Research and Management Techniques for the Conservation of Sea Turtles

Prepared by IUCN/SSC Marine Turtle Specialist Group

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In 1995 the IUCN/SSC Marine Turtle Specialist Group (MTSG) published *A Global Strategy for the Conservation of Marine Turtles* to provide a blueprint for efforts to conserve and recover declining and depleted sea turtle populations around the world. As unique components of complex ecosystems, sea turtles serve important roles in coastal and marine habitats by contributing to the health and maintenance of coral reefs, seagrass meadows, estuaries, and sandy beaches. The *Strategy* supports integrated and focused programs to prevent the extinction of these species and promotes the restoration and survival of healthy sea turtle populations that fulfill their ecological roles.

Sea turtles and humans have been linked for as long as people have settled the coasts and plied the oceans. Coastal communities have depended upon sea turtles and their eggs for protein and other products for countless generations and, in many areas, continue to do so today. However, increased commercialization of sea turtle products over the course of the 20th century has decimated many populations. Because sea turtles have complex life cycles during which individuals move among many habitats and travel across ocean basins, conservation requires a cooperative, international approach to management planning that recognizes inter-connections among habitats, sea turtle populations, and human populations, while applying the best available scientific knowledge.

To date our success in achieving both of these tasks has been minimal. Sea turtle species are recognized as “Critically Endangered,” “Endangered” or “Vulnerable” by the World Conservation Union (IUCN). Most populations are depleted as a result of unsustainable harvest for meat, shell, oil, skins, and eggs. Tens of thousands of turtles die every year after being accidentally captured in active or abandoned fishing gear. Oil spills, chemical waste, persistent plastic and other debris, high density coastal development, and an increase in ocean-based tourism have damaged or eliminated important nesting beaches and feeding areas.

To ensure the survival of sea turtles, it is important that standard and appropriate guidelines and criteria be employed by field workers in all range states. Standardized conservation and management techniques encourage the collection of comparable data and enable the sharing of results among nations and regions. This manual seeks to address the need for standard guidelines and criteria, while at the same time acknowledging a growing constituency of field workers and policy-makers seeking guidance with regard to when and why to invoke one management option over another, how to effectively implement the chosen option, and how to evaluate success.

The IUCN Marine Turtle Specialist Group believes that proper management cannot occur in the absence of supporting and high quality research, and that scientific research should focus, whenever possible, on critical conservation issues. We intend for this manual to serve a global audience involved in the protection and management of sea turtle resources. Recognizing that the most successful sea turtle protection and management programs combine traditional census techniques with computerized databases, genetic analyses and satellite-based telemetry techniques that practitioners a generation ago could only dream about, we dedicate this manual to the resource managers of the 21st century who will be facing increasingly complex resource management challenges, and for whom we hope this manual will provide both training and counsel.

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# Table of Contents

## 1. Overview

An Introduction to the Evolution, Life History, and Biology of Sea Turtles ........................................... 3  
*A. B. Meylan and P. A. Meylan*

Designing a Conservation Program ........................................................................................................... 6  
*K. L. Eckert*

Priorities for Studies of Reproduction and Nest Biology ................................................................. 9  
*J. I. Richardson*

Priorities for Research in Foraging Habitats ....................................................................................... 12  
*K. A. Bjorndal*

Community-Based Conservation ............................................................................................................. 15  
*J. G. Frazier*

## 2. Taxonomy and Species Identification

Taxonomy, External Morphology, and Species Identification ............................................................ 21  
*P. C. H. Pritchard and J.A. Mortimer*

## 3. Population and Habitat Assessment

Habitat Surveys ........................................................................................................................................ 41  
*C. E. Diez and J. A. Ottenwalder*

Population Surveys (Ground and Aerial) on Nesting Beaches ............................................................ 45  
*B. Schroeder and S. Murphy*

Population Surveys on Mass Nesting Beaches .................................................................................... 56  
*R. A. Valverde and C. E. Gates*

Studies in Foraging Habitats: Capturing and Handling Turtles .......................................................... 61  
*L. M. Ehrhart and L. H. Ogren*

Aerial Surveys in Foraging Habitats ...................................................................................................... 65  
*T. A. Henwood and S. P. Epperly*

Estimating Population Size ................................................................................................................... 67  
*T. Gerrodette and B. L. Taylor*

Population Identification ......................................................................................................................... 72  
*N. FitzSimmons, C. Moritz and B. W. Bowen*
4. Data Collection and Methods

Defining the Beginning: the Importance of Research Design ....................................................... 83
J. D. Congdon and A. E. Dunham

Data Acquisition Systems for Monitoring Sea Turtle Behavior and Physiology .............................. 88
S. A. Eckert

Databases ....................................................................................................................................... 94
R. Briseño-Dueñas and F. A. Abreu-Grobois

Factors to Consider in the Tagging of Sea Turtles ....................................................................... 101
G. H. Balazs

Techniques for Measuring Sea Turtles .......................................................................................... 110
A. B. Bolten

Nesting Periodicity and Internesting Behavior ................................................................................ 115
J. Alvarado and T. M. Murphy

Reproductive Cycles and Endocrinology ........................................................................................ 119
D. Wm. Owens

Determining Clutch Size and Hatching Success ............................................................................. 124
J. D. Miller

Determining Hatchling Sex ......................................................................................................... 130
H. Merchant Larios

Estimating Hatchling Sex Ratios .................................................................................................. 136
M. Godfrey and N. Mrosovsky

Diagnosing the Sex of Sea Turtles in Foraging Habitats ............................................................... 139
T. Wibbels

Diet Sampling and Diet Component Analysis .............................................................................. 144
G. A. Forbes

Measuring Sea Turtle Growth ..................................................................................................... 149
R. P. van Dam

Stranding and Salvage Networks .................................................................................................. 152
D. J. Shaver and W. G. Teas

Interviews and Market Surveys .................................................................................................. 156
C. Tambiah
5. Reducing Threats

Reducing Threats to Turtles ............................................................................................................... 165
M. A. G. Marcovaldi and C. A. Thomé

Reducing Threats to Eggs and Hatchlings: *In Situ* Protection ....................................................... 169
R. H. Boulon, Jr.

Reducing Threats to Eggs and Hatchlings: Hatcheries ....................................................................... 175
J. A. Mortimer

Reducing Threats to Nesting Habitat .................................................................................................. 179
B. E. Witherington

Reducing Threats to Foraging Habitats .............................................................................................. 184
J. Gibson and G. Smith

Reducing Incidental Catch in Fisheries ............................................................................................ 189
C. A. Oravetz

6. Husbandry, Veterinary Care, and Necropsy

Ranching and Captive Breeding Sea Turtles: Evaluation as a Conservation Strategy ....................... 197
J. P. Ross

Rehabilitation of Sea Turtles .............................................................................................................. 202
M. Walsh

Infectious Diseases of Marine Turtles ................................................................................................. 208
L. H. Herbst

Tissue Sampling and Necropsy Techniques ......................................................................................... 214
E. R. Jacobson

7. Legislation and Enforcement

Grassroots Stakeholders and National Legislation ............................................................................... 221
H. A. Reichart

Regional Collaboration ......................................................................................................................... 224
R. B. Trono and R. V. Salm

International Conservation Treaties .................................................................................................. 228
D. Hykle

Forensic Aspects ................................................................................................................................. 232
Determining Hatchling Sex

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Introduction
Sexual differentiation in mammals depends on the transformation of the undifferentiated gonad into a testicle. The gene that controls this initial event is SRY and is localized in the short arm of the Y chromosome (Koopman et al., 1990). In non-mammalian vertebrates, SRY related genes of the SOX family are found in both sexes and are independent of the presence of sex chromosomes (Tiersh et al., 1991). The identity of the gene or genes that control sexual differentiation in non-mammalian vertebrates is still unknown. However, as in placental mammals, morphological sexual differentiation in the embryo seems to initiate in the embryo’s gonad. It is reasonable to postulate that the factor or factors required for sexual differentiation act primarily at the level of this organ, controlling its transformation into an ovary or a testicle.

Although the physiological mechanism by which temperature or other environmental factors influence sexual differentiation is unknown, vertebrates have been divided into two groups: (1) Organisms in which environmental factors have no influence over their sexual differentiation are classified as having Genotypic Sex Determination (GSD); (2) Organisms whose sex determination is influenced by the environment, undergo an Environmental Sex Determination (ESD) (Bull, 1983). In sea turtles, determination of sex by temperature has been found in Caretta caretta (Yntema and Mrosovsky, 1980), Chelonia mydas (Miller and Limpus, 1981), Dermochelys coriacea (Rimblot et al., 1985), Lepidochelys olivacea (Morreale et al., 1982), Lepidochelys kempii (Shaver et al., 1988), and Eretmochelys imbricata (Dalrymple et al., 1985).

Identifying Sex in Hatchlings
There is a thermosensitive period (TSP) for sex determination during development placed around the second third of total time of incubation. TSP is defined as “that time span or developmental-stage span outside of which temperature manipulations do not exert any influence on sexual phenotype” (Mrosovsky and Pieau, 1991). Furthermore, in all species of marine turtles, there are no external morphological characters which may be used to determine the sex of organisms at hatching stage, and only through dissection and direct observation of the gonads is this possible.

Invasive Methods (Dissection)
Three procedures, based on morphological observations, are available: (1) direct observations of the gonads in situ; (2) clearing technique of gonads in toto; (3) histological study of the gonads.

Criteria based on (1) are concentrated on gonadal morphological details observable immediately after the viscera that cover them (e.g., intestines, liver, stomach) are removed. Gonads appear as two clear bands that extend along the length of the kidneys (mesonephros). McCoy et al. (1983) attempted to sex L. olivacea gonads based on the fact that ovaries tend to have a wrinkled surface and are larger in size than the testicles. As this criteria is questionable, van der Heiden et al. (1985) proposed method (2) which requires the dissection of the urogenital complex (gonad and kidneys) and fixation in 10% formalin. Afterward, the gonad is separated from the kidney and submerged in 100 ml of 4% formalin solution and 5 ml glycerol (a few drops of copper sulphate should be added to avoid fungal contamination). Using a dis-
section microscope, these authors sexed *L. olivacea* and *C. mydas* hatchlings. Besides the gross external morphology, from which the sexes can be clearly distinguished when the material is processed while still fresh (ovaries have a wrinkled surface and are larger than testicles), gonads in their interior also show clear differences. This is particularly true in the anterior and posterior ends (since they are narrower there) where more detailed observation is possible. According to the authors, testicles are distinguishable by a granular appearance that possibly corresponds to the presence of seminiferous tubules.

In spite of the ease with which the previous procedures can be performed to determine the sex of hatchlings, some authors have expressed serious concerns of their value, proposing that the only reliable criterion is provided by a histological study of the gonad (Mrosovsky and Benabib, 1990; Mrosovsky and Godfrey, 1995). In this case, gonads need to be fixed, dehydrated, embedded in wax or plastic from which stained sections can be obtained and observed under the microscope. An adequately equipped laboratory is required for this procedure.

For the purpose of exemplifying key features, a detailed interpretation of hatchling gonad histology, based on an embryological study of *L. olivacea* hatchling gonads (see Merchant-Larios et al., 1989), is presented below:

In males (Figure 1), the surface epithelium is flat, monolayered and frequently contains various germinal cells. The medullary cords appear separated from the surface epithelium although some remain attached to it. The medullary cords, surrounded by a basal membrane, are formed by epithelial type cells with abundant lipid droplets. Germ cells are scarce and there is no lumen within the medullary cords that would justify the name of “seminiferous tubules”, the correct term would be “seminiferous cords”. Among these formations and bordering the surface epithelium there is a basal membrane and abundant stromal tissue, formed largely by extracellular matrix, fibroblasts cells and blood vessels. Ovaries are distinguished by a conspicuous thickening of the surface epithelium (Figure 2). It appears as a columnar epithelium, one or more cells in thickness. It contains a thick basal membrane that separates the surface epithelium from the medullary region of the gonad. The medullary cords are vestigial and appear as small groups of epithelial cells surrounded by a basal membrane. Stromatic tissue is abundant in the medullary region.

In our experience it is possible to combine procedures (2) and (3) sequentially, taking advantage of the practical advantages of the first and the precision of the second. In cleared gonads, the sex can be easily identified but only if they are well differentiated and the preservation is satisfactory. However, in difficult cases of inter-sexes (alternating regions along the gonad with well developed cortex and medullary cords) or indetermination of the gonads (when both medullary cords and surface epithelium remain poorly developed), the same cleared gonads can be dehydrated in ethanol and embedded in paraffin or plastic for a further histological analysis. Preservation is excellent as can be seen in Figure 3.

Figure 1. *Lepidochelys olivacea* testicle fixed 3 days after hatching. One may clearly appreciate the seminiferous cords (Sc) formed by epithelial cells with dense cytoplasm due to numerous lipid granules. Some germ cells are situated in the cords (arrows) and others in the epithelial surface (arrowheads). Semithin section (2um) fixed with paraformaldehyde-glutaraldehyde (Karnovsky, 1965), post-fixed with OsO4 and embedded in Epon. 200X magnification.
Non-invasive Methods
(Radioimmunoanalysis)

A non-invasive method for the diagnosis of sex in recently hatched organisms has been attempted. Gross et al. (1995) reported the possibility of sexing C. caretta hatchlings using radioimmunoanalysis (RIA) of the chorioallantoic and amniotic fluids (CAFs). They found that in males the ratio of estradiol (E) to testosterone (T) concentrations is significantly lower than in females, allowing them to predict sex with acceptable precision. The same authors found a similar E:T ratio in plasma from hatchlings of the same species. In olive ridley turtles we have carried out RIA of serum prior to and post-hatching of various steroid metabolites including E and T (Merchant-Larios and Salame-Méndez, unpubl. observ.). Unfortunately, no significant differences could be found in any of the metabolites which would permit a distinction of sex in this species. It is possible that in C. caretta, the endocrine activity of the gonads is more advanced than in L. olivacea. This is suggested by the presence of Mullerian ducts at hatching in the latter species while in C. caretta they have disappeared almost totally (Yntema and Mrosovsky, 1980). Bearing in mind these significant differences in developmental timing between species, it is recommended that hatching hormonal patterns for the species under study and its correlation with sex be established before RIA is used as a method for sex identification.

Estimating Sex Ratios in Hatchlings

In order to make an estimation of the proportion of the two sexes (sex ratio) present in a natural population in the field, it is convenient to know beforehand the “pivotal” or “threshold” temperature. Once this is known, in situ nest temperatures of nests chosen from representative zones in the beach can be used to extrapolate the overall temperature range and, from these, derive the sex ratio for the rookery during the particular season nesting season.

The pivotal temperature is defined as the incubation temperature at which the resulting sex ratio in the clutch is 1:1. Experimentally, this value is obtained from incubating groups of eggs at various constant temperatures and determining the resulting proportion of male and female hatchlings. At a range of temperatures still allowing normal development (around 24-34°C), one can determine the masculinizing and feminizing temperature range (which produce 100% males or females, respectively) and estimate the pivotal temperature (50% of each sex). Precise, constant values are difficult to determine because of genetic variations of the individual specimens in each experimental group (see Mrosovsky and Pieau, 1991, for a major discussion on this point). For sea turtles species which have been studied, the pivotal temperature reported is around 30°C. Studies of C. caretta (Mrosovsky, 1988) specimens have shown variations of pivotal temperature as high as one degree centigrade, depending on nest size and genetic factors.

It has been suggested that temperature possibly counteracts a genetic control of gender. If this is true, then the pivotal temperature could be taken as the condition under which the genetic sex is expressed without external alterations (Mrosovsky and Pieau, 1991). Considering that in a nest or in a population at a beach there is a variable ratio of genotypical males to females and that response to temperature varies according to genetic sex, the pivotal temperature may vary as much as one degree centigrade (Mrosovsky, 1988). Thus, an estimation of pivotal temperature among the different turtle populations that nest in different beaches is recommended. A minimum of 5-6
nests per beach per season, following up for at least three consecutive years, would render values in close approximation of the true range of the pivotal temperature for the population being studied.

Although pivotal temperature is a reliable indicator when estimating sex ratios under natural conditions, a knowledge of “transition ranges of temperature” (TRT) is also convenient. This parameter refers to the difference in values between the low temperature producing 100% males and the high that results in 100% females (Mrosovsky and Pieau, 1991). As with pivotal temperature, TRT does not have a fixed value. No doubt there will be some variations depending on the sample size. Hence, as with pivotal temperature, estimation of standard deviation in a particular population will be optimized by measuring temperatures in as many nests as possible in different places on the beach in question and repeating the study over several years.

The equipment necessary for measuring temperature on the beach has been fully discussed by Godfrey and Mrosovsky (1994). They designed a module that memorizes maximum and minimum temperatures. Apart from the equipment being economical and resistant, it withstands burial, reducing the possibility of theft. The core sensor is a commercial memory thermometer (Radio Shack 277-302 or 630/1020) protected inside a Plexiglas box.

Converting sand temperatures into hatchling sex ratio is not as straightforward as it may seem. Pivotal temperatures of turtles are generally derived from incubation of the eggs at constant temperatures in the laboratory, unlike field incubation conditions (Bull, 1985). Microenvironmental factors, such as metabolic warming of the eggs, can cause nest temperatures to differ from sand temperatures (Mrosovsky and Yntema, 1980; Godfrey et al., 1997). This factor should be taken into account when measuring temperatures on the beach, and sensors should be positioned as close as possible to the nest or among the eggs in each nest in order to derive the sex ratio. Also, inter-clutch variation in pivotal temperatures can complicate conversions of beach temperature to sex ratio; hence, adequate sample sizes are important (see above). Nevertheless, sand temperature profiles are useful, particularly in assessing the impact of management techniques on sex ratio (see Godfrey and Mrosovsky, this volume).

**Final Considerations**

The gonads of different species of sea turtle reveal variations in the degree of differentiation at hatching. The most and least differentiated gonads are those of *C. caretta* and *D. coriacea* respectively, while an intermediate differentiation may be observed in *L. olivacea* and *C. mydas*. However, the gonads may be considered morphologically and physiologically immature in all sea turtle species. In vertebrates, differentiated ovaries contain oocytes surrounded by follicular cells and differentiated testes have seminiferous tubules and Leydig cells. In sea turtle hatchlings, as in other species, the onset of meiosis is delayed and there is no follicle formation in the ovaries. In the testes, no differentiated seminiferous tubules are present and only medullary cords, with few germ cells, are found. In most species, some germ cells remain in the surface epithelium and

**Figure 3.** *Dermochelys coriacea* ovary (A), testicle (B) and undifferentiated gonad C previously treated with the clarifying technique (see text). Glycerol was eliminated with a phosphate buffer and the samples were treated as for Figure 1 and 2 material.
no differentiated Leydig cells have been observed (Figure 1). Therefore, complete differentiation of gonads must occur sometime after hatching and when this happens still has to be established.

Genetic variations among different sea turtle populations and the varying environmental conditions of beaches located at different latitudes means that differences in pivotal temperature in turtles of the same species are expected. Therefore, estimations of the parameters for each beach is recommended and freely extrapolating from the results obtained in other beaches should be avoided.

Finally, it is important to consider the relative frequency of gonads referred to as “inter-sex”. In these samples, the medullary cords are conserved in some regions, as in the testicles, and the surface epithelium appears enlarged, in other regions, as in the ovaries. In other cases, the gonad remains “undifferentiated” and there is no clear development towards either sex. Considering the gonad’s vulnerability to temperature and its immaturity at the hatchling stage, it is not surprising to find variations in its development, probably in response to abrupt changes in temperature during the sensitive period.

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