

Breeding Ecology of the Red-Billed Tropicbird Phaethon aethereus Under Contrasting Environmental Conditions in the Gulf of California

Authors: Castillo-Guerrero, José Alfredo, Guevara-Medina, Miguel A.,

and Mellink, Eric

Source: Ardea, 99(1): 61-71

Published By: Netherlands Ornithologists' Union

URL: https://doi.org/10.5253/078.099.0108

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at www.bioone.org/terms-of-use.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

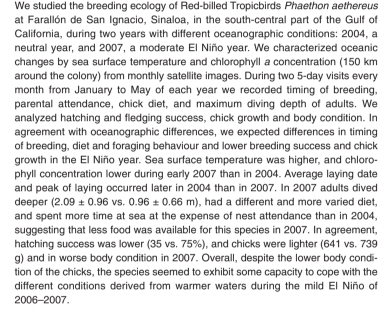
BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Breeding ecology of the Red-billed Tropicbird *Phaethon aethereus* under contrasting environmental conditions in the Gulf of California

José Alfredo Castillo-Guerrero^{1,2,*}, Miguel A. Guevara-Medina¹ & Eric Mellink¹



Castillo-Guerrero J.A., Guevara-Medina M.A. & Mellink E. 2011. Breeding ecology of the Red-billed Tropicbird *Phaethon aethereus* under contrasting environmental conditions in the Gulf of California. Ardea 99: 61–71.





Key words: breeding ecology, environmental fluctuation, El Niño Southern Oscillation, chick growth, Sinaloa, Mexico

¹Centro de Investigación Científica y de Educación Superior de Ensenada B. C.,
 Apdo. Postal 360, Ensenada Baja California, México (U.S. mailing address:
 CICESE P.O. Box 434844, San Diego, California 92143-4844);
 ²Instituto de Ciencias del Mar y Limnología, UNAM. Unidad Académica
 Mazatlán. Av. Joel Montes Camarena S/N, Apdo. Postal 811 C.P. 82040,
 Mazatlán, Sinaloa, México;

*corresponding author (acastillo@ola.icmyl.unam.mx)

In tropical areas, seasonal changes in the marine environment, like monsoons or seasonal upwelling can occur, and although seasonality in food availability is less marked than at higher latitudes, it does influence seabird breeding cycles and breeding ecology (Harrison *et al.* 1983, Morris & Chardine 1992, Le Corre 2001, Le Corre *et al.* 2003, Jaquemet *et al.* 2007, Weimerskirch 2007, Catry *et al.* 2009). Seabirds in the tropics can also face variability in physical and trophic conditions

between years due to anomalies such as the El Niño Southern Oscillation (ENSO). Seabirds have to cope with these inter-annual changes, for instance by changing their foraging areas and diets, but very bad conditions may lead to adults dying and partial or complete reproductive failures (e.g. Schreiber & Schreiber 1984, Valle *et al.* 1987, Anderson 1989, Cruz & Cruz 1990, Ramos *et al.* 2004). However, whether a given species is able to cope (to survive and reproduce), is likely to

depend on the magnitude of the variations and on intrinsic factors, such as species or population specific foraging habits. Pelagic species and those that have short foraging ranges and/or are little able to change their diet are more vulnerable to environmental changes than species with large foraging areas and a less specialized diet (Devney et al. 2009, Catry et al. 2009). Even within a given species, different sex/age groups can be affected in different ways by inter-annual variability, especially in recruitment and survival (Oro et al. 2010). Thus, long-lived tropical seabirds may have to cope with seasonal and inter-annual variation in environmental conditions.

Understanding the adaptations of seabirds to the unpredictability of their food resources is a central theme in seabird ecology (Ashmole 1971, Ramos *et al.* 2002, Weimerskirch 2007), especially in tropical oceans where productivity is lower, prey more patchily distributed, and food resources show less seasonal variation than in temperate or polar environments (Weimerskirch 2007). Knowledge of the breeding ecology of tropical seabirds under different environmental conditions is still necessary to clarify the topic. In addition, in the context of climatic change it is of interest to study the response of different species and populations to changes in their environment, to determine their vulnerability to local and regional interannual changes in the ocean (Monticelli *et al.* 2007).

In this study, we characterized the breeding biology of a tropical seabird, the Red-billed Tropicbird *Phaethon aethereus*, at a colony in the southern Gulf of California, Mexico, a region where the species' ecology is not well known, and that exhibits seasonal and inter-annual environmental fluctuations in its physical properties derived mainly from seasonal upwelling and ENSO cycles (Soto-Mardones *et al.* 1999).

Red-billed Tropicbirds are found in the tropical zones of the Pacific and Atlantic Oceans, and in the north of the Indian Ocean (Orta 1992). Despite their wide distribution, the world population of Red-billed Tropicbirds is small, estimated to be less than 8000 pairs (Lee & Walsh-McGehee 2000), including about 4900 breeding pairs in the Pacific (Spear & Ainley 2005a). In the Pacific Ocean, tropicbirds nest on rocky islands along the coast of the Americas, from the northern Gulf of California and Revillagigedo Islands to the Galápagos Islands and, possibly, Plata Island, Ecuador (Spear & Ainley 2005a,b).

Both at Ascension Island in the Atlantic Ocean and at the Galápagos Islands in the Pacific, Red-billed Tropicbirds breed throughout the year, with peaks in some months and a breeding cycle of about 11 months (Stonehouse 1962, Snow 1965, Harris 1969). They lay only one egg that is incubated for 42 days, and chicks fledge at between 12 and 15 weeks. Documented breeding success has been 32–55% at Galapagos and 51% at Ascension, but these studies were carried out under El Niño conditions (Stonehouse 1962, Snow 1965, Harris 1969). Other studies of this species were limited to an estimate of their numbers and habitat use in the Pacific Ocean (Spear & Ainley 2005a, b), and some local counts and anecdotal reports (e.g. González-Bernal *et al.* 2002, Mellink & Riojas-López 2005).

In Mexico, Red-billed Tropicbirds are considered threatened (SEMARNAT 2002). Along the Pacific coast of Mexico, colonies occur at least as far south as Guerrero (Mellink & Riojas-López 2005), and the population in the Gulf of California has been estimated at 500–1000 pairs (Everett & Anderson 1991). The colony at Farallón de San Ignacio, with up to 250 pairs, is one of the two largest known colonies in the Gulf of California (Guevara-Medina *et al.* 2008).

We studied the breeding ecology of Red-billed Tropicbirds at Farallón de San Ignacio Island, a region with seasonal upwelling in the Gulf of California, during two years with different oceanographic conditions: 2004, considered a neutral year, and 2007, a moderate El Niño year. Our objectives were to determine how feeding conditions, feeding behaviour and reproduction differed between the two years. We expected that Red-billed Tropicbirds should exhibit (1) delayed laying and a shorter breeding season, (2) change in their diet and foraging behaviour, (3) lower hatching and fledging success, and (4) lower chick growth in 2007 compared to 2004.

METHODS

Study area

Farallón de San Ignacio (25°26'N, 109°22'W) is located 36 km off the coast in northern Sinaloa (Fig. 1). The island rises 140 m above sea level. Its base encompasses about 16 ha and its top approximately 3.5–4 ha, and is nearly flat (González-Bernal et al. 2002). The island is surrounded by water 200–500 m deep immediately offshore. In addition to Red-billed Tropicbirds, Brown and Blue-footed boobies Sula leucogaster and S. nebouxii, Double-crested Cormorants Phalacrocorax auritus, and Heermann's Gulls Larus heermanni nest on the island (González-Bernal et al. 2002, Guevara-Medina et al. 2008).

The climate of the region is warm and dry, with a mean annual temperature of 25°C (10.5°C in January,

and over 36°C in July), and 300 mm of rain that falls in the summer (June–September) and early autumn (October; INEGI 2007). The area has a clear seasonal oceanographic pattern, with warm Equatorial Surface Waters during the summer (July–September), and wind-driven upwelling of Pacific Subsurface Waters during the winter (December–March), causing the lowering of sea surface temperatures by about 10°C (Soto-Mardones *et al.* 1999). During summer, the concentration of photosynthetic pigments is low, whereas in winter, when northerly winds produce local upwelling, productivity is high (Alvarez–Borrego 2002).

During the 2004 breeding season (beginning in late 2003) sea surface temperature anomalies were close to 0 and this year was considered an ENSO-neutral year. In contrast, during the 2007 breeding season, sea surface temperature anomalies in the Pacific exceeded 1°C, which represent moderate El Niño conditions. This event lasted from late 2006 to March 2007 and, afterwards, conditions returned to ENSO-neutral (http:// www.cpc.ncep.noaa.gov/products/expert assessment/ ENSO DD archive.shtml). El Niño conditions are characterized by a positive Oceanic Niño Index (ONI) greater than or equal to +0.5°C and persisting for three consecutive months. The ONI is based on sea surface temperature departures from average in the Niño 3.4 region (equatorial central Pacific), and is a principal measure for monitoring, assessing, and predicting ENSO. The maximum historical ONI value is 2.5 (in 1998), whereas it was 1.1 in 2007 (NOAA 2009).

Oceanographic conditions and timing of the breeding season

To characterize oceanographic conditions, we used monthly Sea Surface Temperature (°C) and chlorophyll a concentrations (mg/m³). We derived sea surface temperature and chlorophyll data between coordinates 25.60° N, 110.45° W and 24.7° N, 109.45° W (150 km around the colony; Fig. 1), with 4 × 4 km and 5.5 × 5.5 km resolution, respectively, from data recorded by the aqua-MODIS satellite (http://oceancolor.gsfc.nasa.gov).

We made two 5-day visits per month to the island from January to May in 2004 and 2007. We searched for nests in all accessible parts of the island. For each nest, we obtained height above sea level, depth, width of the nest entrance, and orientation (N, E, S, W, NE, SE, SW, or NW). We measured burrow characteristics because in many seabird species these affect the parents' ability to protect the eggs and chicks from predators and thermal stress. We monitored all nests found throughout the season (46 in 2004, and 55 in



Figure 1. Geographic location of the study area: Farallón de San Ignacio Island (FSI), south-central Gulf of California, Mexico. Shading represents the area from which monthly sea surface temperature and chlorophyll concentration were calculated.

2007). We marked all eggs with waterproof ink to note if they were replaced. Chicks were colour-coded individually with plastic tarsal bands. The onset of the breeding season was determined by subtracting 42 days (mean incubation time, Stonehouse 1962, Snow 1965, Harris 1969) from the date the earliest chicks hatched. If any nests failed during November and December, they were not detected. However, we visited the island in August and October 2004 and observed no breeding activity, so the likelihood that we missed asynchronously early nests was probably small.

Parental attendance, foraging behaviour and diet

We used the proportion of times that parents were present at nests (two daily checks, one in the morning and one in the afternoon) throughout incubation and during the first three weeks after hatching as an index of parental presence.

The maximum diving depth of adult tropicbirds was determined using capillary tubes (0.8 mm internal diameter, 15 cm long, internally covered with icing sugar, and sealed at one end; Burger & Wilson 1988), attached to one of the central rectrices with water-proof tape. We attached these to adults with chicks less than 3 weeks old during January and February in 2004 and

2007. Tubes were recovered an average of 38.4 ± 11.9 (SD) hrs after being deployed, when adults returned from foraging trips. Burger & Wilson (1988) tested the accuracy and applicability of this method and determined that differences between real and estimated depths averaged <3% in single immersions to any depth up to 140 m. With multiple immersions, errors were usually <10% and always <25%. The main disadvantages of the method are that only maximum diving depth is recorded and that high environmental humidity makes reading the tubes difficult. We recovered 35 capillary tubes (23 in 2004 and 12 in 2007) with clearly readable immersion marks.

To characterize diet, we examined 24 regurgitations each year obtained mainly from chicks when handled or from adults trying to feed chicks. Prey items were identified with the aid of Fischer *et al.* (1995), and those that were complete were measured (total length and weight). All mass regurgitated was given back to the chick, to reduce adverse effects on chick growth and body condition.

Breeding success and chick growth

During each year, breeding success was assessed as hatching success (eggs hatched / eggs laid) and fledgling success (chicks reaching flight / hatched eggs). We monitored the growth of 36 chicks in 2004 and 31 chicks in 2007 by daily measuring their culmen and tarsus with callipers (± 0.1 mm) and their mass with a digital balance (± 1 g). Using chicks of known age (seven in 2004 and eight in 2007), we estimated the age of those that hatched between visits (always smaller than 8-day periods).

Body condition of chicks was calculated from measurements of body mass, by correcting for body size. Body mass was regressed upon a body size index (PC1 from culmen and ulna measures, 97% of variance explained) and the residual, expressed as a proportion of the predicted value, was used as an index of body condition (Hamer & Hill 1993). Since we had multiple measurements for each chick, we calculated their average body condition.

Statistical analyses

Average laying date (days after 1 October) was compared between years using a General Linear Model (GLM) considering an identity-link and normal distribution. We compared the percent of parental presence during incubation and during post-hatching in 2004 and 2007 with a normal-identity link GLM. Year was used as a categorical factor and laying date was included as covariate. Maximum diving depths were

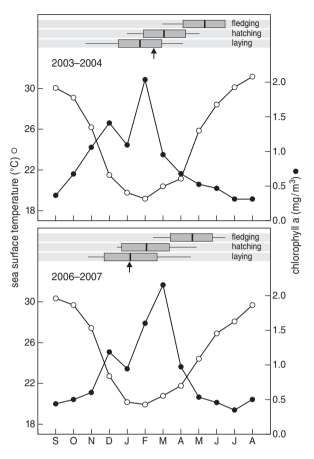


Figure 2. Timing of the breeding season, sea surface temperatures (°C) and chlorophyll a concentrations (mg/m³) during an El Niño-neutral year (2003–2004) and a moderate El Niño year (2006–2007). The three horizontal bars on top of each panel show the mean \pm SD (box) and range (lines) of the timing of breeding: egg-laying; hatching of eggs; fledging of chicks, see Table 1 for details and statistical tests). Peaks of laying are indicated by the arrows.

compared between years with the same model, but we used year and reproductive stage (incubation or brooding) as categorical factors, and date (days since 1 October) as a covariate.

To compare diet composition between the two years we applied the non-parametric analysis of similarity (ANOSIM), using PRIMER statistical software (version 5.2.9). The data on regurgitates were grouped in four categories (flying fish, other fishes, cephalopods and crustaceans) and the number of prey in each group present within each sample was converted to percentage. A similarity matrix using the Bray–Curtis measure was generated. ANOSIM tests were run on the matrices using 999 permutations to test for statistically significant differences in diet composition between samples.

In addition, we predicted the amount of prey species in each year through the relation between sampling effort and the prey species accumulation curve. To build this accumulation curve we resampled the data through (n=100) using EstimateS, Ver. 8.0 (Colwell 2006). Afterwards we adjusted the model: prey-species = $(a \times no. samples)/(1+(b \times no. samples))$ using Statistica 7.1 (StatSoft 2005) and predicted richness as a function of a and b from the equation of the curve (Jiménez-Valverde & Hortal 2003).

To compare hatching and fledging success we used a logit-link GLM due to the binomial distribution of the response variable. First, we used a model containing only year effects for both hatching and fledging success; then we added percent of parental attendance during the corresponding period (incubation, chick tending), and laying date as continuous predictors in both models. All the nest characteristics were removed because they did not significantly influence the response variables in preliminary analyses.

Culmen length at fledging (70–80 days) and maximum mass (60–70 days) were analyzed through identity-link GLM, with year (2004 or 2007) as a categorical factor, and laying date as a continuous predictor. Average body condition was analyzed in a similar way, but adding survival (survived, died) as an additional categorical factor.

RESULTS

Timing of the breeding season

Mean sea surface temperature during the 2004 and 2007 breeding season (October–May) was 22.9°C and 23.4°C respectively. At the beginning and middle of the breeding season (October–April), sea surface temperature was lower in 2004 than 2007, with a maximum difference of 1.2–1.3°C in November and December (Fig. 2) and, at the end of breeding season (May–June), warmer in 2004 than in 2007 (Fig. 2). Chlorophyll concentration exhibited the inverse pattern, with maximum values in February and March. In the first half of the breeding season (October–February), chlorophyll concentration was higher in 2004 than in 2007, but from March to April it was higher in 2007. The maximum peak of chlorophyll concentration was shifted one month between years (Fig. 2).

Red-billed Tropicbirds nested at Farallón de San Ignacio from late October through early June in 2004 and 2007 (Fig. 2). In both years, the onset of the breeding season coincided with the time at which the sea surface temperature dropped below 26–27°C (Fig. 2). The first documented egg was laid on 21 and 28 October, respectively. Average laying date and peak of laying occurred later in 2004 than in 2007 (Fig. 2, Table 1).

Table 1. Pairwise comparisons of 2004 vs. 2007 (means \pm SE, sample sizes in brackets) in the timing of the breeding season, breeder foraging behaviour and breeding parameters of Red-billed Tropicbirds at Farallón de San Ignacio, Mexico. Statistical tests: df = 1 in each case, except for Diet composition and Predicted number of prey species (see text for more details).

Parameter	2004	2007	Statistical test	
Laying date (days since 1 October)	113 ± 5.2 (49)	97 ± 5.5 (62)	$\chi^2 = 4.4, P = 0.035$	
Parental attendance during incubation (%)	$88 \pm 3.1 (36)$	$68 \pm 3.4 (40)$	$\chi^2 = 10.3, P = 0.001$	
Parental attendance post-hatching (%)	$75 \pm 5.2 (24)$	$58 \pm 5.4 (21)$	$\chi^2 = 3.1, P = 0.08$	
Maximum diving depth (m) of adults	0.96 ± 0.14 (23)	2.09 ± 0.27 (12)	$\chi^2 = 26, P < 0.001$	
Diet composition (%):	(24)	(24)	R = 0.084, P = 0.025	
Flyingfish	40 ± 8.7	37.5 ± 10.1		
Other fishes	13 ± 6.8	43.7 ± 10.1		
Squid	40 ± 8.96	14.6 ± 7.0		
Pelagic red crab	7 ± 4.4	4.2 ± 4.2		
Predicted number of prey species	11 ± 1.4 (24)	$24 \pm 1.4(24)$	$a = 1.05, b = 0.09, r^2 = 0.98$ $a = 0.96, b = 0.04, r^2 = 0.99$	
Hatching success (%)	$75 \pm 7.3 (36)$	$35 \pm 7.6 (40)$	$\chi^2 = 11.43, P < 0.001$	
Fledging success (%)	$78 \pm 6.8 (36)$	$77 \pm 7.2 (36)$	$\chi^2 = 0.016, P = 0.89$	
Culmen length at fledging (mm)	51.56± 0.37 (25)	50.51 ± 0.57 (23)	$\chi^2 = 2.08, P = 0.15$	
Maximum mass (g) attained by chicks	$724 \pm 10.5 (25)$	$600 \pm 21.8 (23)$	$\chi^2 = 22.3, P < 0.001$	
Chick body condition	0.11 ±0.028 (25)	-0.12 ± 0.020 (23)	$\chi^2 = 51.5, P < 0.001$	

Parental attendance, foraging behaviour and diet

Parental nest attendance during incubation was lower in 2007 than in 2004 (Fig. 3A, Table 1), and post-hatching parental attendance was also lower in 2007 than in 2004, but not significantly so (Fig. 3A, Table 1). Adult tropicbirds dived significantly shallower in 2004 than in 2007 (Fig. 3B, Table 1). In 2004 and 2007, adults delivered 8 and 12 different species of prey to their chicks, respectively, but the two main prey species (Sharpchin Flyingfish *Fodiator acutus rostratus* and Dart Squid *Loliolopsis diomedeae*) were the same (Appendix 1). According to the prey species / sampling effort curve (see Methods), there were 11 prey species in 2004 and 24 in 2007 (Table 1). ANOSIM tests revealed slight but significant differences in diet composition

between 2004 and 2007 (Table 1). This difference was due to an increase in the amount of fish with a corresponding decrease in the importance of squid during the 2007 breeding season (Fig. 3C).

Reproduction and chick growth

All clutches (n=101) consisted of one egg. Hatching success was significantly higher in 2004 than in 2007 (Fig. 4A, Table 1), whereas fledging success did not differ between the years (Fig. 4B, Table 1). When parental attendance was added to the model containing only year effects and laying date, both hatching success (Wald $\chi^2 = 14.2$, df = 1, P < 0.001) and fledging success (Wald $\chi^2 = 6.97$, df = 1, P = 0.008) were explained entirely by parental attendance, whereas

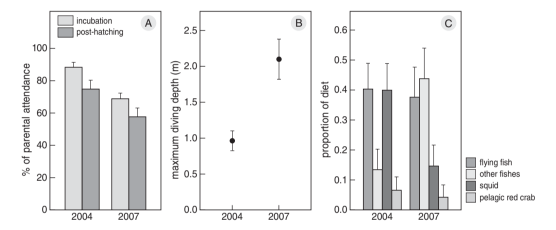


Figure 3. (A) Parental attendance during incubation (%, mean + SE) and post-hatching, (B) maximum diving depth (m, mean \pm SE), and (C) diet contribution (proportion, mean + SE) of flying fish, other fishes, squid and pelagic red crab during an El Niñoneutral year (2004) and a moderate El Niño year (2007). See Table 1 for details and statistical tests.

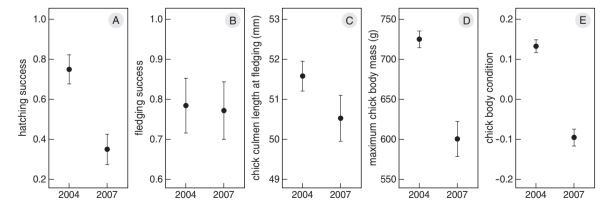


Figure 4. Breeding performance during an El niño-neutral year (2004) and a moderate El Niño year (2007, means \pm SE): (A) hatching success, (B) fledging success, (C) culmen length of chicks at fledging, (D) maximum body mass of fledged chicks, and (E) body condition of fledged chicks. See Table 1 for details and statistical tests.

year was no longer significant for hatching success (Wald $\chi^2 = 2.14$, df = 1, P = 0.14) and fledging success (Wald $\chi^2 = 1.88$, df = 1, P = 0.17).

During both years, chicks fledged at 83 days after hatching (\pm 7.8 in 2004, n = 19, and \pm 10.4 in 2007, n = 22) and did not differ in culmen length at fledging (Fig. 4C, Table 1). However, in 2004 the maximum body mass attained by chicks was higher in 2004 compared to 2007 (Fig. 4D, Table 1). Also, chicks that fledged in 2004 were in better body condition than those in 2007 (Fig. 4E, Table 1). In both years, chicks that died were in poorer condition than those that fledged (Wald χ^2 = 10.5, df = 1, P = 0.001).

DISCUSSION

We studied the breeding biology of red-billed tropicbirds by comparing two different years: a normal year (2004) and a moderate El Niño year (2007). Timing of the breeding season, feeding conditions, and parental attendance between these two years differed markedly, probably primed by differences in sea surface temperature (lower in 2004 than in 2007), and accordingly differences in patterns of primary productivity (sea water chlorophyll content was higher in the pre-breeding and early breeding season in 2004 than in 2007). Inter-annual changes in feeding conditions and parental attendance suggested better conditions in 2004 than in 2007, and accordingly breeding success, chick growth, and chick body condition were higher during 2004 than in 2007.

Timing of the breeding season

The breeding of Red-billed Tropicbirds at Farallón de San Ignacio in our study was markedly seasonal, beginning in November and continuing until early June when the last fledglings left their nests. This seasonality coincides with the annual pattern of low summer (June-August) to high winter (December-February) productivity that characterizes the east margin of the Gulf of California (Alvarez-Borrego 2002). The relatively rapid decrease in sea surface temperature (change of trend) before the onset of reproduction indicates a strong modification of the environment, which preceded the increase in marine productivity. Other tropical seabird species in the Indian Ocean, such as Sooty Tern, Red-footed Booby, Audubon's Shearwater and Redtailed Tropicbird, exhibit seasonal patterns when they inhabit areas that have seasonal, and predictable, changes in their physical conditions (mostly indicated by sea surface temperature; Le Corre 2001, Jaquemet

et al. 2007). Hermann's Gull, Elegant Tern (Velarde et al. 2004), and Blue-footed Booby (J.A. Castillo-Guerrero, unpubl. data) colonies in the Gulf of California have a seasonal breeding season related to the upwelling season, when the abundance of small pelagic fishes increases. So the Red-billed Tropicbird colony in our study contrasts with colonies at Isla Ascension in the south Atlantic (Stonehouse 1962) and Isla Daphne, Islas Galapagos (Snow 1965, Harris 1969), which lack clear oceanographic seasonality, and exhibit breeding throughout the year.

In some marine birds, the onset of egg-laying has been linked to sea surface temperature (Diamond 2003, Jaquemet et al. 2007, Tomita et al. 2009), probably because of its direct and indirect effects on food availability (Vlietstra et al. 2005, Frederiksen et al. 2006, Tomita et al. 2009). In tropical areas, surface enrichment in chlorophyll leads to a higher abundance of zooplankton and fish (Piontkovski & Williams 1995), which are largely preyed on by tropical seabirds. Redtailed Tropicbirds (P. rubricauda) nesting on Aldabra and Europa Islands in the Western Indian Ocean bred seasonally in austral summer when sea surface temperature increased above 27°C (Prys-Jones & Peet 1980, Le Corre 2001). At Europa Island, the influx of warm water appeared to increase the availability of food due to the presence of tropical fish that were unavailable at other times of the year (Le Corre et al. 2003). In contrast, Red-billed Tropicbirds at Farallón de San Ignacio in our study initiated breeding when sea surface temperature dropped below 27°C, marking the onset of seasonal upwelling. In the Eastern Pacific, Redbilled Tropicbirds use waters with shallow thermoclines and low salinity near upwelling regions (Spear & Ainley 2005a). Although the patterns indicated above are opposite, in both cases breeding is driven by seasonal oceanographic changes that increased prey availability.

Inter-year difference in mean average laying date and time of laying peak could be a result of an adaptive response by females to synchronize the energy demands of offspring production and provisioning with the period of most favourable environmental conditions (Frederiksen *et al.* 2004, Reed *et al.* 2009). Alternatively, the onset of breeding could have been constrained by the availability of resources during the pre-breeding period rather than the result of an optimal decision by the birds (Drent & Daan 1980). Although at Farallón de San Ignacio breeding was seasonal and coincidental with the time of marine upwelling, in both years it lasted for about 8 months, which allowed for an asynchrony of up to four months between the first and last breeders. So, differences in average laying date

between years appear to reflect that favourable conditions for egg-laying lasted longer. We found that in 2004 the peak of laying occurred in February, coinciding with the peak in chlorophyll concentration and the lowest sea surface temperature, whereas in 2007 the peak of laying occurred in January, two months before the peak in chlorophyll concentration. Most likely, the lower food availability in 2007 reduced the incidence of late egg-laying and thereby advanced the average egglaying date compared to 2004. Hence, tropicbirds appear to be able to "predict" the marine environmental factors affecting the timing and magnitude of food productivity, which would allow them to time their reproduction to optimize their breeding success and sometimes disfavouring egg-laying late in the season (as in 2007).

Parental attendance and foraging behaviour

The 2007 season in our study was under the influence of a weak El Niño episode (NOAA 2009). There was a sea surface temperature positive anomaly, and less chlorophyll earlier in the season than in 2004. Related to these changes, parental nest attendance, diet and diving depth of adult Red-billed Tropicbirds breeding at Farallón de San Ignacio exhibited inter-annual variability, reflecting less available food in 2007 than in 2004. When El Niño anomalies occur in the Gulf of California, there are changes in phytoplankton communities and fish populations (Kahru *et al.* 2004, Espinosa-Carreón & Valdez-Holguín 2007), and less food available to seabirds (Velarde *et al.* 2004).

At Farallón de San Ignacio, Red-billed Tropicbirds forage only in the upper three meters of the water column and prey mainly on squid and flying fish, as in previous general reports for the family (Le Corre 1997, 2001, Ramos & Pacheco 2003). However, maximum diving depth in 2007 was more than double that in 2004 and differences between both years likely reflected local and regional changes in prey availability as already shown for other species (Croxall et al. 1999, Mellink et al. 2001, Tremblay & Cherel 2003, Jaquemet et al. 2008, Catry et al. 2009). A more varied diet, deeper plunge dives and a reduction in the proportion of the main prey items in regurgitates suggests that prey were less available in 2007 and that Red-billed Tropicbirds compensated by increasing their intake of secondary prey items. In some tropical marine birds, plasticity in feeding habits reduces their vulnerability to changes and allows them to breed under varied circumstances (Jaquemet et al. 2008, Catry et al. 2009). Although limited, our data suggest that Red-billed Tropicbirds can cope with variation in food availability,

in an environment that is variable and displays little predictability at inter-annual scales.

A lower presence of parents at the nest in 2007 suggests that adults devoted more time to foraging in detriment of nest and chick protection. Flexible time-activity budgets have been supported by studies of a wide variety of seabird species, showing that as food supply declines birds spend less time at the colony and more time foraging (Montevecchi 1993, Litzow & Piatt 2003). This flexible time allocation is recognized as an important adaptation for trading off seabird breeding success and breeder survival against the variability in food supply (Litzow & Piatt 2003).

Reproduction and chick growth

Changes in foraging and parental behaviour of red billed Tropicbirds at FSI in 2007 were insufficient to compensate for the low food availability in that year, and consequently parents hatched fewer eggs, chicks fledged with a lower mass and in a worse body condition compared to 2004. Our analyses indicate that parental attendance seems the main driving factor explaining differences in reproductive success between the two years. When feeding conditions change, tropic-birds adjust their parental attendance at the expense of their success: when food is in short supply they spend more time foraging in detriment of parental attendance. This implies less protection and incubation time for eggs, and less protection and possibly food supply for chicks.

The differences in parental attendance and reproductive performance were congruent with differences in oceanographic conditions. There were differences in hatching success (early breeding season), but not in fledging success (late breeding season). So, although high chlorophyll concentration occurs at the end of the season in 2007, the chicks were in worse condition than in 2004, perhaps because the peak of chlorophyll occurred too late (i.e. before it reaches the higher trophic levels) to benefit the tropicbirds.

The response of seabirds to changes associated with El Niño events vary in agreement with the intensity of the event, the geographic area, and the seabird species. In intense events, like that of 1982–83, the response of most central and eastern Pacific seabirds was dramatic: breeding was suspended, nesting areas were deserted and some species experienced adult mortality (Schreiber & Schreiber 1984, Valle *et al.* 1987). During a moderate El Niño event (1986–1987), at Galapagos Islands some species were impacted heavily and abandoned their colonies (Blue-footed Booby and Wedgerumped Storm Petrel), others abandoned only some

colonies (Nazca Booby), and some other species did not show evident effects (Red-footed Booby and Great Frigatebird, Anderson 1989). At Farallón de San Ignacio, during the moderate El Niño event of 2007, we observed different effects on seabird species breeding. Blue-footed and Brown boobies, and Heermann's Gull had a very low breeding success, and more than 80% of the prior abandoned the colony, at the time that foraging trips were longer and most chicks died of starvation (J. A. Castillo-Guerrero, unpubl. data). Red-billed Tropicbirds, at this location, suffered the least from adverse oceanographic conditions, and they were capable of producing and raising chicks.

Coincidently to our results, Red-billed Tropicbirds in Daphne and Plaza (in the Galápagos Islands, Snow 1965, Harris 1969) and Ascension (Stonehouse 1962), lack catastrophic responses and successfully breed and raise chicks during El Niño years. We lack the conclusive data to assess why Red-billed Tropicbirds are generally less impacted by El Niño conditions compared to many other seabirds. However, based on the duration of foraging trips and diet, Red-billed Tropicbirds seem to be able to forage much farther from the colony than boobies and gulls. This, and the fact that they forage alone (Spear & Ainley 2005a), might allow them to better exploit small, isolated food patches, and not depend on large concentrations of prey (schooling fishes). In contrast, boobies, Heermann's Gull and Elegant Tern appear to depend on small pelagic schooling fish (see Mellink et al. 2001, Velarde et al. 2004). Coinciding with the former, seabirds with short foraging intervals close to the coast were more vulnerable to changes in the trophic structure in the Indian Ocean (Catry et al. 2009).

In summary, Red-billed Tropicbirds at Farallón de San Ignacio breed seasonally, coinciding with the winter upwelling period, when they can profit from a higher oceanographic productivity. A number of facts suggest that the higher sea surface temperature of 2007, in comparison to 2004, was related to lower food availability for this species at this colony. In 2007, adults spent more time foraging, dived deeper and obtained different prey species, causing a lower nest attendance and lower hatching success, compared to 2004. The smaller body mass and worse condition of chicks at fledging reflected these conditions. Overall however, despite a potential handicap imposed by El Niño conditions, the species breeds successfully compared with other seabirds in the same colony and exhibits some capacity to cope with the harsher conditions derived from warmer waters during mild El Niño conditions.

ACKNOWLEDGEMENTS

We are grateful to M. Prado E. Peñaloza and E. González for assistance during field work, to J. Moreno, Sr., and J. Moreno, Jr., for providing transportation and monitoring the weather while we were on the island, and to the González-Bernal family for logistic support. Dik Heg and two anonymous reviewers provided editorial advice and helped to improve this article. This work was funded by the Centro de Investigación Científica y de Educación Superior de Ensenada, and by the Consejo Nacional de Ciencia y Tecnología (CONACyT) through a grant to E. Mellink and graduate scholarships to M. A. Guevara and J. A. Castillo-Guerrero.

REFERENCES

- Alvarez-Borrego S. 2002. Physical oceanography. In: Case T.J., Cody M.L. & Ezcurra E. (eds) A new island biogeography of the Sea of Cortez. Oxford University Press, New York, pp. 41–69.
- Anderson D.J. 1989. Differential responses of boobies and other seabirds in the Galapagos to the 1986–87 El Niño-Southern Oscillation event. Mar. Ecol. Prog. Ser. 52: 209–216.
- Ashmole N.P. 1971. Seabird ecology and the marine environment. In: Farner D.S. & King J.R. (eds) Avian biology, vol. 1. Academic Press, New York, pp. 223–286.
- Burger A.E. & Wilson R.P. 1988. Capillary-tube depth gauges for diving animals: an assessment of their accuracy and applicability. J. Field Ornithol. 59: 345–354.
- Catry T., Ramos J.A., Jaquemet S., Faulquier L., Berlincourt M., Hauselmann A., Pinet P. & Le Corre M. 2009. Comparative foraging ecology of a tropical seabird community of the Seychelles, western Indian Ocean. Mar. Ecol. Prog. Ser. 374: 259–272.
- Colwell R.K. 2006. EstimateS, Version 7: Statistical estimation of species richness and shared species from samples (Software and User's Guide). Freeware for Windows and Mac OS. http://viceroy.eeb.uconn.edu/EstimateS
- Croxall J.P., Reid K. & Prince P.A. 1999. Diet, provisioning and productivity responses of marine predators to differences in availability of Antarctic krill. Mar. Ecol. Prog. Ser. 177: 115–131
- Cruz J.B. & Cruz F. 1990. Effect of El Niño-Southern Oscillation conditions on nestling growth rate in the Dark-rumped Petrel. Condor 92: 160–165.
- Devney C.A., Short M. & Congdon C. 2009. Sensitivity of tropical seabirds to El Niño precursors. Ecology 90: 1175–1183.
- Diamond A.W. 2003. Seabirds as indicators of changes in marine ecosystems: ecological monitoring of Machias Seal Island. Environ. Monit. Assess. 88: 153–175.
- Drent R.H. & Daan S. 1980. The prudent parent: energetic adjustments in a avian breeding. Ardea 68: 225–252
- Espinosa-Carreón T.L. & Valdez-Holguín J.E. 2007. Variabilidad interanual de clorofila en el Golfo de California. Ecología Aplicada 6: 86–92.
- Everett W.T. & Anderson D.W. 1991. Status and conservation of the breeding seabirds on offshore pacific islands of Baja California and the Gulf of California. International Council of Bird Protection Tech. Pub. 11: 115–139.

- Fischer W., Krupp F., Schneider W., Sommer C., Carpenter K.E. &. Niem V.H. 1995. Guía FAO para la identificación de especies para los fines de pesca. Pacifico centro-oriental. FAO. Roma.
- Frederiksen M., Harris M.P., Daunt F., Rothery P. & Wanless S. 2004. Scale-dependent climate signals drive breeding phenology of three seabird species. Global Change Biol. 10: 1214–1221.
- Frederiksen M., Edwards M., Richardson A.J., Halliday N.C. & Wanless S. 2006. From plankton to top predators: bottom-up control of a marine food web across four trophic levels. J. Anim. Ecol. 75: 1259–1268.
- González-Bernal M.A., Mellink E. & Fong-Mendoza J.R. 2002. Nesting birds of Farallón de San Ignacio, Sinaloa, Mexico. Western Birds 33: 254–257.
- Guevara-Medina M.A., Castillo-Guerrero J.A. & González-Bernal M.A. 2008. Presencia y abundancia de aves de la isla Farallón de San Ignacio, Sinaloa. Huitzil 9: 23–32.
- Hamer K.C. & Hill J.K. 1993. Variation and regulation of meal size and feeding frequency in Cory's Shearwater Calonectris diomedea. J. Anim. Ecol. 62: 441–450.
- Harris M.P. 1969. Factors influencing the breeding cycle of the Red-billed Tropicbird in the Galapagos Islands. Ardea 57: 148–157.
- Harrison C.S., Hida T.S. & Seki M.P. 1983. Hawaiian seabird feeding ecology. Wildlife Monogr. 85: 1–71.
- INEGI 2007. Anuario Estadístico del Estado de Sinaloa. Instituto Nacional de Estadística, Geografía e Informática, México. México D.E.
- Jaquemet S., Le Corre M. & Quartly G.D. 2007. Ocean control of the breeding regime of the Sooty Tern in the southwest Indian Ocean. Deep Sea Res. Pt I 54: 130–142.
- Jaquemet S., Potier M., Cherel Y., Kojadinovic J., Bustamante P., Richard P., Catry T., Ramos J.A. & Le Corre M. 2008. Comparative foraging ecology and ecological niche of a superabundant tropical seabird: the Sooty Tern *Sterna fusca*ta in the southwest Indian Ocean. Mar. Biol. 155: 505–520.
- Jiménez-Valverde A. & Hortal J. 2003. Las curvas de acumulación de especies y la necesidad de evaluar la calidad de los inventarios biológicos. Revista Ibérica de Aracnología 8: 151–161.
- Kahru M., Marinone S.G., Lluch-Cota S.E., Parés-Sierra A. & Mitchell B.G. 2004. Ocean-color variability in the Gulf of California: scales from days to ENSO. Deep Sea Res. Pt II 51: 139–146.
- Le Corre M. 1997. Diving depths of two tropical Pelecaniformes: the Red-tailed Tropicbird and the Red-footed Booby. Condor 99: 1004–1007.
- Le Corre M. 2001. Breeding season of seabirds at Europa Island (southern Mozambique Channel) in relation to seasonal changes in the marine environment. J. Zool. 254: 239–249.
- Le Corre M., Cherel Y., Lagarde F., Lormee H. & Jouventin P. 2003. Seasonal and inter-annual variation in the feeding ecology of a tropical oceanic seabird, the Red-tailed Tropic-bird *Phaethon rubricauda*. Mar. Ecol. Prog. Ser. 255: 289–301
- Lee D.S. & Walsh-McGehee M. 2000. Populations estimates, conservation concerns, and management of tropicbirds in the Western Atlantic. Caribb. J. Sci. 36: 267–279.
- Litzow M.A. & Piatt J.F. 2003. Variance in prey abundance influences time budgets of breeding seabirds: evidence from pigeon guillemots Cepphus columba. J. Avian Biol. 34: 54–64.

- Mellink E. & Riojas-López M.E. 2005. Breeding seabirds of Morros El Potosí, Guerrero, México. Western Birds 36: 59–63.
- Mellink E., Domínguez J. & Luévano J. 2001. Diet of Eastern Pacific Brown Boobies *Sula leucogaster brewsteri* on Isla San Jorge, north-eastern Gulf of California, and an April comparison with diets in the Middle Gulf of California. Mar. Ornithol. 29: 39–44.
- Montevecchi W.A. 1993. Birds as indicators of change in marine prey stocks. In: Furness R.W. & Greenwood J.J. D. (eds) Birds as monitors of environmental change. Chapman & Hall, London, pp. 217–266.
- Monticelli D., Ramos J.A. & Quartly G.D. 2007. Effects of annual changes in primary productivity and ocean indices on breeding performance of tropical Roseate Terns in the western Indian Ocean. Mar. Ecol. Prog. Ser. 351: 273–286.
- Morris R.D. & Chardine J.W. 1992. The breeding biology and aspects of the feeding ecology of Brown Noddies *Anous stolidus* nesting near Culebra, Puerto Rico, 1985–1989. J. Zool. 226: 65–79.
- NOAA. 2009. ENSO Cycle: Recent Evolution, Current Status and Predictions. http://www.cpc.noaa.gov/products/analysis_monitoring/lanina/enso_evolution-status-fcsts-web.pdf.
- Oro D., Torres R., Rodríguez C. & Drummond H. 2010. Climatic influence on demographic parameters of a tropical seabird varies with age and sex. Ecology 91: 1205–1214.
- Orta J. 1992. Family Phaethontidae (tropicbirds). In: del Hoyo J., Elliott A. & Sargatol J. (eds) Handbook of the birds of the world. Lynx Edicions, Barcelona, Spain. pp. 280–289.
- Piontkovski S.A. & Williams R. 1995. Multiscale variability of tropical ocean zooplankton biomass. ICES J. Mar. Sci. 52: 643–656.
- Prys-Jones R.P. & Peet C. 1980. Breeding periodicity, nesting success and nest site selection among Red-tailed Tropicbirds *Phaethon rubricauda* and *Phaethon lepturus* on Aldabra Atoll. Ibis 122: 76–81.
- Ramos J.A. & Pacheco C. 2003. Chick growth provisioning of surviving and nonsurviving White-tailed Tropicbirds (*Phaethon lepturus*). Wilson Bull. 115: 414–422.
- Ramos J.A., Maul A.M., Ayrton V., Bullock I., Hunter J., Bowler J., Castle G., Mileto R. & Pacheco C. 2002. Influence of local and large-scale weather events and timing of breeding on tropical roseate tern reproductive parameters. Mar. Ecol. Prog. Ser. 243: 271–279.
- Ramos J.A., Maul A.M., Bowler J., Monticelli D. & Pacheco C. 2004. Laying date, chick provisioning, and breeding success of Lesser Noddies on Aridae Island, Seychelles. Condor 106: 887–895.
- Reed T.E., Warzybok P., Wilson A.J., Bradley R.W., Wanless S. & Sydeman W.J. 2009. Timing is everything: flexible phenology and shifting selection in a colonial seabird. J. Anim. Ecol. 78: 376–387.
- Schreiber R.W. & Schreiber E.A. 1984. Central Pacific seabirds and the El Niño Southern Oscilation: 1882 to 1983 perspectives. Science 225: 713–716.
- SEMARNAT 2002. Norma Oficial Mexicana NOM-059-ECOL-2001, Protección ambiental Especies nativas de México de flora y fauna silvestres Categorías de riesgo y especificaciones para su inclusión, exclusión o cambio Lista de especies en riesgo. 6 de Marzo de 2002. México D.F.

Snow D.W. 1965. The breeding of the Red-billed Tropicbird in the Galapagos Islands. Condor 67: 210–214.

Soto-Mardones L., Marinone S.G. & Parés-Sierra A. 1999. Variabilidad espaciotemporal de la temperatura superficial del mar en el Golfo de California. Cienc. Mar. 25: 1–30.

Spear L.B. & Ainley D.G. 2005a. At-sea distribution and abundance of tropicbirds in the eastern Pacific. Ibis 147: 391–407.

Spear L.B. & Ainley D.G. 2005b. At-sea behaviour and habitat use by tropicbirds in the eastern Pacific. Ibis 147: 353–366.

Stonehouse B. 1962. The tropicbirds (Genus *Phaethon*) of Ascension Island. Ibis 103: 124–161.

Tomita N., Niizuma Y., Takagi M., Ito M. & Watanuki Y. 2009. Effect of interannual variations in sea-surface temperature on egg-laying parameters of black-tailed gulls (*Larus crassirostris*) at Teuri Island, Japan. Ecol. Res. 24: 157–162.

Tremblay. Y. & Cherel Y. 2003. Geographic variation in the foraging behavior, diet and chick growth of Rockhopper Penguins. Mar. Ecol. Prog. Ser. 251: 279–297.

Valle C.A., Cruz F, Cruz J.B., Merlen G. & Coulter M.C. 1987.
The impact of the 1982–1983 El Niño-Southern Oscillation on seabirds in the Galapagos Islands, Ecuador. J. Geophys. Res. C92: 14437–14444.

Velarde E., Ezcurra E., Cisneros-Mata M.A. & Lavín M.F. 2004. Seabird Ecology, El Niño anomalies, and prediction of sardine fisheries in the Gulf of California. Ecol. Appl. 14: 607–615.

Vlietstra L.S., Coyle K.O., Kachel N.B. & Hunt Jr. G.L. 2005. Tidal front affects the size of prey used by a top marine predator, the Short-tailed Sheawater (*Puffinus tenuirostris*). Fish Oceanogr. 14: 196–211.

Weimerskirch H. 2007. Are seabirds foraging for unpredictable resources? Deep Sea Res. Pt II 54: 211–223.

SAMENVATTING

De auteurs onderzochten de broedbiologie van de Keerkringvogel Phaethon aethereus op het eiland Farallón de San Ignacio in de Golf van Californië voor de kust van Mexico. De omstandigheden op zee (oppervlaktetemperatuur en concentratie van chlorofyl a) werden in een van twee jaren (2007) beïnvloed door El Niño, in een ander jaar (2004) was dat niet het geval. Van januari tot en met mei werd tweemaal per maand gedurende vijf dagen verbleven op het eiland. De registraties omvatten het tijdstip van begin van broeden, de aanwezigheid van de ouders bij de jongen, en de maximale duikdiepte door de ouders. Daarnaast werd onderzocht wat de jongen aan voedsel kregen, hoe snel ze groeiden en wat hun lichaamsconditie was. Tevens werd het uitkomstpercentage van de eieren vastgesteld, en het percentage van de jongen die vliegvlug werden. In 2007 was de temperatuur van het zeewater hoger dan in het controlejaar, en bovendien was de concentratie van chlorofyl lager. De vogels begonnen eerder met de eileg in 2007. De metingen gaven aan dat er minder voedsel beschikbaar was in 2007: de vogels doken dieper (2,09 m tegen 0,96 m in 2004), de voedselsamenstelling was gevarieerder, en ze bleven langer op zee ten koste van de tijd bij de jongen. Ook was het uitkomstpercentage lager (35% tegen 75% in 2004), en de jongen waren lichter (641 g tegen 739 g) en in slechtere conditie. (DH)

Corresponding editor: Dik Heg Received 29 May 2010; accepted 10 January 2011

Appendix 1. Frequency of occurrence of prey species and mean length and weight (\pm SE) in regurgitations of Red-billed Tropicbirds at Farallón de San Ignacio, Mexico, during the 2004 (24 regurgitates) and 2007 (24 regurgitates) breeding seasons. - = not determined.

Prey species	2004	2007	Length (mm)	Weight (g)
Cephalopods				
Dart Squid Loliolopsis diomedeae	13	4	93±8	-
Crustaceans				
Pelagic Red Crab Pleuroncodes planipes	3	1	-	-
Fishes				
Sharpchin Flyingfish Fodiator acutus	11	7	162±5	49±10
Flyingfish Hirundichthys sp.	1	1	132	31
Sailfin Flyingfish Parexocoetus brachypterus	1	0	214	-
Ornamented Flyingfish Cypselurus callopterus	0	1	-	-
Mackerel Scomber sp.	2	1	-	-
Longfin Halfbeak Hemiramphus saltator	1	4	181	-
Bigwing Halfbeak Oxyporhampus micropterus	0	2	-	-
Mackerel Scad Decapterus macarellus	0	2	-	-
Shortjaw Leatherjack Oligoplites refulgens	1	1	145	-
Pacific Thread Herring Opisthonema libertate	0	1	-	37
Mojarra Gerres sp.	0	1	132	38