White GC (2008) Closed population estimation models and their extensions in Program MARK. Environ Ecol Stat 15: 89–99.

31. Burnham KP, Anderson DR (2002) Model selection and multimodel inference. A practical information-theoretic approach. Second edition. Springer-Verlag, New York, USA.

32. Burnham KP, Anderson DR, White GC, Brownie C, Pollock KH (1987) Analysis and design methods for fish survival experiments based on release-recapture. Am Fish Soc Monogr 5.

33. Leslie PH (1943) On the Use of Matrices in Certain Population Mathematics. Biometrika 33: 183–212

34. Spangerberg JA, and Jangck JR (2005) Leslie/LeKvovich Matrix Models for Age or Stage-structured Populations 1.0, BioQUEST Curriculum Consortium, Beloit College, http://bioquest.org/esteem/esteem_details.php?product_id = 210

35. Crowder LB, Crouse DT, Heppell SS, Martin TH (1994) Predicting the Impact of Turtle Excluder Devices on Loggerhead Sea Turtle Populations. Ecological Applications 4: 437–445.

36. Campbell R (1974). The Dictionary of Birds in Color. The Viking Press.

37. Reilly T, Waadill B (1965) Marabou Stork Leptoptilos crumeniferus (Lesson) breeding in Swaziland. Ostrich 36: 96.

38. Brown L, Pomeroy DE (1984) The age structure of populations of wild birds in tropical Africa, as demonstrated by plumage characteristics and marking techniques. Proc V Pan-Afr orn Congr 97–119.

39. Borkhataria RR, Frederick PS, Hyton R, Bryan Jr AL, Rodgers JA (2008) A Preliminary Model of Wood Stork Population Dynamics in the Southeastern United States. Waterbirds 31: 42–49.

40. Kanyamibwa S, Schierer A, Pradel R, Lebreton J-D (1990) Changes in adult annual survival rates in a western European population of the white stork Ciconia ciconia. Ibis 132: 27–35.

41. Schaub M, Kania W, Koppen U (2005) Variation in primary production during winter induces synchrony in survival rates in migratory white storks Ciconia ciconia. J Anim Ecol 74: 656–666.

42. Hylton RA, Frederick PC, de la Fuente TE, Spalding MG (2006) Effects of nesting health on postfledging Survival of wood storks. Condor 108: 97–106.

43. Jovani R, Tella JF. (2004) Age-related environmental sensitivity and weather mediated nesting mortality in White Storks Ciconia ciconia. Ecology Geograph 27: 611–619.

44. Kahl MP (1966) A contribution to the ecology and reproductive biology of the Marabou Stork in East Africa. J Zool, Lond 148: 289–311.

45. Clutton-Brock TH, Allon SD, Guinness FE (1983) Parental investment and sex differences in juvenile mortality in birds and mammals. Nature 313: 131–133.

46. Seber GAF, Elton R (1981) Tag loss and the Petersen mark-recapture experiment. Program MARK. Environ Ecol Stat 15: 89–99.

47. Rotella JJ, Hines JE (2005) Effects of tag loss on direct estimates of population growth rate. Ecology 86: 821–827.

48. Whyte J, Otto JPA du T, Barton B (1993) Marabou nesting in the Kruger National Park. Ostrich 64: 186.

49. Stephens PA, Sutherland WJ, Freckleton RP (1999) What is the Allee effect? Oikos 87: 185–190.

50. Carroll C, Miquelle DG (2006) Spatial viability analysis of Amur tiger Panthera tigris altaica in the Russian Far East: the role of protected areas and landscape matrix in population persistence. J Appl Ecol 43: 1056–1068.

51. Barnes K (1998) The important bird areas of southern Africa. Johannesburg: BirdLife South Africa.

52. Sarrazin F, Baggiolini C, Pinna JL, Danchin E, Clobert J (1994) High survival estimates of griffon vultures (Gyps fulvus fulvus) in a reintroduced population. Auk 111: 853–862.