Population Dynamics in Two Nesting Groups of the Leatherback Turtle, *Dermochelys Coriacea* and Implications for Management

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POPULATION DYNAMICS IN TWO NESTING GROUPS OF THE
LEATHERBACK TURTLE, DERMOCHELYS CORIACEA,
AND IMPLICATIONS FOR MANAGEMENT

M. TUNDI AGARDY

A DISSERTATION SUBMITTED IN PARTIAL FULFILLMENT
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OF

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ABSTRACT

The leatherback sea turtle, *Dermochelys coriacea*, is a highly migratory and pelagic endangered species. Leatherbacks are globally distributed, with pantropical nesting populations frequenting the Caribbean, Atlantic, Indian and Pacific oceans. These nesting groups appear to have cohesive and predictable occurrence, but the true demic structure of any population subgroup has never been demonstrated. Using two independent iso-electric focusing methods to analyze blood proteins, this study showed that the North Atlantic population of leatherback turtles is indeed subdivided into at least two, though probably many more, genetically distinct stocks.

That the Atlantic population of *Dermochelys*, previously assumed to be a contiguous, panmictic assemblage, is subdivided has important implications for management. First, since nesting groups appear to exhibit no gene flow, each nesting population needs to be treated as a separate management unit. Second, managers will have to look at demes objectively to determine where a concentration of effort and funds will produce the greatest recovery for the species. Lastly, demes will have to be monitored to determine if they are approaching minimum viable population size and to quantify the level of inbreeding.
Information on the structure and movement of populations is critical for predictive modeling of population growth and for attempted management of the species based on such population models. For example, a demographic model such as the Leslie matrix can demonstrate how a defined population will react to decreased mortality in any population sector. The results of theoretical modeling in this study show that for the North Atlantic leatherback turtle, as for the loggerhead (Caretta caretta) population modeled by Crouse et al. (1987), protection of large juveniles is the most critical management action that can be taken. The implications of this finding are of major importance to management, since current conservation measures for this and all other sea turtle species focuses almost exclusively on protection of eggs at the nesting beaches.
ACKNOWLEDGMENTS

Like the best major professors, Dr. C. Robert Shoop has provided inspiration, guidance, technical advice, and good friendship throughout the course of this dissertation research and writing. Unlike many such professors, however, he has managed to do so with an unfailing ability to instill his sense of humor and wonderful spirit into the work, and this has made the process utterly enjoyable. Many heartfelt thanks to you, Dr. Shoop. Special thanks are also due to Dr. Saul Saila, who also provided inspiration and help, with a modicum of patience besides. Dr. Peter August, who planted the seeds for so much of my thinking on these and other subjects, deserves special credit. Thanks also to Suzanne Ayvazian, who stood at my side at the best and the worst of times, and made graduate school an experience that I will always remember with fondness and a smile.
This dissertation follows manuscript format and is presented as three separate papers. The first paper describes the genetic variability in two North Atlantic populations of the leatherback turtle as derived from electrophoretic studies, and was written for submission to the herpetological journal *Copeia*. The second paper is a short description of a blood sampling technique and will be submitted as a note to *Copeia*. The third and final paper describes more generally the population dynamics of these two stocks and the management implications of the genetic and demographic work, and was written for submission to the journal *Conservation Biology* or *Biological Conservation*. 

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PROBLEM STATEMENT AND INTRODUCTION

TO DISSERTATION
The leatherback turtle, *Dermochelys coriacea*, is one of the largest and least studied reptilian species. It shares many ecological characteristics with the other seven species of sea turtles found worldwide, including numerous adaptations for life at sea in which the only normal contact with land is during nesting or at hatching. The land-based activities of sea turtles are relatively well known, but we know little about their lives at sea. The gap in our knowledge includes some very basic life history data such as life span and intrinsic rate of increase in any population, information that is sorely needed for management. Since the leatherback turtle has endangered species status and risks imminent extinction, these basic questions must be answered.

Taxonomically, the leatherback is the monotypic member of the family Dermochelyidae, while the other seven living sea turtle species are in the family Cheloniidae. Significant differences between the leatherback and the other species include large adult size and an ability to regulate body temperature (Greer et al., 1973). *Dermochelys coriacea* adults undergo large-scale migrations with home ranges that commonly exceed 5000 kilometers (Pritchard, 1976; Lazell, 1980). To withstand the severe environmental conditions of migrations from tropical breeding sites to temperate and sub-arctic feeding grounds and diving to great depths,
leatherbacks have evolved a semi-homeothermic capability. Individuals are able to maintain body temperatures up to 18°C above the ambient water (Standora et al., 1984) with an efficient countercurrent heat exchanger in the circulatory system (Friar et al., 1972).

*Dermochelys* has such unusual anatomical features that some systematists argue for a taxonomic separation at the subordinal level from the other sea turtles (Smith and Smith, 1979). The most obvious morphological characteristic unique to leatherbacks is the seemingly "soft" shell. The carapace, although present, is reduced to a relatively thin mosaic-like shell under the tough but flexible skin. This adaptation may have evolved in response to two ecological features: (1) the enormous size of adults, sometimes exceeding 900 kilograms (Eckert and Luginbuhl, 1988), which may discourage the predation that would make a shell necessary, and (2) deep diving behavior, exceeding 470 meters (Eckert et al., 1986), possibly enhanced by tissues which can tolerate compression under great pressures. Indeed, Rhodin (1985) has shown that leatherback bone is different from that of other reptiles, being much more flexible and compressible than the bone of other sea turtles.

Osteological studies of leatherbacks have also led to some surprising and counter-intuitive theories about their rate of growth. The large size and high fecundity of *Dermochelys* may suggest a demographic pattern of slow maturation and long life span that characterize many other chelonid species. However, Rhodin (1985) postulated that leatherbacks grow very rapidly, attaining
reproductive maturity in two to three years. If Rhodin's estimate is correct, it has serious implications for projections of population growth and, correspondingly, for management. How such a rapid growth rate, an over 60,000-fold increase in weight in only 730 days, could be sustained by an animal adapted to a diet of soft-bodied coelenterates comprised mostly of water, is unknown. The metabolic requirements of such rapid growth are probably such that hatchling and juvenile leatherbacks would have to eat foods with fairly high protein content (Bels et al., 1989). Despite the fact that other sea turtle species are opportunistic feeders when young (Carr, 1987), they exhibit much slower growth than postulated for leatherbacks by Rhodin.

Knowledge of individual growth rate and population replacement rate are required before recovery of this endangered species can be comprehensively undertaken. Even without concrete information about these parameters, assumptions about growth rate, fecundity, and survivorship can be used to formulate theoretical models of population dynamics that can be verified or disproved through the accumulation of survivorship data. However, such models are not realistic if the population to be modelled has not been clearly defined.

I argue that the critical information necessary to promote the recovery of this species is the definition of the population to be recovered. Leather-backs are not equally endangered throughout the world, as some "populations" (e.g. the French Guiana nesting group) are quite large and apparently suffer little human-induced mortality. Thus, how leatherbacks are organized into demes (which correspond to
management units) and which demes are in gravest danger of extinction is of critical importance.

Administrators or natural resource managers responsible for the recovery of leatherback populations ought to be concerned with the efficiency of their efforts, given that time and funds are limited. Ideally, the scientific program should aim to provide the following:

1. Identification of the management unit. This identification must be comprehensive to include all age classes and both sexes within the deme.
2. Determination of population sizes and trends for each management unit. Tagging studies, growth curve generation, and other research should aim to establish the population replacement rate ($R$).
3. Identification of the most sensitive stocks and demographic groups to target for management. Since resources for conservation are limited, the most effective and fastest results will come from this sort of allocation.

The manuscripts included in this dissertation describe techniques and results that elucidate stock structure, genetic heterozygosity, and theoretical population dynamics in Atlantic leatherback turtles and discuss why such demographic information is crucial for efficient and timely management. So little is known about the biology of the leatherback that every bit of information gained is potentially important to conservation. Owing to logistical considerations, researchers have spent an inordinate proportion of time studying a very narrow aspect of this species' ecology, namely nesting. Much more work is needed on stock delineation, population
dynamics, movement patterns, and resource requirements at sea, to further elaborate the demography of this enigmatic endangered species.
LITERATURE CITED


MANUSCRIPT #1:

GENETIC VARIABILITY IN TWO SEASONALLY ALLOPATRIC POPULATIONS
OF THE LEATHERBACK TURTLE, *DERMOCHELYS CORIACEA*
GENETIC VARIABILITY IN TWO SEASONALLY ALLOPATRIC POPULATIONS
OF THE LEATHERBACK SEA TURTLE, DERMOCHELYS CORIACEA

ABSTRACT

Two-dimensional polyacrylamide gel electrophoresis was used to analyze blood samples from adult female leatherback sea turtles (Dermochelys coriacea) to elucidate the stock structure and genetic composition of two North Atlantic nesting populations. The isozyme analyses were used to estimate both overall levels of heterozygosity in the North Atlantic population and extent to which the two sampled populations could be identified as genetically distinct. Blood samples were collected from nesting female leatherback turtles in St. Croix, U.S. Virgin Islands, and Parismina, Costa Rica, two widely separated sites representing extremes of the Caribbean nesting range for D. coriacea. Isoelectric focusing was utilized in both specific isozyme identification and total protein densitometry analysis to determine whether the two populations sampled represented two true demes.

Both the non-parametric tests performed on the binary data, indicating presence or absence of a particular allozyme, and multivariate discriminant analyses of the total protein densitometry data established a clear genetic separation of the stocks. In addition, the genetic baseline information showed how animals stranded or trapped in temperate areas, whose stock identities could not be otherwise determined, could be matched to the most probable nesting population of origin, using a maximum likelihood estimator.
and a nearest neighbor classification. Electrophoretic analysis of blood proteins was thus used in two independent ways to elaborate not only stock structure and demic integrity, but also as a tool for determining the stock affinity of untagged individuals found far from their nesting grounds.
The leatherback turtle (Dermochelys coriacea), one of five sea turtle species found in the northern Atlantic and one of eight species in existence, is both federally and internationally listed as endangered (Groombridge, 1982). As such, its population numbers are considered low enough for the species to be in danger of extinction (Meylan et al., 1985; Ross 1982), and in some areas, high levels of egg harvesting and habitat loss preclude any immediate hope of recovery (Bustard, 1972). The entire worldwide population of mature females, the only segment of the population easily censused because only breeding females come ashore and because they appear at the same place in regular intervals, has been roughly estimated at only 115,000 individuals (Pritchard, 1982).

The leatherback is the largest of the sea turtles and the largest of extant marine reptiles, commonly exceeding 600 kilograms in weight (Mrosovsky, 1987). A recent stranding of an adult turtle in Wales, Great Britain, weighed a record 916 kilograms (Eckert and Luginbuhl, 1988). The leatherback is the most widely distributed marine reptile, with individual home ranges exceeding 5000 kilometers (Pritchard, 1976). It thus frequents both tropical areas and temperate to sub-arctic areas (Brongersma, 1972; Shoop et al., 1981).

Despite its large size and impressive movements, the life history and behavioral ecology of Dermochelys remain poorly understood. The highly migratory behavior and pelagic habits of the leatherback have contributed to the lack of information about the species. In
addition, *Dermochelys* is sensitive to disturbance, elusive to observe, and with one documented exception (Bels et al., 1989) virtually impossible to raise in captivity (P. Lutz, pers. comm.). These factors, coupled with the relative rarity of the animal, explain some of the paucity of existing biological data.

Our knowledge of the leatherback sea turtle is least solid in those aspects where information is most needed. While certain aspects of the reproductive biology of the circumtropical leatherback nesting populations have been well documented as annual female fecundity and egg to hatchling survivorship (e.g. Bacon et al., 1984; Balasingham, 1967, Carr and Ogren, 1959; Eckert et al., 1985; Fretey, 1980; Limpus and McLachlan, 1984; Meylan et al., 1985; Pritchard, 1971, 1976; Ross, 1982 and Schulz, 1982), virtually nothing is known about the population dynamics and ecology of the ocean inhabiting segments of the worldwide population. One important and as yet unanswered question concerns whether male leatherbacks undergo the extensive seasonal migrations that lead breeding females from temperate feeding grounds to tropical nesting areas. Since the movements of males have not been fully documented and since positive evidence of mating in the proximity of nesting beaches is lacking (Eckert and Eckert, 1988), whether geographically separated nesting groups represent genetically distinct demes is not known.

Leatherbacks inhabiting the North Atlantic Ocean utilize tropical nesting beaches throughout eastern Central America, the Antilles, and the northern shores of the South American continent (Figure 1). The largest concentrations of nesting leatherbacks in
Figure 1. Major nesting beaches used by *Dermochelys coriacea* in the Caribbean region.
this region occur in Costa Rica, French Guiana, and Guyana (Pritchard, 1976), although smaller populations return seasonally to use beaches in other parts of the Caribbean such as Puerto Rico, the U.S. Virgin Islands, and Dominican Republic (Meylan et al., 1985). The latter habitats are increasingly under risk from encroachment by beachfront resort developments, so the survival of these already small populations groups may be particularly threatened (Baker, 1980).

Questions concerning stock structure and population dynamics of Dermochelys previously have been ignored or answered in general and hypothetical terms (Brongersma, 1972; Carr, 1952;), save the recent hypothesis on Virgin Island stock structure that emerged from data on epifaunal encrustation patterns evident on nesting females (Eckert and Eckert, 1988).

The Atlantic leatherback turtle population, for instance, was assumed to constitute a largely panmictic assemblage, with open gene flow among nesting groups (Lazell, 1980). The lack of evidence of mating near nesting habitats seemed to point to some central breeding area away from the nesting beaches (J. Lazell, pers. comm.), perhaps in the temperate or subarctic feeding grounds. In this hypothesized scenario, leatherbacks inhabiting the Atlantic Ocean would be considered as constituting a single population, with only seasonal population subdivision occurring when the breeding females move to their widely separated nesting beaches to deposit eggs.

The question of stock cohesion and degree of gene flow between and within regional sub-groups of the leatherback population is important for management of the species. Reduced genetic variability
may hinder the species' ability to adapt to rapidly changing environments (Levins, 1968). The tentative link between low levels of heterozygosity and reduced fitness has been argued for many species of vertebrates (see review by Allendorf and Leary, 1986), although the mechanisms depressing fitness are rarely elucidated. However, restricted population size and genetic exchange with other demes may lead to inbreeding and a corresponding increase in expression of deleterious mutations, as has been shown in studies of the immune responses of endangered populations of cheetahs (O'Brien et al., 1986). Inbred vertebrates may have lowered resistance to stressful conditions (Parsons, 1971), and can exhibit significant behavioral changes that are considered "abnormal" (Lynch, 1971). Professional breeders of rare species have long recognized the deleterious nature of reduced genetic variability and actively promote increased heterozygosity via outcrossing (Kleiman, 1980). Frankel and Soule (1981) stated that inbreeding always reduces fitness in animals, and claimed that a 10% increase in the coefficient of inbreeding will be expected to produce a 5-10% decline in reproduction.

Despite some controversy about what levels of heterozygosity are acceptable in managed populations (see Templeton, 1986 for an extended discussion), genetic analysis remains a necessary prerequisite to endangered species management. This point has been eloquently stated in the case of endangered fish management (Meffe, 1986), but has yet to be established as critical in the management of other marine vertebrates, although marine mammalogists are now
beginning to look at the problem (P. Tyack, pers. comm.). An analysis of genetic variability exhibited by North Atlantic leatherback populations is used here as a tool to define stock structure and the corresponding units of management, to elucidate breeding behavioral patterns, and to focus questions concerning leatherback biology.

That stock identification is important for sea turtle management cannot be overstated. Not only is it important to identify unit stocks, defined by some level of geographical separation in time and/or space that occur with some corollary biological discreteness, it is also important to determine what factors contribute to stock delineation. Description of these features of managed populations is important in understanding the true limits of population interactions. Clearly, any management measure instituted to protect or rehabilitate a population will have drastically different effects if applied to small segments of a population, as opposed to the comprehensive population as a whole (Brown et al., 1987).

Electrophoretic techniques remain one of the most valuable tools for detecting intraspecific differences between members of a population (Hartl, 1980), and are in many ways more cost-effective than higher resolution techniques such as DNA sequencing and mitochondrial DNA analysis. Morphometric and meristic data are important parameters for detecting such differences, but these characters are more frequently influenced by external factors and may thus distort true unit stock delineations (Saila, 1987). Furthermore, comparisons between quantitative genetic and morphometric data may be invalid due to uneven statistical
assumptions (Lewontin, 1983). However, a combination of genetic data inferred from electrophoretic analysis and direct observation (for example as provided by tagging studies) can provide highly rigorous estimates of demic structure (Ihssen et al., 1981).

Fisheries biologists have generated huge amounts of quantitative data from biochemical samples to delimit population structure in previously uncharacterized fish stocks. For instance, stocks of highly migratory fish such as salmon, herring, and halibut have been extensively defined using electrophoretic methods (Utter et al., 1987; Grant and Utter, 1984; and Grant et al., 1984). With respect to similarly highly migratory sea turtles, some attempt has been made to quantify genetic variability in loggerhead and green sea turtles (Caretta caretta, Chelonia mydas, respectively) using starch gel electrophoresis (Smith et al., 1977), but as the authors were interested primarily in the average rate of genetic heterozygosity, genetic delineation of demes was not attempted. Bowen et al. (1989) examined the mitochondrial DNA (mtDNA) of nesting female green turtles and found genetic cohesiveness of nesting populations, however since mtDNA is maternal in origin these results do not describe the genetic composition of the population as a whole.

My study is the first to describe the population genetic structure and degree of inbreeding exhibited by leatherback turtle populations. It targets two geographically distinct groups of nesting female turtles found at the longitudinal extremes of the wider Caribbean nesting region. Blood samples taken at each site are compared using several different methods of biochemical and
statistical analyses; the results are used to give credence to the hypothesis that the Atlantic population of *Dermochelys coriacea* is not a panmictic assemblage with open gene flow among nesting groups but is instead divided into independent demes.
MATERIALS AND METHODS

Blood samples were collected from adult female leatherback turtles during their nocturnal emergences on the beach. To minimize disturbance to the nesting sea turtles, blood samples were collected immediately after the females finished egg-laying and before the active phase of nest covering began. During this time the animals are relatively still and seem less responsive to external stimuli.

Blood was drawn using a 5 cm 13-gauge spinal tap needle inserted into the paravertebral sinus of the neck (Agardy, 1989). This technique is generally reliable and minimizes trauma to the animal. For the technique to be successful the needle must be inserted close to the vertebral column (Figure 2) and kept vertical during insertion and blood withdrawal. Because of the extremely viscous nature of Dermochelys blood, both the needle and internal syringe barrel were coated in heparin, and collected samples were immediately placed in vacutainers containing EDTA as an anticoagulant. Blood samples were placed on ice following collection and were frozen at -5°C within 3-4 hours after collection.

Sampling at the beaches near Parismina, Costa Rica, took place during the period from 10-20 April 1986 and at Sandy Point, St. Croix, U.S. Virgin Islands, from 28 April to 7 May 1986 (Figure 3), under U.S. Endangered Species Permit # 703758 and CITES Permit # 698138. These two sites were selected because they represent geographical extremes of Dermochelys nesting in the Caribbean area, and because research expeditions to these locations were logistically convenient. The number of adult leatherback turtles using these
Figure 2. Needle insertion point for blood sampling in adult leatherback turtles.
Figure 3. Sampling sites in Costa Rica and the U.S. Virgin Islands.
beaches during the peak of nesting (approximately 200 on the beach in immediate proximity to Parismina (Hirth and Ogren, 1987) and 30 on the Sandy Point beach (Agardy, 1981) ensured that the small sample sizes needed for the analysis could be obtained in a short period. The sampling dates were planned in close proximity to decrease the probability of sampling the same female in two different locations. In fact, samples from the Virgin Islands group were taken from tagged and thus individually identified animals, so duplication could not have occurred.

Fifteen blood samples were collected at Parismina and twelve at Sandy Point, from twelve individual female turtles at each locality. When possible, 8-10 ml of blood were collected, however 2-3 ml proved sufficient for electrophoretic separation and staining. Frozen blood samples were shipped to the University of Rhode Island for storage and analyses.

Several blood samples were collected from stranded leatherback turtle adults in Rhode Island and Massachusetts. Only two of these samples were fresh enough to be usable: one 5 ml sample obtained from a live female captured in a pound net off Newport on 5 August 1986, the other obtained by Dr. Gregory Early of the New England Aquarium from a dead female adult found on Horseneck Beach, Massachusetts, in July 1986. The samples from the stranded individuals were frozen and stored with the other samples until analysis.

The entire collection of samples from Costa Rica and the Virgin Islands may have been subjected to a partial thaw in November 1986 when the electrical power at the Department of Zoology, University of
Rhode Island, was lost for a 36-hour period. Work done by the National Marine Fisheries Service (S. Galloway, pers. comm.) had suggested that the most stable of sea turtle blood proteins can resist denaturing, even after successive freeze/thaw episodes, although inherently unstable proteins may be prone to configurational change and thus operational dysfunction. Indeed, subsequent IEF analyses and staining of the samples showed that the relevant proteins had not denatured, since the reactions went to completion (see Appendix A for enzymatic reactions). These reactions, utilizing an electron transfer method to produce stains on specific IEF bands, require that the targeted enzyme be intact and fully functional for staining to be visible.

Prior to electrophoretic analysis, the blood was thawed at room temperature and centrifuged at approximately 5000 rpm for 20 minutes. The supernatants were then diluted with distilled water to a 1:1 concentration. The prepared samples were micropipetted in 15 μl quantities onto LKB brand paper wicks, which were then applied to polyacrylamide electrophoretic gels for separation.

The blood proteins were analyzed using an iso-electric focusing apparatus (MRS-150 Electrofocusing tank and E-C-500 constant power supply coupled with a VWR constant temperature refrigerated bath). All the samples were run together on polyacrylamide gels, thereby eliminating possible variability in running conditions between samples. Premade polyacrylamide gels (PAG plates) with a pH range of 3.5-9.5 (LKB Manufacturers) were used throughout the study.

Each of the ten gels containing 26 sea turtle blood samples was run for 2.5 hours at a maximum voltage of 1.5 kilovolts, a maximum
power of 25 watts, and a maximum current of 70 milliamps. A fresh sample of human blood was run in Lane 1 of each gel because the separation of proteins was more clearly visible in an unfrozen sample and running time could be more accurately gauged using the same human blood as a reference point each time.

Specific enzyme staining complexes were used to detect the presence of allozymes produced at distinct alleles. The specific stains were of two general classes: chemical detection stains, which employ chemical reagents to produce colored compounds at the sites of enzyme activity, and electron transfer dyes reduced by electron donors to produce a stain. The enzyme complexes assayed included lactose dehydrogenase (LDH), hexokinase (HK), peptidases B and D (PEP), glyceraldehyde-phosphate dehydrogenase (GAPDH), phosphoglucomutase (PGM), phosphogluconate dehydrogenase (PGD), and esterases (EST). In addition, detection of red cell acid phosphatase was attempted without success. Specific staining reactions are diagrammed in Appendix A.

The protocol for detecting these enzymes was devised specifically for Caretta caretta and Chelonia mydas sea turtle blood by S. Galloway and T. Inabnett (pers. comm.) of the Charleston Laboratory of the National Marine Fisheries Service and were generously provided by them for this study. Running times were extended by 10-20 minutes per assay and other slight modifications in buffer solution composition were made for D. coriacea application. Since these protocols have not yet been published, the buffer and stain recipes are noted briefly below:
**LDH** : Incubate gel at 37°C for 30-45 minutes in .6 g lactic acid, .04g B-NAD, and 50 ml .05M tris/glycine at pH 8.4. Pour off buffer and add to it .02 g NBT and .005 g PMS, pour over gel and soak until purple bands are visible.

**HK** : Mix .05 g MTT and .01 g PMS into the following buffer:
3 ml .1 M tris/HCl at pH 7.5, .18 g glucose B-D+, .04 g ATP (disodium salt), 0.16g NADP, 80 µl G-6-PDH, and .0041 g MgCl2 • 6H2O. (This is a dark reaction, therefore stains should be added to the buffer in darkness). Pipet this solution onto the gel and roll with a glass rod, then incubate at 37°C in complete darkness for 4 hours.

**PEP** : Pipette and roll onto the gel the following: 2 ml .02 M NaPO at pH 7.5, .04 g peptide, .02 g peroxidase, .5 ml .1 M MnCl2, 200 µl L-amino acid oxidase, and .028 g 3-amino-9-ethyl carbazole.
Incubate at 37°C for 2.5 hours.

**GAPDH** : Incubate gel at 37°C in 50 ml .05 M tris/HCl at pH 7.5, 40 ul D-glyceraldehyde-3-phosphate (free acid), .06 g NAD, .05 g arsenic acid, .05 g pyruvic acid, .0075 g MTT and .0025 g PMS for 1 hour.

**PGM** : Incubate gel in .48 g glucose-1-phosphate, .08 g MgCl2, .008 g NADP, .028 ml G6PDH (140 µ/ml), and 50 ml .5 M tris/HCl at pH 7.1 for one hour. Pour off and add .02 g NBT and .004 g PMS.

**PGD** : Mix 50 ml .5 M tris/HCl at pH 8.0, .1 g 6-phosphogluconate (disodium), 2.5 ml .2 M MgCl2, and 50 mg NADP. Add .075 g MTT and .015 g PMS; incubate gel at 37°C for two hours.

**EST** : Add to 50 ml .1 M tris/HCl (pH 7.0) the following: .02 g α-naphthyl acetate, .02 g β-naphthyl acetate, .02 g α-naphthyl propionate, 432 ul α-naphthyl butyrate mixed with acetone. Add
.02 g α-naphthyl phosphate, then .07 g Fast Blue BB. Incubate, without shaking or bumping staining tray, for 30 minutes.

Each stained gel was scored for presence or absence of allelic products and was then preserved and photographed (Figure 4). The binary data, indicating the presence of proteins coded by each hypothesized locus, were used to measure overall genetic variability in the populations sampled as well as intra-nesting group differences. A preliminary chi-square analysis was performed on the pooled binary values for each nesting group to determine the extent to which data differed from the null hypothesis expectation that the two populations would exhibit similar values for genetically-encoded allozyme composition. A non-parametric analysis of the binary data was then used to determine the extent to which samples could be classified into either of the two nesting populations. This analysis, using a nearest neighbor classification technique described by James (1985), assigns each sample to a group based on a type of reverse histogram method. The BASIC computer program outlined by James (1985) was also used to assign unknown samples to either of the two groups. Using another technique, the mean allozyme pattern of each population was also used as the basis for maximum likelihood estimation (MLE) and assignment of unknown samples (Millar, 1987).

In addition to the two specific enzyme detection methods indicated above, two additional gels were treated with a general protein stain employing Coomassie Brilliant Blue R-250 (Fabrizio, 1986) to detect all proteins contained in the samples. The additional gels (Figure 5) were then scanned with an EC-910
Figure 4. Example of specific enzyme stained isoelectric-focusing (IEF) gel.
Figure 5. Example of total protein stained isoelectric-focusing (IEF) gel.
integrated transmission densitometer. From the 30 to 34 proteins detected by this method, eight reference measurements of the most prominent peaks were chosen for comparison. The concentrations of these eight proteins, given as a proportion of the total integrated area of the protein profile, were then used to perform a standard discriminate analysis (Fabrizio, 1986).

The data obtained from the densitometer readings were first calculated as percentage of total area, and the fractions were arcsine transformed to mathematically accentuate the existing differences between values (Snedecor and Cochran, 1980). The transformed values for the eight selected proteins in the Virgin Islands and Costa Rica nesting population samples were then used in a stepwise discriminate analysis, which indicates discreteness of the two groups and also highlights those measurements most useful in discriminating between groups.
RESULTS

Samples were scored according to presence or absence of each of the seven most prominent metabolic allozymes commonly detected through electron transfer dye staining methods. This scoring was binary, the value one representing presence of the common allozyme and zero representing its absence (Table 1). These data were used to estimate overall levels of genetic heterozygosity in samples of the North Atlantic leatherback turtle population and to delineate stocks according to genetic differences.

All seven consistently detectable metabolic enzymes investigated: hexokinase (HK), lactate dehydrogenase (LDH), 6-phosphogluconate dehydrogenase (PGD), peptide (PEP), esterase a (EST), phosphoglucomutase (PGM), and glyceraldehyde phosphate dehydrogenase (GAPDH), were polymorphic in Dermochelys coriacea. Average heterozygosity across all seven enzyme systems of all samples (N=24) was 0.33 (Table 2). Although both Costa Rica and Virgin Islands subgroups had similar levels of heterozygosity, the samples from each population were heterozygous at different loci.

A chi-square analysis was performed on the binary data by pooling the samples in each subgroup and comparing mean values of the number of times the common allozyme for each enzyme system was present. This group-group comparison showed a clear distinction between stocks. The null hypothesis, that the seven polymorphic proteins occur independently of the population subgroup, was rejected at the 95% confidence level ($X^2_{0.05, 6}=15.32 > 12.59$).
Table 1. Binary scored data on presence or absence of the most prominent allozymes (0=absence; 1=presence) in seven enzyme systems: lactate dehydrogenase (LDH), phosphoglucomutase (PGM), 6-phosphogluconate dehydrogenase (PGD), peptide (PEP), esterase A (EST), hexokinase (HK), and glyceraldehyde phosphate dehydrogenase (GAPDH) from electrophoretically separated and stained blood samples taken from adult leatherback turtles in three locations.

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Table 2. Estimated heterozygosity values (%) in two nesting populations of *Dermochelys coriacea* as determined by electrophoretic separation of seven polymorphic proteins and specific staining. For full names of proteins see Table 1.

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<th>Loci Represented</th>
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Because of small sample size and no assumption of a normal distribution of the binary values, a non-parametric nearest neighbor classification was used to arrange samples into the demic groups based on allozyme patterns. The test assigned samples into the two groups with a Bayes error of 22.73 % (Table 3). The Bayes error value suggests the maximum rate of misclassification using these parameters for discrimination is less than one in four.

The total protein densitometry analysis, in which the entire complement of proteins present in the sample were stained and read by densitometer, resulted in protein profiles having 28 to 48 detectable peaks (Figure 5). Some of these peaks were artifacts caused by unusually wide or heterogeneous IEF bands. Of the 28 consistently apparent peaks, eight of the most prominent were selected as representative bands for analysis to make the data set manageable (Figure 5). The eight variables, representing the integrated areas of the largest peaks, were then arc-sin transformed (Table 4) and were used in a stepwise linear discriminant analysis (Saila and Martin, 1987). This analysis, like the nearest neighbor classification, assigns samples to groups according to the eight variables; results showed a clear genetic separation of stocks.

The total protein densitometry data were assumed to have a normal distribution for the multivariate discriminant analysis. Thus, in creating the group one and group two covariance matrices, the computer analysis treated them as statistically identical. Using cross-validation, the program chose only the most highly discriminatory variables and then performed the grouping using this function.
Figure 6. Example of densitometry readings for a blood sample taken from an adult leatherback turtle in Costa Rica.
Table 3. Nearest neighbor (NN) classification (James, 1985) of electrophoretically separated and specifically stained leatherback blood proteins, using binary data from Table 1 with 22 cases using 7 variables. Asterisks denote misclassified cases.

Group I: Costa Rica Samples

NN to 1 from Group 1 is case 15 from Group 2 *
NN to 2 from Group 1 is case 9 from Group 1
NN to 3 from Group 1 is case 8 from Group 1
NN to 4 from Group 1 is case 5 from Group 1
NN to 5 from Group 1 is case 4 from Group 1
NN to 6 from Group 1 is case 1 from Group 1
NN to 7 from Group 1 is case 16 from Group 2 *
NN to 8 from Group 1 is case 3 from Group 1
NN to 9 from Group 1 is case 2 from Group 1
NN to 10 from Group 1 is case 11 from Group 1
NN to 11 from Group 1 is case 12 from Group 1
NN to 12 from Group 1 is case 11 from Group 1

Group II: Virgin Island Samples

NN to 15 from Group 2 is case 1 from Group 1 *
NN to 16 from Group 2 is case 7 from Group 1 *
NN to 17 from Group 2 is case 21 from Group 2
NN to 18 from Group 2 is case 19 from Group 2
NN to 19 from Group 2 is case 20 from Group 2
NN to 20 from Group 2 is case 1 from Group 1 *
NN to 21 from Group 2 is case 22 from Group 2
NN to 22 from Group 2 is case 21 from Group 2
NN to 23 from Group 2 is case 24 from Group 2
NN to 24 from Group 2 is case 23 from Group 2

(Numbers 13 and 14 are unknowns and appear in Table 7)

Error Rate = 22.72728 %
Table 4. Transformed densitometry data for the 8 most prominent protein variables taken from electrophoretically separated and stained blood proteins of adult leatherbacks nesting in Costa Rica (samples 1-12) and the Virgin Islands (samples 15-24). Also included are two samples from two stranded leatherbacks of unknown origin (samples 13 and 14). The identities of the proteins represented by the data are not known.

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37
In the interest of parsimony, only the most highly
discriminatory densitometry variables were used for maximum
likelihood solutions of the discrimination function. Using the two
most discriminatory variables (protein peaks 3 and 5), or those that
were identified as most powerful in the stepwise analysis by the
program, linear discrimination analysis showed a clear division
between subgroups. Adding one more variable (the third-most
discriminatory as identified through the discrimination exercise) to
the discrimination function caused a better fit; the null hypothesis
that the densitometry values occurred independently of the subgroups
was rejected at the $p=.05$ level (Table 5). When posterior
probabilities were determined as a cross-validation, only two of the
samples in the first group (20%) and three of the samples from the
second (25%) were misclassified using two discrimination variables
(Table 6). Thus, two variables were powerful enough to classify over
75% of the samples correctly using discriminant analysis. The error
rate was reduced significantly when more variables were introduced
into the discriminate function; however, given the small sample
sizes, the risk of overfitting the data remains great if using more
than two variables (A. Solow, pers. com.).

In a separate but related analysis aimed at exploring the
discriminatory power of the protein parameters given samples of
unknown origin, two samples were taken from untagged adults stranded
on a Massachusetts beach and captured in a Rhode Island pound net.
These unknown samples were statistically matched to the most probable
stock through a maximum likelihood estimator (MLE) computer program
Table 5. Discriminant functions generated from total protein stained isoelectric-focusing densitometry data from two groups of adult female leatherback turtle blood proteins, using the 3 most discriminatory variables (protein bands 3 \( a \), 5 \( b \), and 7 \( c \)).

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<td>U.S.V.I.</td>
<td>[ y = 4.03a + 0.20b + 5.37c - 74.21 ]</td>
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Table 6. Results of forward stepwise discriminant analysis of sea turtle blood protein data from densitometry readings. The results presented as number of correctly classified observations in each group. Stepwise discrimination analysis removes one case from the data set at a time, then uses best fit to the discriminant functions (Table 5) to match the case to a class. The three variables used in this stepwise analysis corresponded to protein bands 3, 5, and 7 of the eight most prominent peaks selected. Circled values indicate number of correctly classified cases for each nesting group.

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</tr>
</tbody>
</table>
designed by P. Ihssen and F. Utter (Millar, 1987). Using the binary specific enzyme data (Table 1), both unknowns were matched with 50% likelihood to the Virgin Islands population but showed no statistical affinity to the Costa Rica population. Given the small number of unknown samples used, this MLE test was used only to show the potential practicality of this technique.

The same unknown samples were classified using the nearest neighbor technique mentioned above. Using the nearest neighbor classification technique on the densitometry data, the first unknown (Sample # 13) was classified with the Costa Rican stock while the second (sample #14) was classified with the Virgin Island stock (Table 7).
Table 7. Classification of two blood samples from stranded adult leatherback turtles of unknown origin. The blood proteins were separated electrophoretically, stained for total proteins, and grouped on the basis of densitometry data by a nearest neighbor classification algorithm (James, 1985). Values used for the classification appear in Table 4.

<table>
<thead>
<tr>
<th>Sample #</th>
<th>Nearest neighbor</th>
<th>Sample group of nearest neighbor</th>
</tr>
</thead>
<tbody>
<tr>
<td>13</td>
<td>Case 3</td>
<td>Group 1, Costa Rica</td>
</tr>
<tr>
<td>14</td>
<td>Case 16</td>
<td>Group 2, Virgin Islands</td>
</tr>
</tbody>
</table>
DISCUSSION

There are several lines of inquiry that my data address, all of which may have relevance to leatherback sea turtle population recovery and species management. The results can be used to address three fundamental questions about *D. coriacea* population dynamics: 1) Is the North Atlantic Ocean leatherback population a panmictic assemblage or are these leatherback populations instead divided into not only seasonally spatial, but also genetically distinct demes between which there is little gene flow? 2) If the North Atlantic leatherback population is divided into true stocks, what levels of genetic variability exist in the smaller subgroups and are those levels as dangerously low as they are in other fragmented populations of endangered species? 3) Do existing differences in genetic makeup of stocks allow use of blood proteins as markers for stock identification when tagging studies are limited?

At least two, possibly more, stocks of leatherbacks occur in North Atlantic waters. The demic structure I found in my genetic investigation is also supported by tag returns which indicate that adult females show very little nesting site interchange. One leatherback tagged on Sandy Point, St. Croix, U.S.V.I. did visit the Culebra nesting beaches, but these areas are separated by less than 80 kilometers and this episode was deemed a rare event by researchers studying these nesting group intensively (Eckert and Eckert, 1988). Analysis of epifaunal encrustation of females nesting at Sandy Point also supports the notion of independent demes with little interchange and a high degree of nest site fidelity (Eckert and Eckert, 1988).
The genetic information in my study suggests that not only do nesting females form geographic subgroups but that breeding males do so with the females. Such behavior directly contradicts the theory that leatherbacks assemble at central mating areas where members of different nesting subgroups interbreed as suggested by Lazell (pers. comm.). That an evolutionary strategy in which males exhibit migratory patterns different from those of females, and thus have to search for females in the open ocean, has developed in leatherbacks is difficult to imagine, and has no precedent in any other highly migratory marine species. More likely, males accompany females to the general vicinity of the nesting beaches, where they mate one or more times during the nesting season. The fact that copulating pairs have not been seen near nesting beaches, excepting one case near St. Croix where a presumed mating pair of leatherbacks was seen by a Division of Fish and Wildlife employee who was patrolling offshore (M. Tobias, pers. comm.) only suggests that leatherbacks may not mate close to shore. Because my data show clear genetic distinction between the Virgin Islands and Costa Rican nesting colonies, reproductive separation must exist between males of these colonies.

The results of this study should not be used to make generalizations about the demic structure of leatherback sea turtle populations elsewhere. Two important qualifications limit the general applicability of the findings: first, that samples were not collected from all major nesting populations; second, that sample sizes were small out of regulatory necessity. The optimal sample size question is difficult for many researchers, particularly those
that work with endangered or locally rare species. In this work with endangered and thus federally-protected *D. coriacea* populations, the sample size was restricted by legislative mandate and could not be increased. Regardless, Gorman and Renzi (1979) showed definitively that sample sizes in genetic studies such as mine were less critical than the number of protein variables measured.

In my study of leatherback demic structure, although the assumption of normality could not be made with certainty, the relatively large number of parameters investigated in comparison to the number of samples obtained ensured that the discrimination analysis was justified despite small sample sizes. It must also be stressed that the sample sizes represented relatively large proportions of the nesting populations. Thus 6% of the estimated population nesting in the vicinity of Parismina and a highly significant 25% of the estimated St. Croix nesting group were sampled.

The sample size difficulty was amplified with my use of only two samples from turtles of unknown origin. My purpose in doing the MLE and nearest neighbor classifications exercises was not to draw broad conclusions about the migratory movements of leatherbacks, but rather to demonstrate the usefulness of the discrimination analyses when good baseline information exists. The quality of this baseline information would be greatly improved by larger sample sizes from each of several additional nesting groups. This improved information would make possible assignment of any captured, moribund, or freshly dead animal of unknown origin to the most probable stock, based on a small blood sample.
Since the nearest neighbor method is preferable to the MLE when using small sample sizes (James, 1985), I place greater confidence in the nearest neighbor results and would recommend the analysis for future studies. Again, the small sample size for the unknown group, and secondarily, the sampled stocks, reduces the probability of a definitive outcome for the analyses performed in this study. However, these methods hold great promise for stock identification of stranded leatherbacks, and the use of such analyses should be encouraged whenever possible. This approach can lead to an increased understanding of foraging areas for different stocks, migratory patterns, and the genetic makeup of sympatric demes.

That the Atlantic population of leatherbacks is subdivided into discrete stocks has important implications for management. Bolstering production in one segment of the population, for instance adult females of the small St. Croix - Puerto Rico stock, will probably have little effect on the inclusive population size of the species. In other words, beach-specific management efforts will have only localized effects when population subdivision exists. Any recovery plan for the species, such as is mandated by the U.S. Endangered Species Act of 1973, must acknowledge this natural fragmentation. Also important is the concern that the local area has the resources to support and sustain a population increase if efforts are directed at increasing population size through so-called head-starting and hatchery programs. Many Caribbean nesting beaches have become marginal leatherback nesting sites because of development and recreational use (Meylan et al., 1985). If the population sizes are
suddenly increased, the leatherbacks returning as adults to the natal beach may not find the beach of a quality that permits successful reproduction. This concern has already been voiced with respect to the olive ridley (Leptochelys olivacea) population nesting in Nancite, Costa Rica (Cornelius, 1986). Although the latter species nests en masse in events known as "arribadas" and may thus not serve as a valid basis for comparison, the future resource requirements of leatherbacks should be considered in any management program.

Consequently, each leatherback stock must be treated as a separate entity. In fact, because genetic distinction is so clearly exhibited by the two subgroups investigated, each stock should probably be preserved in an effort to maintain the greatest genetic diversity in the species as a whole. Lande (1988) described how population subdivision can lead to maintenance of genetic variability in even drastically reduced populations. Such a mechanism could account for the relatively high level of heterozygosity in Dermochelys (see below), and strengthens the argument for maintaining each nesting population as a discrete entity. Resource managers should assess the genetic composition of each hypothesized stock and its potential for recovery, and then act in the most efficient way possible to ensure that each stock is preserved. Such research and management action would be an assured way of realizing the goal of maximum likelihood of this species' survival.

There are other grounds for opting for a management strategy in which each population sub-group is protected. That Dermochelys coriacea populations utilize a geographically wide and ecologically
diverse array of nesting beaches could counter the potentially
disasterous effect caused by the loss of a single critical nesting
area, such as could occur in the event of a natural catastrophe such
as a severe storm or extreme sea-level rise. The Kemp’s ridley sea
turtle (<i>Lepidochelys kempii</i>) may be so severely endangered precisely
because it now relies on one major nesting beach, demonstrating the
vulnerability of a restricted nesting locality strategy.

That the North Atlantic leatherback population is subdivided
also suggests that the local stocks are closer to minimum viable
populations (MVPs) in each subgroup than previously thought.
Quantitative values for MVPs in leatherbacks have not been
calculated, due in large part to the discrepancies in estimating age
at first breeding, which have been hypothesized to be as low as 2-3
years (Rhodin, 1984) and as high as 30 years (Carr, 1952). However,
each fragment of the population must be closer to any hypothetical
MVP than the sum total representing the Atlantic population as a
whole (Soule, 1980). For this and the aforementioned reasons, I
argue for increased urgency for the protection of each nesting
population, despite strong local pressures to harvest leatherbacks
for various products in some areas (Groombridge, 1982).

Within the two stocks studied, genetic variability of the
polymorphic proteins was surprisingly high when compared to other sea
turtle species. The mean heterozygosity value as averaged across
seven loci, 0.33, is far greater than found for green (0.12) or
loggerhead (0.03) turtles (Smith et al., 1977). I cannot make direct
comparisons since the higher resolution technique I used should
reveal greater variability than the simpler starch gel
electrophoresis technique of Smith et al. (1977).

Even small stocks such as the St. Croix nesting population show little evidence of inbreeding depression. In addition, the relatively high genetic variability found in the Dermochelys populations studied indicates that no historical bottlenecking effect occurred, an inference consistent with what is known about historical exploitation of the species (Carr and Ogren, 1959).

The results of this study vary in their accordance with predictions made about adaptive variability described in theoretical models. The "coarse-grained" versus "fine-grained" models of adaptation presented by Levins (1968), models which were later adapted for vertebrate comparisons by Valentine (1976), predict that large, highly migratory carnivores such as the leatherback should be highly homozygous. However, other theories such as those elaborated by Selander and Kaufman (1973) predict that the leatherback, being an endotherm (Greer et al., 1973), would have a higher genetic variability than ectotherms in general. In my estimation, validation of these theoretical models using only a few proteins is not possible. Such comparisons are better made using extremely high resolution techniques such as DNA sequencing. Unfortunately, using even such highly objective analyses to prove or disprove evolutionary theory can be misleading, since the level of genetic variability observed is to some extent influenced by the choice of methodology employed by the researcher, who has an a priori expectation of how much variability there ought to be (R. Levontin, pers. comm.)

The levels of resolution obtained in my study of the genetic
variability of leatherback populations is sufficient to delimit stocks. Both of the methods used: the specific enzyme identification and total protein densitometry, supported the same conclusions about demic integrity of Atlantic subpopulations. However, since the methods are backed by different theoretical models and assumptions, it is useful to compare them. Although the total protein densitometry analysis yielded more quantitative information than the specific enzyme staining techniques, the specific staining data showed statistically better discriminating power. More importantly, the specific enzyme staining technique highlighted the differences in gene products. This quantitative information encoded by the binary data reflected a difference with a discernable genetic basis. The specific staining data were, therefore, more appropriate for estimating the stock affinity of the unknown stranded and trapped individuals. For these reasons, I suggest that others may wish to use the specific enzyme staining method, particularly when sample sizes are necessarily small (i.e. from endangered populations).

The analyses used here should be expanded to more rigorously define the population structure of this and other sea turtles species. For the Atlantic population of Dermochelys coriacea, samples should be obtained from Panama, Columbia, French Guiana, Suriname, Guyana, and the Dominican Republic, all important and geographically distinct nesting sites. The blood samples should be analyzed using specific staining techniques and should be run under published conditions, thereby facilitating cross-comparison.

Also needed is a clear definition of the behavior of male leatherbacks, including the extent of emigration from one population
subgroup to another, thus increasing gene flow. A related problem is whether females are polygamous within a single nesting season. One way to address this question would be to sample hatchlings from separate clutches laid by the same female and determine whether inter-clutch genetic variability is greater than within-clutch variability. If so, one could infer that the offspring were fathered by different individuals, presuming that females do not store the sperm of multiple matings together, which is also possible. Similar questions about parentage have been addressed using electrophoretic analysis of blood proteins in other vertebrates (McCracken and Bradbury, 1977). Again, such a question has implications not only for gaining greater knowledge of the species but also for developing more effective management.

*Dermochelys coriacea* may be in danger from both localized population extinctions and gradual but irreversible extirpation. Perhaps more than any other sea turtle species, it requires a diversity of intact habitat types to survive, ranging from highly productive subarctic and temperate open ocean areas to warm, shallow water and coastal tropical habitats. Comprehensively managing threats to this species could mean establishing corridors linking small protected segments of each habitat together in a system providing access to the entire set of marine ecosystems utilized throughout the individual home range. Although not explicitly stated, this is what many conservationists are already attempting to do to through their diverse sea turtle recovery efforts. Each beach protection program, each legislative action mitigating the impact of
fishing and shipping activities on feeding turtles, and each action taken to prevent marine habitat degradation effectively protects one link in the vital chain of habitats needed by endangered sea turtles. But none of these efforts can be systematically beneficial if the most basic question of all, namely, "What is the population to be managed and how does it vary in time and space?" is left haphazardly to idle speculation.
I express gratitude to those people that helped make my fieldwork enjoyable, especially John and Virginia Kollman, Robert Brandner and Sue Brasford, and the Earthwatch volunteers in St. Croix. Special thanks to C. Robert Shoop, Saul Saila and Suzanne Ayvazian, whose tireless offers of help and guidance did not go unheeded. This work was supported in part by a fellowship from the International Women’s Fishing Association, a grant from the Society of Sigma Xi, the Pew Charitable Trusts and the Marine Policy Center of the Woods Hole Oceanographic Institution.
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MANUSCRIPT # 2:

AN EFFECTIVE METHOD TO OBTAIN BLOOD SAMPLES FROM LARGE SEA TURTLES
Obtaining blood samples from sea turtles is becoming increasingly crucial activity for addressing questions about physiology, reproductive biology, genetics, and population dynamics. For small species, such as the ridley turtles (*Lepidochelys kempii* and *Lepidochelys olivacea*), or for juveniles and subadults of larger species, blood may be obtained using the technique of Owens and Ruiz (1980). Their method involves using 21-25 gauge hypodermic needles and an optional angled restraining table in sampling blood from the dorsal cervical sinus. Such blood sampling clearly preferable to cardiac puncture via limb (Friar, 1977) or plastral insertion (Dozy et al., 1964), or to post-sacrificial bleeding, since both of these formerly common methodologies involve significant trauma. However, the blood sampling technique of Owens and Ruiz (1980), using relatively small gauge needles for sampling from the neck region, is not effective for large loggerhead (*Caretta caretta*) or leatherback (*Dermochelys coriacea*) sea turtles, due to large neck girths and the depth of the cervical sinuses. Attempts to obtain blood from the paracervical sinus of leatherbacks were so often unsuccessful that the U.S. Fish and Wildlife Service actively discouraged researchers from continuing sampling efforts (S. Furness, pers. comm.). Blood work on this species has thus been limited.

A relatively easy and extremely efficient method to withdraw blood from *Dermochelys* and *Caretta* is to use a 13 gauge spinal tap needle. These needles are 5 centimeters long, have a highly beveled
tip, and can be inserted into the neck of any large sea turtle specimen without any apparent discomfort to the animal. This is an especially important consideration since samples are most easily obtained during the period when females are depositing eggs. Traumatizing females during the breeding season might cause physiological or behavioral interference with reproduction, which must be avoided with endangered species.

Both the needle and the syringe barrel should be coated in heparin before insertion to avoid clot formation, since sea turtle blood coagulates very quickly when exposed to air. Sampling may be done only on relatively still individuals, such as nesting females that have deposited their eggs but have not begun covering the nest. The needle should be inserted in the paracervical region of the neck just lateral to the vertebral column (Figure 1). The needle must be perpendicular to the dorsal plane of the neck (approximately vertical when the animal’s head is resting on the ground). Suction should be applied to the syringe as the needle is inserted to its base and during withdrawal. Blood is most easily obtained from the paracervical sinus as the syringe is pulled upwards and the needle travels through the sinus.

Twenty-four, 10 ml blood samples were obtained from adult female leatherbacks nesting in Costa Rica and the U.S. Virgin Islands and over thirty 5 ml samples were collected from loggerheads nesting in Mexico (Agardy, 1988) using this method. The majority (40) of the turtles sampled were tagged and observed up to 2.5 months after sampling, and none showed after-effects or complications from the
Figure 1. Needle insertion point for blood sampling from adult leatherback turtles.
technique. Blood samples can be stored for later use in vacutainers containing ETDA as an anticoagulant.
LITERATURE CITED


MANUSCRIPT # 3:

WHY INFORMATION ON POPULATION DYNAMICS IS CRITICAL TO THE CONSERVATION
OF ENDANGERED SPECIES: LESSONS FROM SEA TURTLE RECOVERY ATTEMPTS
WHY INFORMATION ON POPULATION DYNAMICS IS CRITICAL TO THE
CONSERVATION OF ENDANGERED SPECIES:
LESSONS FROM SEA TURTLE RECOVERY ATTEMPTS

ABSTRACT

Managers of endangered animal populations are often faced with
the task of making management decisions without the luxury of having
complete knowledge about the species. The problem is especially acute
in cases where the body of knowledge about an animal’s ecology
pertains only to a certain portion of its life cycle. Sea turtle
recovery efforts are a perfect case in point: managers worldwide are
struggling to prevent local and sometimes global extinctions while
knowing little about the species’ demographic parameters, beyond that
which is know about nesting females and emergent hatchlings. However,
the total time that a sea turtle spends on land within the realm of
convenient study is less than one percent of its life span.

Given that resources to study sea turtle ecology are limited
and that time, especially for some critically endangered species, is
short, managers must be presented with the kind of information
necessary for formulating efficient recovery attempts. In the case
of highly migratory marine animals such as sea turtles, the most
important ppopulation data are the following: 1) what is the size
and extent of the population unit to be managed?, and, 2) what is
that population’s intrinsic rate of increase and how is it prevented
from being fully realized? Without answers to these basic
prerequisite questions, other data collected on the biology of the animals cannot be framed in an accurate context for developing sound management.
INTRODUCTION

There is a Hungarian colloquialism that can be loosely translated to mean "those that work the hardest are those that have to". In Hungary the expression was used long ago to justify a certain smugness exhibited by those who lived quite well without seeming to have expended much effort. But it can also mean the reverse that in critical situations, hard work is demanded, and that in such situations half-hearted or misdirected efforts are insufficient and sometimes even counterproductive. For those who focus their attention on the recovery of endangered species, hard work is an unconditional and accepted necessity. Buying time to gain more complete knowledge is a luxury that may be afforded in basic research but which is of little use in developing the applied scientific tools needed for crisis management. Those working to save a species from extinction are under more pressure to be efficient despite a limited information base, and they thus work harder because they must.

The management of sea turtles, those pantropically threatened marine reptiles which have become the center of much conservation activity and environmentalist interest, provide a salient example of how several recovery attempts and a great deal of hard work may be wasted because critically important information about species dynamics is lacking. I illustrate this possibly unpopular opinion with a description of the case of the leatherback turtle, Dermochelys coriacea, in the North Atlantic Ocean.
The leatherback is extraordinary in many regards. Unlike the other species of shelled sea turtles that comprise the family Cheloniidae, the leatherback sea turtle is the only living species in the family Dermochelyidae and has a skin-covered shell. However, it shares some morphological characteristics with its super-familial cohorts, namely a streamlined body and flippers as adaptations for aquatic locomotion, a non-retractable head, and terrestrial egg deposition and incubation. Beyond these similarities the leatherback is distinctive from other sea turtle species in the following features: a "soft" shell which lacks the keratinized laminae found in other sea turtle species (Frazier, 1987), a tough dermis including a mosaic pattern of bones embedded in the tough connective tissues of the skin, a counter-current heat exchanger which allows the animal to maintain its body temperature at least 18°C higher than the ambient water temperature (Greer et al., 1973), large keratinized buccal and esophageal papillae, an elongated esophagus used for food storage, proportionally large anterior flippers, and the enormous size of the adult animal, sometimes exceeding 900 kilograms (Eckert and Luginbuhl, 1988).

There are also behavioral characteristics exhibited by the leatherback not common to the green, Kemp's ridley, hawksbill, or loggerhead turtles with which it shares Atlantic waters. These include more extensive migrations from tropical nesting grounds to temperate and even sub-arctic waters (Lazell, 1980), its extensive feeding on medusoid jellyfish such as *Cyanea capillata* (Brongersma, 1972), and deep-diving to depths exceeding 780 meters (Eckert et al., 1986). The long distance movements and predominantly pelagic
lifestyle of the species, are directly related to its unusual food preference, explain why even less is known about the biology of leatherbacks than about the other sea turtle species.

What is known about the leatherback turtle comes from disparate bits of information about its marine ecology (Eckert and Eckert, 1988; Lazell, 1980); extensive data recording certain aspects of its terrestrial ecology such as nesting behavior and egg survivorship (Deraniyagala, 1939; Whitmore and Dutton, 1985; Hirth and Ogren, 1987; and Pritchard, 1971), especially in some well-studied areas; and fragmentary historical records about its abundance and apparent population decline (Carr, 1952; Ross, 1982). The animal is considered endangered throughout its range in all the world's oceans (Groombridge, 1982), although estimates of historical population size and even current numbers are extremely tenuous. What estimates have been made (Pritchard, 1982) are based on beach surveys during the nesting season and some short-term time series data obtained from tagging studies. But it is not certain that all the world's major leatherback nesting beaches have been discovered, and we know nothing about the population sizes of the majority of demographic sectors, including adult males, non-breeding females, juveniles, and hatchlings beyond one to one day of age.

The sequential revisions of *Dermochelys* world population estimates made in the last thirty years illustrate our ignorance of the species. In 1961, Fitter (1961) estimated the population as being composed of only 1000 breeding pairs. By 1971, that estimate had been revised upward over fifty-fold when Pritchard estimated the
number of breeding females at 29,000 to 40,000 individuals. Following the discovery of a major nesting ground in Pacific Mexico, Pritchard (1982) changed his estimate to 115,000 mature females worldwide. Verification of these estimates is lacking, however.

A quick glimpse at this kind of demographic record might give the mistaken impression that the leatherback turtle population is growing at a fast rate, rather than diminishing. However, the apparent increase reflects more the elusive nature of the critical data rather than any trend in population size. Ross (1982) extensively documented the decline of the leatherback in specific breeding aggregations, and he and other authors (Meylan et al., 1985; Pritchard, 1982) conceded that the leatherback’s endangered status is warranted.

The decline of *Dermochelys coriacea* may be attributed to several compounding factors. In addition to the grave impacts caused by accelerated loss of nesting habitat, possible interference with migratory and feeding behavior, and probable increases in the natural predation induced mortality of eggs and young, the leatherback has suffered increasing rates of slaughter at the hands of man. Although never hunted in large numbers for its meat, leatherback oil has been treasured as a cure for everything from arthritis to head colds, and is commonly used as a lubricant (Bustard, 1973; Carr, 1971). Inhabitants of many Caribbean islands have now turned to leatherback meat as a supplement to ever-decreasing supplies of meat from preferred species (Meylan, et al., 1985). Leatherback meat may even be eaten by Inuits of Baffin Island in lean times (Shoop, 1980). Around the world leatherback eggs carry the cross-cultural and
decidedly dubious distinction of commanding a high price as aphrodisiacs, and both the legal and black market trade in eggs is thriving (Meylan et al., 1985). Furthermore, anthropogenically-induced, indirect impacts on the population, such as collisions with ships, entrapment in fishing gear, morbidity and mortality from contact with oil and tar, and death from the ingestion of plastic and other debris, continue to rise at an alarming rate (Balasz, 1985; Carr, 1987; and Laist, 1987).

The leatherback turtle appears threatened with extinction. But if this or any other sea turtle species does go extinct, it will not be for want of human concern. Much recent environmentalist interest is focused on sea turtles (Canin, 1989), which may now rival whales as the symbolic object of pity and advocacy (R. McManus, pers. comm.). The rise in public awareness and the international interest in endangered species protection has helped promote sea turtle conservation projects around the globe (IUCN General Assembly Report, 1988). But are these efforts aimed at recovering sea turtle populations as efficiently and quickly as possible? I argue that in the case of *Dermochelys coriacea* they are not, because some of the most fundamental and critical questions about the species have neither been asked nor answered.

Management practices determine where problems are and aim to deduce what the causes of those problems might be. In the case of endangered species management, the problem is obvious: a downward demographic trend that threatens to bring the population dangerously near the minimum size needed for its self-perpetuation (the minimum
viable population size). The problem statement is accepted a priori. This is the case in the management of the leatherback turtle in the North Atlantic region. But the target for possible management is unknown. Whether it is the worldwide population, the species as a whole, the population found within an ocean basin, such as the Atlantic, or the female population that returns on one to three year cycles to a particular nesting beach has not been elucidated. Without defining the population to be recovered as a management unit, with respect to its size, extent and dynamics, a manager cannot hope to begin the kind of efficient recovery plans that the situation desperately demands.
POPULATION DYNAMICS INFORMATION AS CRITICAL TO MANAGEMENT

The field of population biology concerns itself with how a group of organisms move, change, and replace themselves in space and time. These are the dynamics that need to be investigated for a population, or management unit, to be defined. We are exceedingly ignorant about the population biology of the leatherback turtle. Whether the North Atlantic population is contiguous or comprised of independently-functioning demes is unknown. We know that breeding females exhibit nest site fidelity, are iteroparous, and return to the same beach to lay eggs, sometimes over several seasons. In this regard, then, we know that at least some of the time segments of the Atlantic population are subdivided. But until recently it was not known whether the subdivision corresponded to true demic segregation: a genetically rather than temporally defined subdivision.

We also do not know what the rate of population replacement is in any population we define. At least some of the factors that prevent this natural population replacement rate (whatever it may be) from being realized have been elucidated, albeit not systematically. But the two most fundamental questions are those concerning the definition and intrinsic dynamics of the population to be recovered, and these have been ignored in the fervor to save this species from extinction. And because they have not been addressed, we cannot judge the efficiency of any management activities to know if they are helping to boost the Atlantic population as a whole, or merely protecting a small local population, or, in the extreme, not helping at all.
For the management of the Atlantic population of leatherbacks to be planned effectively with a sound rationale, a study of the species' population dynamics is a first order-exercise. Assuming consensus concerning the nature and severity of the decline in numbers, a program aimed at providing the necessary information to managers might aim to address the following questions in the following order:

1) What is the size and extent of the population to be controlled or recovered - in other words, what is the management unit?

2) What is the population replacement rate of this management unit?

3) What are the natural and anthropogenic factors interfering with the natural rate of replacement? And what factors must be considered to enhance or maintain normal genetic variability in populations with reduced sizes, recognizing that the fitness of a population is as, if not more, important as its size?

4) Which of the above factors can be controlled through management measures and which of those will yield the fastest, most tractable results?

Complete knowledge about a species is not a prerequisite for elaborating the demography of a population or modeling trends in population size. The population units themselves can be defined using sophisticated techniques that do not require enormous amounts of data collection time, and models can be developed relatively easily which can be used to generate predictions of the population's
dynamics over time. Conservation biologists already acknowledge the value of such work, however in most cases this message has not been relayed to natural resource managers or administrators (Lande, 1988).
LEATHERBACK SEA TURTLE POPULATION STRUCTURE

Leatherback turtles, although relatively rare in occurrence, are ubiquitous in North Atlantic and Caribbean waters (Shoop et al., 1981). During the spring and summer breeding season, leatherback females come ashore to nest in Costa Rica (Hirth and Ogren, 1987), Panama (Meylan et al., 1985), Columbia (Ross, 1982), French Guiana (Fretey and Lescure, 1979), Dominican Republic (Ross and Ottenwalder, 1983), and some of the Lesser Antilles (Figure 1). Several other Caribbean localities support fewer numbers of nesting females (Groombridge, 1982; Pritchard, 1971). Aerial surveys show that leatherbacks are also found in offshore waters from Cape Hatteras to Nova Scotia (Shoop, 1987; Shoop et al., 1981), Cape Hatteras to Key West (Thompson and Shoop, 1983), and in the Gulf of Mexico (Fritts et al., 1983). In the summer and fall months, some leatherbacks become entrapped in various kinds of fishing gear set in temperate waters (Goff and Lien, 1988). During the same time of year, dead and moribund animals are found washed ashore along the eastern seaboard of the United States and Canada, particularly in New Jersey, New York, New England, Nova Scotia, and Newfoundland. The normal home range for this species thus appears to extend over 5000 kilometers (Pritchard, 1976), with hypothesized migration routes (Lazell, 1980) carrying individuals from tropical nesting sites to sub-arctic feeding areas (Figure 2).

The North Atlantic population of leatherback turtles has been assumed to be a panmictic assemblage with only seasonal segregation related to breeding patterns. Conservation attempts in the Atlantic
Figure 1. Major nesting beaches used by Dermochelys coriacea in the Caribbean region
Figure 2. Hypothesized migration routes for the leatherback turtle (after Lazell, 1980).
and Caribbean region, on the other hand, have always been piecemeal and directed as if towards discrete but local population units. The discrepancy between these two underlying assumptions is obvious. Yet no literature record exists which defines any Dermochelys population as a management unit.

I attempted to resolve this problem for two widely separated nesting populations using electrophoretic analysis of blood proteins to determine whether separate leatherback nesting populations were genetically distinct. The resultant data and analyses did not support the null hypothesis that the Costa Rica and Virgin Islands nesting groups constituted geographical extremes of a contiguous Atlantic population (Agardy, 1989a). My data suggested the Atlantic population of leatherbacks is actually comprised of at least two, although probably many more, discrete demes. This clearly implies that any management efforts must take the genetic delineation into consideration. More than one management unit exists, but how many units requiring independent but coordinated conservation efforts has not been determined. The definition of management units certainly merits immediate attention.

How can the Atlantic populations of Dermochelys coriacea best be delineated and defined? I suggest that the most cost-effective and time efficient method is to employ techniques that elucidate genetic structure of the nesting populations. Fisheries biologists commonly use genetic analyses to elucidate stock structure, but their task is often simplified by the ease with which enormous numbers of samples can be collected. However, there are statistical tools which make it
possible to hypothesize stock structure with a high degree of confidence even when sample sizes are small. SIMCA, a multivariate discrimination program developed for analytical chemistry, is one such method (Saila and Martin, 1987). There are numerous other discrimination techniques that also may be applied, including the stepwise discrimination method that I chose for the leatherback stock identification work (Agardy, 1989a). Although the sample sizes were necessarily small in the leatherback study due to regulatory restrictions on number of samples permitted, the analysis proved rigorous. Additional support for using such algorithms despite few sample is provided by Gorman and Renzi (1979) who contend that small sample sizes will not distort estimates of genetic heterozygosity as long as many loci are investigated.

Electrophoretic analysis of blood proteins can be an effective way to look at demography through the eyes of genetics. Blood can be sampled easily from leatherbacks using a 13-gauge spinal tap needle inserted into the paravertebral sinus of the neck (Agardy, 1989b). Only small amounts of blood are needed for electrophoretic analysis (3-5 ml), and sampling appears to have no ill-effect on adult turtles. Blood is most easily drawn from nesting females that have completed egg-laying, but blood may also be sampled from captured or stranded animals.

There are a number of alternatives a researcher can choose when analyzing blood proteins electrophoretically; these include looking for known allelic variants of metabolic enzymes (specific staining technique) or investigating all the proteins present in the sample without knowing specifically what their genetic correspondence might
be (general protein densitometry). The advantages of the former method are that immediate visual results are produced and that the data can be used in estimating inbreeding coefficients. On the other hand, the advantage of the total protein method is that it generates a large data set, making more statistically powerful statements possible.

Aspects of population genetics are extremely important in the applied field of conservation biology. When population numbers are low, as they are de facto in endangered species, inbreeding and genetic drift effects which lead to increased genetic homozygosity must be monitored. Though demography is probably more important a science for conservation than genetics (see Lande, 1988), genetic research is one way to describe the demographic characteristics of open populations.

Do using the tools of genetics constitute the best way to gain critical information about sea turtle demography? Probably yes, if efficiency is a consideration. There are other ways to define sea turtle populations, however, and some have been practiced for many years. The most common method used to discern stock structure is through the tagging of adult females. Monel or Inconel cattle ear-tags are affixed to the trailing edge of either the front or rear flipper (or both) of a terrestrially nesting female. Long experience with tagging has refined the methods used, so now large numbers of turtles are tagged each year with little disturbance to the animal and, typically, only moderate tag loss. The problem with this technique is that it is time intensive. True stock structure can
only be determined through saturation tagging (marking all available individuals) over an extensive area and "recapturing" marked individuals year after year to determine adult turnover rate (ars of replacement or recruitment). And since only nesting females can be tagged in large numbers, tagging provides little information about the other segments of the population.

Recent technological advances could allow marking hatchling sea turtles with implanted microchip or passive transponders. Though feasible, the methods are inherently limited. Hatchling and juvenile sea turtle natural mortality is presumably so high that huge numbers of young need to be tagged for the recapture of even a few adults, and these microchip tags are currently very expensive. Add to that the time horizon needed to obtain results; leatherbacks may take thirty years or more to mature (their growth rate is highly controversial). Thus saturation tagging of hatchlings requires the presence of researchers at the natal and nearby nesting beaches many years after tagging. In the meantime, improper management measures could theoretically hinder the species' recovery.

There are tracking methods which allow migratory and local movements of a tagged individual to be monitored directly. These include conventional radiotracking methods from the surface or by satellite (e.g. Standora et al., 1984). The drawback of these methods is that they require extensive field observation and cannot be performed on large numbers of animals at once without exorbitant operating costs. Furthermore, location alone will not provide adequate information to delineate stocks or elucidate demography.
Another method being currently investigated as a possible tool for defining stock structure of sea turtles is mitochondrial DNA (mtDNA) analysis (Bowen et al., 1989). The underlying assumptions and the basic techniques used are similar to those used in conventional electrophoretic studies, but the genetic material of cells rather than their products is analyzed. Mitochondrial DNA is easily obtained from sea turtle tissues, especially from high metabolic tissues such as heart, liver, and other organs (Avise, 1987). Since mtDNA analysis is a higher resolution technique than conventional electrophoresis, and since it looks directly at the genome, it is necessarily less subjective than the latter technique. However, mtDNA analysis has two serious shortcomings when used to study sea turtle population structure: 1) sampling usually requires sacrifice of the animal, and 2) it does not shed light on the population genetics of males since mtDNA is maternally inherited.

Electrophoretic analysis of the metabolic enzymes present in Dermochelys coriacea blood remains the fastest, most effective method of elucidating stock structure and defining the demographic unit in need of sound management. When this information is supplemented by data generated from other kinds of study, including mtDNA analysis and tag return statistics, a valid and rigorous basis for instigating management can be established.
ESTIMATING THE NET REPLACEMENT RATE OF SEA TURTLE POPULATIONS

Once the management unit has been identified and a clear picture of population movements emerge, the next information needed by a resource manager is an estimate of the rate of replacement for the population ($R_0$). For an organism that spends each stage of its life within the realm of scientific observation, the natural mortality in each age group can be readily quantified. The case is infinitely more complex for sea turtles, since the animals spend only a fraction of their lifetime in areas where they can be easily studied, and since offspring cannot yet be followed to maturity. For some species such as the green turtle, *Chelonia mydas*, some life table information can be obtained from intensive studies at nesting beaches and in places where the animals are kept in captivity (Bjorndal, 1980; Thompson, 1980). Since leatherback sea turtles cannot be held in captivity for long periods (Bels et al., 1988) and because they are highly migratory, little definitive information on survivorship exists.

Given the crucial need for this type of demographic information and the lack of knowledge about leatherback biology, significant scientific effort should be invested in demographic studies. Simulation models with different mortality rates at different life history stages or different rates of growth lead to astonishing predictions and further emphasize the need for verified demographic estimates. For instance, if one assumes that the survivorship pattern in leatherbacks is similar to that of other sea turtles that
are better known, such as the loggerhead, *Caretta caretta*, (Frazer, 1983), a hypothetical life history table can be constructed for modeling work (Table 1). From Leslie matrix model simulations using these data (Appendix B), results indicate that the most important population sectors for increasing population growth are the hatchling, juvenile and newly mature age classes. For instance, by changing the survivorship parameter for hatchlings by a mere 10% increase in value, the simulated population growth trend changes dramatically from a gradually declining population heading towards extinction to a relatively stable but slightly increasing population (Figure 3). Various scenarios incorporating different survivorship values for all stage classes are given in Table 2; estimated lambda values less than one indicate population decrease and eventual extinction, greater than one population increase and recovery.

These results are consistent with those demonstrated in loggerhead sea turtle population simulations (Crouse et al., 1985). For this more rigorous modeling exercise, a data set on the reproductive biology of loggerheads nesting on Little Cumberland Island, Georgia, was combined with survivorship estimates calculated for the same population (Frazer, 1983) to generate a stage-based population model. This work also implies that if the hypothesized demographic parameters used for modeling are accurate, the most critical component of the population for replacement value is the large juvenile age class.

In my hypothesized leatherback population simulations, the results are based on the assumption that the rate of growth is slow, with leatherback females taking 20 years or more to achieve
Table 1.

Hypothetical life history parameters for *Dermochelys coriacea*, used as the basis for Leslie matrix simulations. This table includes: \( l(x) \), survivorship from the egg to any subsequent stage; \( p(x) \), stage to stage survivorship; \( m(x) \), number of eggs deposited every two years by each breeding female divided by two (assuming a 1:1 sex ratio of offspring); and \( R_0 \), the finite rate of population increase, where \( R_0 = l(x)m(x) \).

<table>
<thead>
<tr>
<th>Stage</th>
<th>( l(x) )</th>
<th>( p(x) )</th>
<th>( m(x) )</th>
<th>( l(x)m(x) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Egg</td>
<td>1.00</td>
<td>.85</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Hatchling</td>
<td>.0425</td>
<td>.05</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Young Juveniles</td>
<td>.0043</td>
<td>.10</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Older Juveniles</td>
<td>.0006</td>
<td>.15</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Neophyte Breeders</td>
<td>.0005</td>
<td>.90</td>
<td>220</td>
<td>.132</td>
</tr>
<tr>
<td>Remigrants</td>
<td>.0004</td>
<td>.80</td>
<td>220</td>
<td>.110</td>
</tr>
<tr>
<td>3rd Time Breeders</td>
<td>.0003</td>
<td>.70</td>
<td>220</td>
<td>.088</td>
</tr>
<tr>
<td>4th Time Breeders</td>
<td>.0002</td>
<td>.60</td>
<td>220</td>
<td>.044</td>
</tr>
</tbody>
</table>

\( R_0 = .374 \)
Figure 3. A comparison of simulated population trends, first (A) using hypothetical survivorship estimates given in Table 2; then modified (B) to increase juvenile survivorship by 10% and decrease egg survivorship by 25%.
Table 2. Survivorship values for leatherback turtle stage classes used in the Leslie matrix simulations and resultant eigenvalue (lambda) values. A lambda of less than one indicates eventual extinction.

<table>
<thead>
<tr>
<th>Stage class</th>
<th>Stage-specific survivorships</th>
</tr>
</thead>
<tbody>
<tr>
<td>P[0] eggs</td>
<td>.85  .85  .85  .60  .85  .85</td>
</tr>
<tr>
<td>P[1] emergent hatchlings</td>
<td>.05  .05  .15  .15  .15  .05</td>
</tr>
<tr>
<td>P[2] young juveniles</td>
<td>.10  .10  .10  .10  .10  .10</td>
</tr>
<tr>
<td>P[3] older juveniles</td>
<td>.15  .15  .15  .10  .10  .10</td>
</tr>
<tr>
<td>P[4] neophyte breeders</td>
<td>.90  .90  .90  .90  .90  .99</td>
</tr>
<tr>
<td>P[5] remigrant</td>
<td>.80  .50  .50  .80  .50  .99</td>
</tr>
<tr>
<td>P[6] 3rd time breeder</td>
<td>.70  .50  .50  .70  .50  .99</td>
</tr>
<tr>
<td>P[7] 4th time breeder</td>
<td>.60  .50  .50  .60  .50  .99</td>
</tr>
<tr>
<td>P[8] 5th time breeder</td>
<td>.50  .50  .50  .50  .50  .99</td>
</tr>
</tbody>
</table>

Approximate eigenvalues (\(\lambda\))

|     | .894 | .854 | 1.02 | 1.00 | .964 | .948 |

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reproductive maturity. If, however, a faster growth rate of only 3 years to maturity is used (as suggested by Rhodin (1985)), the transition probability for stage to stage survivorship must be altered. If this is done, the results reveal that the hatchling sector is most critical in promoting population increase.

Population simulations are sensitive to errors in parameter estimation, and can generate misleading conclusions. Some parameters are more critical than others; in these simulations, age at maturity and related transition probabilities are extremely critical parameters. If these models are used to justify management, incorrect estimates of survivorship probabilities could lead to disastrous management. One could develop a set of demographic scenarios for differing growth and development rates and then wait for more definitive information on growth to be obtained, but all will be for naught if length of reproductive period of breeding adults remain unknown. Consequently, the major thrust of research efforts should be focused on these aspects of sea turtle biology. Deriving further demographic information on leatherback and other sea turtle species is critical now; without it conservationists and managers of sea turtle populations cannot gauge the success of their work.
MANAGEMENT IMPLICATIONS OF LEATHERBACK TURTLE DEMOGRAPHY

The most obvious implication of the preceding discussion is that further study of sea turtle demographics is desperately needed and is a prerequisite to effective management. However, there are some important messages contained in the demographic information already accruing. First, the non-contiguous nature of the Atlantic population of leatherback turtles casts doubt on the usefulness of treating the North Atlantic leatherback population as a single entity. Each population unit or deme must be managed independently for recovery. If, conversely, the results of the genetic analysis had provided evidence for a panmictic Atlantic population, then managers responsible for the recovery of leatherbacks could set priorities for management areas - and efforts to preserve a relatively minor nesting group such as the Virgin Island population might be discouraged in favor of a better investment elsewhere.

But independent demes do exist in the structure of the North Atlantic leatherback turtle population. Each stock has a unique genetic composition, therefore each gene pool should be conserved, because the genetic character of a population group determines how well that population will be able to adapt to change over evolutionary time. Considering the possibly small demic sizes, genetic drift is a serious possibility. We have no idea what the adaptive landscape, as expressed by the Wright (1969), will present to future leatherback generations in the form of evolutionary challenges. A high degree of heterozygosity within the species or
within a geographical subgroup as a whole will help probably allow an adequate evolutionary response to change. If, through ignorance or misdirected management, small and seemingly insignificant demes become extinct, the genetic variability of the species and its ability to cope with environmental change may be undermined.

Perhaps the logical conclusion is that all sub-populations in the Atlantic region should be managed with equal effort. However, there are complicating factors generated by this newly documented picture of stock structure. If the North Atlantic population is indeed fragmented into discrete demes that roughly correspond to nesting groups, then perhaps some sub-populations must be closer to minimum viable population sizes than assumed. The Virgin Island nesting population, probably comprised of fewer than 100 breeding females, may belong to a deme of less than 200 adults, assuming an equal sex ratio. If one believes the minimum effective population size Rule of 500 (Franklin, 1980), then this deme may be in grave danger of slipping below threshold levels. The situation is made even more critical if one assumes a sex ratio of males to females of less than one and males are not polygamous, or when adult recruitment from a dwindling juvenile population is decreasing.

The logical counterargument is that there is no reason to believe the Virgin Islands stock is near the minimum viable population size, since the population numbers appear to be stable over the short term of seven years (Eckert and Eckert, 1985). Possibly, the deme is comprised of individuals making up the Virgin Islands, Culebra, and possibly the Dominican Republic breeding groups together. Again, the desperate need for further genetic and
demographic studies is emphasized. But one point remains clear: the further fragmenting of an already small population such as the deme outlined above could bring its subgroups precariously near minimum effective population size and thus potential population collapse.

Simulations performed to study how the rate of replacement varies with changing life table parameters may elicit further caution. If the older juvenile sector is the most important in maintaining or increasing population size (Crouse, 1985), then management measures augmenting survivorship of population subgroups, especially of those populations near minimum viable population size, should be implemented. No conservation activities have been deliberately undertaken to protect leatherback turtles of the larger juvenile and newly mature age groups. Even Turtle Excluder Device (TED) regulations aimed at reducing mortality of sea turtles caught incidentally in shrimp fishing operations will not significantly increase survivorship of leatherbacks, since only rarely do fishing operations in the Gulf of Mexico and Southeast U.S. impact leatherback turtles.

At present, enormous quantities of money, coming from both governmental agencies and private foundations, are spent to conserve the earliest age group of sea turtles: eggs and emergent hatchlings. This potentially constitutes a poor choice in allocation of funds. However, the positive outcome of this investment has been to increase both public awareness and interest in sea turtles, which may in turn to increased funding opportunities for better protection.

I know of no data which confirm any specific age class of
Dermochelys coriacea as being most sensitive in altering the rate of population increase. However, there is anecdotal evidence suggesting the mature age classes are more critical than the egg stage of development. On the Caribbean coast of Costa Rica, a 25 kilometer nesting beach between Puerto Limon and Tortuguero, is heavily used by nesting leatherbacks. Egg poaching in this area has occurred at a sustained high rate for many years. My interviews with members of the local communities reveal that this practice goes back many generations, and is an integral component of the culture of inhabitants. Many young poachers conceded that they learned their trade from a parent or grandparent. During the nesting season, scarcely a nest is left in the sand. How, then, is this population able to sustain itself?

One answer, which is purely speculative, is that leatherbacks have evolved their highly fecund reproductive patterns to sustain a high natural mortality of eggs and hatchlings. Prior to coastal settlement in these areas, human predation of eggs was not in the ecological equation for leatherbacks. However, humans in some areas now displace natural predators, including jaguars, seabirds, and coatimundis, in competition for sea turtle eggs. The rates of this human predation have probably increased steadily over the decades of settlement in the area. Leatherbacks may have an adaptive strategy that ensures that at least some clutches will survive to hatching: the apparently unpredictable timing of the onset of nesting. Since villagers said they were unaware that turtles were nesting on their beaches until the season was well underway, many early clutches (and probably the last clutches) were missed and thus may have survived to
hatching.

If some nests are successful, why then are the leatherback populations declining? One answer may be because adult mortality, once extremely low due to the absence of natural predators on adults, may be increasing. More and more large immature and adult leatherbacks are becoming entangled in fishing gear (R. Prescott, pers. comm.) and more may be dying from ingesting plastic and other debris (Balasz, 1985). Perhaps, from a population dynamics perspective, high egg loss is less detrimental than high adult mortality. Such a situation would confirm Pritchard's (1980) comment "I am still convinced that those individuals that have survived the vicissitudes of their long pre-maturity period, namely the breeders, are the most important to protect."
CONCLUSIONS

It has not been my intention to claim that the collective scientific efforts of sea turtle biologists and conservationists have been consciously misguided when it comes to providing the necessary information for the management of *Dermochelys coriacea*. Nor have I meant to belittle the efforts of the myriad of groups working to save this species from extinction. Beach patrols, the guarding of eggs and young, translocations of eggs, and other common conservation practices cannot be considered counterproductive, even if their end result is a negligible increase in population size and incrementally minor additions to our state of knowledge. But program administrators and management agencies should be aware that their conservation efforts will not be efficient, either with respect to time or finances, unless further demographic information on *Dermochelys* is obtained promptly. I believe consideration should be given to the strong possibility that the significant effort and money needed to run costly programs with unknown returns on investment, such as head-starting projects for many sea turtle species where hatchlings are reared a year or more prior to release, should be reallocated to studies which further elucidate stock structure, growth curves, and life history parameters. With limited resources to undertake both scientific study and conservation, and an already short time horizon that is foreshortened with every passing day, we must work hard at being efficient.
ACKNOWLEDGEMENTS

I thank Nat Frazer and Deborah Crouse for sharing their ideas about the importance of sea turtle demography work, and Brooks Martin who so gallantly came to my rescue when I struggled with the population simulations. Thanks also to Karen Eckert, who ran the Earthwatch-sponsored program in St. Croix with great enthusiasm and dedication, who remained supportive throughout my study, and who continues to be an inspiration. Special thanks are in order to Saul Sails and Robert Shoop, who convinced me despite my Hungarian biases that hard work is not only necessary but also fun. This work was supported in part by the Pew Charitable Trusts and the Marine Policy Center of the Woods Hole Oceanographic Institution.
LITERATURE CITED


Agardy, M.T. 1989b. A note on an easy and effective method to obtain blood samples from sea turtles. Ph.D. Dissertation manuscript, University of Rhode Island.


APPENDIX A: STAINING REACTIONS FOR IEF GELS
APPENDIX A: Staining reactions for Iso-electric focusing gels.

The data from this investigation come from two distinct methodologies used to decipher population differences between two geographically distinct nesting groups of leatherback sea turtles. The null hypothesis tested was the same in both investigations: that the two subgroups sampled exhibit no group differences with respect to electrophoretically detectable characteristics. Both methodologies were used on single blood samples taken from each sea turtle.

The first data set (Table 1, Manuscript 1) described whether a particular enzyme variant was present in the blood sample. All sea turtle blood contains the common enzymes needed for metabolism, but genetic variants of these enzymes (allozymes) differ from individual to individual. These allozymes are functionally identical but structurally different in iso-electric point (the pH at which the molecule is electrostatically neutral). Staining the gel with an electron-transfer dye which detects a particular enzyme (and in fact uses that enzyme in the chain reaction needed for staining) results in visual detection of all allozymes present in the sample. The staining reactions for each enzyme investigated are given on the following pages (Figures 1 and 2). The binary data thus correspond to the presence or absence of a particular allozyme as detected through use of an iso-electric focusing apparatus and subsequent staining.
The second data set (Table 3, Manuscript 1) revealed the total protein complement present in the same blood samples. Using this method, the researcher stains the sample for all proteins, rather than for a specific metabolic enzyme. The gel banding patterns are read by a scanning integrated transmission densitometer. The value representing each protein band is an estimate of the areal extent of the band as it sits on the gel. These raw densitometry data were then arc-sine transformed and compiled for discriminant analysis, according to the methods of Fabrizio (1986).
Figure 1. Staining reactions for LDH, PGM, 6-PGD, and PEP.
EST

\[ \text{Carboxylic Acid Anion} \rightarrow \text{Naphthyl Ester (Acetate)} \rightarrow \text{EST} \rightarrow \text{Fast Blue } \beta \rightarrow \text{Dye} \star \]

HK

\[ \text{ATP} \rightarrow \text{Hexokinase} \rightarrow \text{ADP} \rightarrow \text{Glucose} \rightarrow \text{Glucose-6-P} \rightarrow \text{66PDH} \rightarrow \text{6-Phosphogluconate} \rightarrow \text{NADP} \rightarrow \text{NADPM} \rightarrow \text{PMS} \rightarrow \text{MIT} \rightarrow \text{Formazan} \star \]

GAPDH

\[ \text{D-Glyceraldehyde 3-Phosphate} \rightarrow \text{Orthophosphate} \rightarrow \text{GAPDH} \rightarrow \text{3-Phospho-D-Glycerol Phosphate} \rightarrow \text{NAD} \rightarrow \text{PMS} \rightarrow \text{NADH} \rightarrow \text{MIT} \rightarrow \text{Formazan (Blue Stain)} \star \]

Figure 2. Staining reactions for EST, HK, and GAPD
APPENDIX B: LESLIE MATRIX SIMULATIONS
APPENDIX B.

LESLIE MATRIX SIMULATIONS OF A DERMOCHELYS CORIACEA POPULATION USING HYPOTHetically-DERIVED PARAMETERS

INTRODUCTION

Knowledge of demographic trends is critical to understanding how well and how quickly stressed populations can be expected to recover given certain management measures. The value of such demographic information was brought to light in Cole’s (1954) landmark paper, but has been overshadowed by the recent use of population genetics as a management tool (Lande, 1988). Few researchers have studied the population dynamics of sea turtles, with the notable exceptions of Bjorndal (1980), Crouse (1985) and Frazer (1983), although most studies contribute in some way to better understanding of life history parameters. Without this information, only crude estimates of population dynamics can be made and the recovery potential of any population cannot be determined.

In order to investigate the way in which life history estimates influence predictions of an endangered population’s recovery potential, I ran several simulation exercises using a Leslie matrix model. Once again, these simulations require some knowledge about the demography of the species; in cases such as this one where the demographic parameters are unknown, they must be estimated using the best available information. The age at first reproduction, total life
span, reproductive life span, frequency of reproduction, and mortality rates for all life stages beyond the egg stage, are not known for *Dermochelys coriacea*. With the exception of fecundity and nest (hatching) success, virtually all the demographic parameters must be hypothesized. Some of these approximations can be inferred from better data that exist for wild green (*Chelonia mydas*) and loggerhead (*Caretta caretta*) sea turtles (e.g. Bjorndal, 1980; Crouse, 1985; Crouse et al., 1987; and Frazer, 1983;) and captive-reared individuals of these two species. However, *Dermochelys coriacea* is markedly different from these other sea turtle species in anatomy (Greer et al., 1973; Rhodin, 1985), physiology (Eckert et al., 1986; Standora et al., 1984), and behavior (Eckert and Eckert, 1988; Lazell, 1980; Pritchard, 1982) so such extrapolations may not be valid.

Despite these qualifications, leatherback turtle population modeling can lead to at least three important findings: 1) an indication of potential population recovery times given management regimes protecting certain age classes, 2) an assessment of the sensitivity of demographic parameters to changes in value, such that the most critical life history stages can be targetted for management to result in the speediest recovery, and 3) an indication of where more data on demographic parameters are most needed.
A Leslie matrix model (Leslie, 1945) was used to simulate demographic trends in a hypothetical leatherback turtle nesting population comprised of one hundred adults, of which 50 are breeding females. Since the life history stages of sea turtles (e.g. eggs, hatchlings, sub-adult, breeding adult) are of unequal duration, the model chosen is one with a stage- rather than age-class structure (Vandermeer, 1975). This stage-class model has also been used in modeling loggerhead turtle populations with interesting results (Crouse et al., 1987). According to the model, the number of individuals in any stage class in the year t+1 is a function of the number of individuals in that year class in the year t such that

\[ Ax(t) = x(t+1) \]

where \( x(t) \) is a vector of stage classes at time t and A is a square matrix of the form:

\[
\begin{bmatrix}
g(0) & g(1) & \ldots & g(m-1) & g(m) \\
a(0) & 0 & \ldots & 0 & 0 \\
0 & a(1) & \ldots & . & . \\
0 & 0 & \ldots & . & . \\
0 & 0 & \ldots & a(m-1) & . \\
\end{bmatrix}
\]

Where \( g(x) \) is the number of offspring in each stage class that survive that period and \( a(gx) \) is the transition probability describing the number of individuals of one stage class that survive to enter the next.

A paucity of data on mortality of leatherbacks of all ages makes it impossible to generate population models of the species without arbitrarily assigning parameter values. The values used in this study
are based on existing information for loggerhead turtles, with
necessary amendments based on what little we do know about leatherback
life histories. The life history parameters used in the model are the
following:

\[ m(x) = \text{number of eggs produced/female in each stage class} \]
\[ p(x) = \text{stage to stage survivorship (1-stage-based mortality rate)} \]
\[ l(x) = \text{egg to stage survivorship} \]
\[ R(0) = \text{finite rate of population increase} \]
\[ G(0) = \text{gross reproductive rate} \]
\[ r = \text{instantaneous rate of population increase} \]

\[ \text{from the population equation } N_t = N_0 e^r \]

The five life history stages used in the model are:

\( x=0 \) egg stage
\( x=1 \) hatchling stage (terrestrial)
\( x=2 \) juvenile stage (marine)
\( x=3 \) newly mature adults
\( x=4 \) remigrants

The only values for which we have definitive information are \( m(x), \)
\( l(x), \) and the duration of the \( x=0 \) life stage; these values come from
my own work with leatherback turtles and from published sources. All
other parameters must be estimated through extrapolation from other
species. One assumption that must be made to ensure that the Leslie
matrix model, which deciphers population trends for females members of
a population only, is realistic is a one to one sex ratio among adults
(Stancyck, 1982). No information about sex ratios of leatherbacks
exists in the literature, however equal sex ratios have been assumed
in investigations of other sea turtle species (Frazer, 1983; Ross,
1982). Other possibly unrealistic assumptions are that the population
is closed (allowing no consideration for emigration or immigration),
and that density dependent mortality effects do not exist.

The Leslie matrix model is a deterministic model which follows
the female population through time to simulate trends in total
population size. The elements of the matrix include age-specific fecundity in Row 1 and age-specific survival probabilities in the subdiagonals. Vectors built from observed (or in this case hypothesized) data allow matrix multiplications which result in the dominant latent root or eigenvalue. If this value is one, the population can be considered at equilibrium; if less than one the population will decrease; and if greater than one the population will increase.

The specific kinds of data that are needed and should be verified by time-series observation are:

1) the number of eggs produced by every stage class;
2) the mortality rates for each stage class from the start to the end of each time period, from which survival estimates are inferred;
3) the total number of size or stage classes within the matrix;
4) adult sex ratio;
5) the number of remigrants and new recruits to nesting for each time interval.

Using hypothesized values for the life history of the leatherback (Ms. 1, Table 1), a life history table can be constructed for use in a Leslie matrix that simulates population sizes over time for any initial vector. A deterministic Leslie matrix model designed for a LOTUS 1-2-3 software program, designed by B. Martin, was used for the manipulations. The first matrix used in the simulation exercise was the following, based on an simplified version of the hypothetical life history table (Table 1):

\[
\begin{array}{cccccc}
0 & 0 & 0 & 220 & 220 \\
.85 & 0 & 0 & 0 & 0 \\
0 & .217 & 0 & 0 & 0 \\
0 & 0 & .03 & 0 & 0 \\
0 & 0 & 0 & .01 & 0 \\
\end{array}
\]
Using a total adult population size of 2000 individuals, of which 1000 are females, an arbitrary initial vector was used to run the simulations.

After 50 years, the matrix manipulation yielded a dominant eigenvalue, $\lambda$, of 0.896, indicating that the population is declining at an annual rate of -0.014.

For the second set of simulations, the matrix was expanded to include five breeding cycles. Stage classes were used as above, but hatchlings were divided into those on land and those at sea, and a juvenile (labeled subadult in the data printouts) stage class was added. In addition, breeding females were divided into newly mature (neophyte breeders), 2nd time breeders or remigrants, 3rd time breeders or remigrants, 4th time breeders or remigrants, and 5th time breeders or remigrants. The model was also expanded to run for a 200, rather than a 50, year simulation.

Using a conservative estimate of survivorship probabilities as follows:

<table>
<thead>
<tr>
<th>stage class</th>
<th>survivorship</th>
<th>fecundity</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>.85</td>
<td>0</td>
</tr>
<tr>
<td>hatchlings (land)</td>
<td>.05</td>
<td>0</td>
</tr>
<tr>
<td>hatchlings (marine)</td>
<td>.10</td>
<td>0</td>
</tr>
<tr>
<td>juvenile</td>
<td>.15</td>
<td>0</td>
</tr>
<tr>
<td>neophyte</td>
<td>.90</td>
<td>220</td>
</tr>
<tr>
<td>2nd time breeder</td>
<td>.80</td>
<td>220</td>
</tr>
<tr>
<td>3rd time breeder</td>
<td>.70</td>
<td>220</td>
</tr>
<tr>
<td>4th time breeder</td>
<td>.60</td>
<td>220</td>
</tr>
<tr>
<td>5th time breeder</td>
<td>.50</td>
<td>220</td>
</tr>
</tbody>
</table>
an eigenvalue approximation of 0.894 resulted. This simulation assumes an equal average fecundity for all breeding females; if the fecundity is reduced so that 4th and 5th time breeders do not produce viable eggs, the eigenvalue approximation drops slightly to 0.845 (see appended data sheets).

A significant change occurs if hatchling or juvenile survivorship is increased even slightly. If, for instance, hatchling survivorship is increased just 10%, from .05 to .15, the population simulation shows an increasing population over 200 years. Even when the egg survivorship is reduced by 25% while the hatchling survivorship is maintained at the .15 value, an eigenvalue approximation of 1.003 indicates an increasing population (see Figure 1).

Various scenarios using different survivorship values and fecundity parameters were run. The most interesting combinations are given in Table 1, with corresponding eigenvalues. The conclusion from these simulations is that while variability in egg survivorship only slightly changes population projections, manipulating the juvenile parameters significantly alters the demographic trends. The neophyte breeder survivorship is also an important parameter. Thus, the most sensitive variables are the sub-adult stage survivorships.

The implications of the matrix simulation are twofold: first, we may do well to rethink our management and conservation efforts to focus more on the most sensitive age classes (juveniles and newly mature adults); and second, that further basic research is needed to better define demographic parameters. Until we know, or at least have observation-based estimates of, stage class mortalities and growth
### Figure 1. Additional projections from hypothetical Leslie matrix simulations, using various survivorship values

#### Case 1: Equal fecundity all breeders

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>0.854</td>
<td>0.854</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.85</td>
<td>0</td>
<td>0.85</td>
</tr>
<tr>
<td>hatching</td>
<td>0.05</td>
<td>0</td>
<td>0.0425</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.00425</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.000637</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.0008733</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.80</td>
<td>220</td>
<td>0.0009459</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.70</td>
<td>220</td>
<td>0.000321</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.60</td>
<td>220</td>
<td>0.001952</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000096</td>
</tr>
</tbody>
</table>

#### Case 2: 3 Nesting cycles only

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>0.845</td>
<td>0.845</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.85</td>
<td>0</td>
<td>0.85</td>
</tr>
<tr>
<td>hatching</td>
<td>0.05</td>
<td>0</td>
<td>0.0425</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.00425</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.000637</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.0008733</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.80</td>
<td>220</td>
<td>0.0009459</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.70</td>
<td>220</td>
<td>0.000321</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.60</td>
<td>220</td>
<td>0.001952</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000096</td>
</tr>
</tbody>
</table>

#### Case 3: Reduced adult survivorship

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>0.845</td>
<td>0.845</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.85</td>
<td>0</td>
<td>0.85</td>
</tr>
<tr>
<td>hatching</td>
<td>0.05</td>
<td>0</td>
<td>0.0425</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.00425</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.000637</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.0008733</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.80</td>
<td>220</td>
<td>0.0009459</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.70</td>
<td>220</td>
<td>0.000321</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.60</td>
<td>220</td>
<td>0.001952</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000096</td>
</tr>
</tbody>
</table>

#### Case 4: Increase beach survivors to 15%

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>1.020</td>
<td>1.020</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.85</td>
<td>0</td>
<td>0.85</td>
</tr>
<tr>
<td>hatching</td>
<td>0.15</td>
<td>0</td>
<td>0.1275</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.01275</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.001912</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.0001721</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000686</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000430</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.0002153</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000107</td>
</tr>
</tbody>
</table>

#### Case 5: Low nest success but help hatchlings

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>1.005</td>
<td>1.003</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.60</td>
<td>0</td>
<td>0.6</td>
</tr>
<tr>
<td>hatching</td>
<td>0.15</td>
<td>0</td>
<td>0.099</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.00135</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.000637</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.001215</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.80</td>
<td>220</td>
<td>0.000972</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.70</td>
<td>220</td>
<td>0.000490</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.60</td>
<td>220</td>
<td>0.000406</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000204</td>
</tr>
</tbody>
</table>

#### Case 6: Low adult survival, low nest success, but help hatchlings

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>0.964</td>
<td>0.964</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.85</td>
<td>0</td>
<td>0.85</td>
</tr>
<tr>
<td>hatching</td>
<td>0.05</td>
<td>0</td>
<td>0.0425</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.00425</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.000637</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.0008733</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.80</td>
<td>220</td>
<td>0.0009459</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.70</td>
<td>220</td>
<td>0.000321</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.60</td>
<td>220</td>
<td>0.001952</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000096</td>
</tr>
</tbody>
</table>

#### Case 7: Survivorship: high eggs, low hatchlings, very high adult

<table>
<thead>
<tr>
<th>Repro pop</th>
<th>ult/penult</th>
<th>penult/penult</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lambda=</td>
<td>0.948</td>
<td>0.948</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>age class</th>
<th>female surv.</th>
<th>fecundity</th>
<th>year=</th>
</tr>
</thead>
<tbody>
<tr>
<td>eggs</td>
<td>0.85</td>
<td>0</td>
<td>0.85</td>
</tr>
<tr>
<td>hatching</td>
<td>0.05</td>
<td>0</td>
<td>0.0425</td>
</tr>
<tr>
<td>swimmers</td>
<td>0.10</td>
<td>0</td>
<td>0.00425</td>
</tr>
<tr>
<td>subadult</td>
<td>0.15</td>
<td>0</td>
<td>0.000637</td>
</tr>
<tr>
<td>neophyte</td>
<td>0.90</td>
<td>220</td>
<td>0.0008733</td>
</tr>
<tr>
<td>2nd breeder</td>
<td>0.80</td>
<td>220</td>
<td>0.0009459</td>
</tr>
<tr>
<td>3rd breeder</td>
<td>0.70</td>
<td>220</td>
<td>0.000321</td>
</tr>
<tr>
<td>4th breeder</td>
<td>0.60</td>
<td>220</td>
<td>0.001952</td>
</tr>
<tr>
<td>5th breeder</td>
<td>0.50</td>
<td>220</td>
<td>0.000096</td>
</tr>
</tbody>
</table>

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rates, we cannot begin to have an accurate understanding of leatherback turtle population dynamics. And without this understanding, sea turtle management will continue to capitalize on opportunity rather than on what is most effective.

ACKNOWLEDGMENTS

Brooks Martin of U.R.I. developed the LOTUS 1-2-3 Leslie matrix program & was invaluable in assisting with the simulations.

LITERATURE CITED

(See dissertation bibliography)
BIBLIOGRAPHY


