Insights on Immune Function in Free-Ranging Green Sea Turtles (*Chelonia mydas*) with and without Fibropapillomatosis

Justin R. Perrault1,*, Milton Levin2, Cody R. Mott3, Caitlin M. Boveri4, Michael J. Bresette3, Ryan M. Chabot3,4, Christopher R. Gregory5, Jeffrey R. Guertin3*, Sarah E. Hirsch1,*, Branson W. Ritchie5, Steven T. Weege3, Ryan C. Welsh3, Blair E. Witherington3 and Annie Page-Karjian6

1 Loggerhead Marinelife Center, Juno Beach, FL 33408, USA; shirsch@marinelife.org
2 Department of Pathobiology and Veterinary Science, University of Connecticut, Storrs, CT 06269, USA; milton.levin@uconn.edu
3 Inwater Research Group, Jensen Beach, FL 34957, USA; cmott@inwater.org (C.R.M.); mbresette@inwater.org (M.J.B.); rchabot@inwater.org (R.M.C.); jguertin@inwater.org (J.R.G.); sweege@inwater.org (S.T.W.); rwelsh@inwater.org (R.C.W.); bwitherington@inwater.org (B.E.W.)
4 Gumbo Limbo Nature Center, Boca Raton, FL 33432, USA; cbovery@ci.boca-raton.fl.us
5 Infectious Diseases Laboratory, University of Georgia, Athens, GA 30602, USA; crg@uga.edu (C.R.G.); britchie@uga.edu (B.W.R.)
6 Harbor Branch Oceanographic Institute, Florida Atlantic University, Fort Pierce, FL 34946, USA; cpagekarjian@fau.edu
* Correspondence: jprerrault@marinelife.org
† Present address: Volusia County Environmental Management, DeLand, FL 32770, USA.

Simple Summary: Sea turtles are susceptible to several herpesviruses that are linked to dermatologic diseases, including fibropapillomatosis (FP) and lung-eye-trachea disease. Aside from obvious skin lesions, a number of other sublethal impacts occur in response to these diseases, such as reduced immune function. In this study, we found no relationship between disease presence or severity and T-cell proliferation in green turtles from Florida, USA, at least until the moderate stages of FP; however, natural killer cell activity, a measure of innate immune function, was significantly reduced in turtles with FP compared to tumor-free individuals. This is the first study to examine natural killer cell activity in relation to FP, improving upon our understanding of altered immune system function associated with this disease.

Abstract: Chelonid alphaherpesviruses 5 and 6 (ChHV5 and ChHV6) are viruses that affect wild sea turtle populations. ChHV5 is associated with the neoplastic disease fibropapillomatosis (FP), which affects green turtles (*Chelonia mydas*) in panzootic proportions. ChHV6 infection is associated with lung-eye-trachea disease (LETD), which has only been observed in maricultured sea turtles, although antibodies to ChHV6 have been detected in free-ranging turtles. To better understand herpesvirus prevalence and host immunity in various green turtle foraging aggregations in Florida, USA, our objectives were to compare measures of innate and adaptive immune function in relation to (1) FP tumor presence and severity, and (2) ChHV5 and ChHV6 infection status. Free-ranging, juvenile green turtles (*N* = 45) were captured and examined for external FP tumors in Florida’s Big Bend, Indian River Lagoon, and Lake Worth Lagoon. Blood samples were collected upon capture and analyzed for ChHV5 and ChHV6 DNA, antibodies to ChHV5 and ChHV6, in vitro lymphocyte proliferation using a T-cell mitogen (concanavalin A), and natural killer cell activity. Despite an overall high FP prevalence (56%), ChHV5 DNA was only observed in one individual, whereas 20% of turtles tested positive for antibodies to ChHV5. ChHV6 DNA was not observed in any animals and only one turtle tested positive for ChHV6 antibodies. T-cell proliferation was not significantly related to FP presence, tumor burden, or ChHV5 seroprevalence; however, lymphocyte proliferation in response to concanavalin A was decreased in turtles with severe FP (*N* = 3). Lastly, green turtles with FP (*N* = 9) had significantly lower natural killer cell activity compared to FP-free turtles (*N* = 5). These results increase our understanding of immune system effects related to FP and provide evidence that immunosuppression occurs after the onset of FP disease.
Keywords: ChHV5; ChHV6; ELISA; herpesvirus; lung-eye-trachea disease; lymphocyte proliferation; marine turtle; mitogen; natural killer cell; seroprevalence

1. Introduction

Green sea turtles (Chelonia mydas) are considered indicators of marine ecosystem health [1–4]. They play important roles in seagrass bed resiliency by preventing overgrowth and reducing nutrient input into these areas [5]. Green turtle populations in the northwestern Atlantic Ocean suffered substantial population declines prior to the 20th century due to overexploitation of eggs and adults for human consumption, and are subject to modern threats including entanglement in fishing gear, habitat loss, beach-front lighting, boat strikes, and pollution [3,6–8]. Despite their current threatened status, green turtle populations in Florida, USA appear to be recovering due to conservation efforts, evidenced by increasing nest counts, hatching production, and captures of immature individuals in foraging areas [3,7,9–12]. Although these population trends are encouraging, immature green turtles in Florida have a high prevalence (>50%) of fibropapillomatosis (FP) [13–18], a transmissible tumor disease associated with chelonid alphaherpesvirus 5 (ChHV5) infection [19]. Extensive research has been conducted in an attempt to better understand FP, yet etiology, pathogenesis, and transmission are still not well understood. Turtles inhabiting nearshore, shallow-water embayments tend to express the highest FP prevalence. Environmental cofactors including agricultural runoff, exposure to biotoxins and toxicants, habitat degradation, warm water temperatures, and UV radiation have been suggested to play a role in FP pathogenesis and maintenance within green turtle populations [17,19–27]. To date, however, no studies have identified confirmed causal relationships between environmental cofactors and FP occurrence.

Since its discovery in the Florida Keys in 1937 [28] and in Hawaii in 1958 [29], FP has become a panzootic in green turtles with observations in all seven sea turtle species in numerous populations around the globe [30–37]. ChHV5 is likely horizontally transmitted via shedding of infected epithelial cells, and possibly in urine and/or bodily secretions [21,38–42]. When FP tumors are present, depending on tumor size, extent, and anatomic location, they can cause physical alterations that lead to reductions in afflicted turtles’ ability to see, breathe, swim, dive, forage, and avoid predators [43]. ChHV5 nucleic acids can be detected in biological samples using polymerase chain reaction (PCR). Previous exposure to, and presumptive infection with ChHV5 can be identified via detection of circulating antibodies in sea turtle blood samples using enzyme-linked immunosorbent assays (ELISA) [4,44]. Changes in blood-based indicators of health have been observed in turtles with FP, including anemia, hypoproteinemia, leukopenia/immunosuppression, and other biochemical alterations [15,17,29,45–51]. More specific associations of FP with afflicted turtles’ immune function include decreased T- and B-cell function, suggesting impacts to cell-mediated and humoral immunity (i.e., adaptive immunity). It remains unknown, however, whether these immunomodulatory alterations are a cause of or sequel to the onset of FP disease [27,36,46,47].

Another disease-causing herpesvirus that affects sea turtles is chelonid alphaherpesvirus 6 (ChHV6) [52], formerly known as lung-eye-trachea virus (LETV), which was first recognized in 1975 in 15–20-month-old, maricultured green turtles residing at the Cayman Turtle Farm (Grand Cayman, British West Indies). Clinical signs included respiratory and buoyancy abnormalities with caseous exudate surrounding the eyes, glottis, and trachea, and a mortality rate of up to 38% [53]. Since its discovery, lesions associated with ChHV6 infection have not been reported in free-ranging turtles; however, antibodies to ChHV6 have been detected in juvenile green turtles from Florida with seroprevalence rates of 10–22%, depending on location [54,55]. Nesting loggerheads (Caretta caretta; 75% seroprevalence: 3/4 from Melbourne Beach, Florida) and green turtles (100% seroprevalence: 9/9 from Melbourne Beach, Florida; 15% seroprevalence: 6/41 from Juno Beach,
Florida) have also tested positive for ChHV6 antibodies via ELISA [4,55], but negative for circulating ChHV6 DNA via PCR [4]. Despite exposure to ChHV6, abnormalities in blood health analytes have not been observed in turtles with ChHV6-specific antibodies [4]. To build upon previous studies associating sea turtle host immunity to gross and molecular diagnostic measures of ChHV5 and ChHV6 infection, the objectives of this study were to compare innate and adaptive immune function in relation to (1) FP tumor presence and severity, and (2) ChHV5 and ChHV6 infection status in free-ranging, juvenile green turtles from three foraging aggregations across Florida.

2. Materials and Methods
2.1. Sampling Sites

2.1.1. Big Bend Region, Florida
The coastal waters of Florida’s Big Bend region (Figure 1; hereafter referred to as Big Bend) extend from the eastern boundary of the Apalachicola–Chattahoochee–Flint River Watershed to Anclote Key in Pinellas County. This area serves as an important foraging habitat for juvenile green and Kemp’s ridley (Lepidochelys kempii) sea turtles as it contains the second largest expanse of seagrass habitat in the eastern Gulf of Mexico [18,56,57], with primarily natural land coverage, conservation protection of much of the coastal and inland areas, low human population density, and minimal anthropogenic threats (e.g., boating, eutrophication, and pollution) [58,59]. Despite this seemingly pristine environment, loss of oyster beds and seagrasses and a high FP prevalence rate (60–70%) in green turtles occur in this area [18,57,59].

![Figure 1. Locations of green turtle (Chelonia mydas) sampling sites across Florida, USA: Big Bend Region, Jennings Cove in the Indian River Lagoon, and the Jupiter Inlet in the northern portion of the Lake Worth Lagoon.](image)

2.1.2. Jennings Cove, Indian River Lagoon, Florida
The Indian River Lagoon (IRL) spans 250 km of Florida’s east coast, was described as one of the most biodiverse estuaries in North America, and has been designated as...
an Estuary of National Significance [60,61]. This system has been impacted by human population growth, species loss, nutrient runoff, pollutants, and harmful algal blooms. Poor water quality and declines in ecological and biological integrity have occurred in the area due primarily to nutrient-rich runoff and habitat loss [62–64]. More than 50 point-source waste sites are permitted to discharge pollutants into the IRL including emissions from power plants, brine discharges, septic tanks, and water treatment plants [61]. The prevalence of FP in green turtles captured at Jennings Cove in the southern portion of the IRL (Figure 1) has varied with no detectable trend over the last 10–15 years, with annual prevalence rates ranging from 33 to 90% (mean ± standard error of 68 ± 6%) [65].

2.1.3. Jupiter Inlet, Lake Worth Lagoon, Florida

The Jupiter Inlet (Figure 1) serves as the northernmost boundary of the Lake Worth Lagoon, which is a heavily urbanized waterway spanning ~32 km in Palm Beach County, Florida. Research on the Jupiter Inlet is lacking; however, the primary environmental issues in the Lake Worth Lagoon system are related to untreated stormwater runoff and septic tank overflow, which can drain into the lagoon, subsequently leading to reductions in water clarity, salinity fluctuations, and excess nutrient deposition [66,67]. Over the last 10–15 years, underground storm water treatment areas have been installed to filter water before it reaches Lake Worth Lagoon. This is part of an overall restoration plan that includes mitigation measures to enhance seagrass, oyster beds, and mangrove habitats in the lagoon. Juvenile green turtles are abundant in the more southern areas of Lake Worth Lagoon and exhibit a high prevalence (~50%) and severity of FP [16], but little is known about FP rates in turtles from the more northern areas of the lagoon (i.e., Jupiter Inlet).

2.2. Animal Capture and Sample Collection and Processing

Green turtles were captured using dip nets, tangle nets, and/or the rodeo method, depending on habitat and environmental conditions [68]. Turtles from Big Bend, Jennings Cove, and Jupiter Inlet were sampled from 27 to 30 August 2020, 20 September 2018 to 8 October 2020, and 17 to 18 September 2020, respectively. Captured turtles were brought onboard the vessel for data collection. Prior to sample collection, turtles were externally examined [69,70], weighed and measured (standard straight carapace length, SCL), assessed for subjective body condition (i.e., emaciated, thin, good, and robust), and Fulton’s body condition index (BCI) was calculated [6]. A complete external visual exam was conducted on all turtles. This included observations for the presence or absence of leeches (Ozobranchus spp.), leech cocoons, barnacles, flipper and carapace damage, and visual assessment and enumeration of FP tumors (i.e., total tumor number). Flipper and carapace damage were defined as missing >5% of the flipper or carapace and/or an injury caused by predation, anthropogenic impact, or abnormal development. Tumors were measured and recorded on a standardized tumor score sheet. Each turtle was assigned a tumor severity category based on tumor burden (0: no tumors; 1: mildly afflicted; 2: moderately afflicted; 3: severely afflicted) [29]. Blood (~10–20 mL; <1% of body weight) was sampled from the external jugular vein using sodium heparin and serum separator Vacutainer® (Becton, Dickson, and Co., Franklin Lakes, NJ, USA) blood collection systems fitted with 21–25-gauge, 1 inch needles, as appropriate based on turtle size. The venipuncture site was swabbed with alternating applications of povidone iodine and 70% isopropyl alcohol prior to and after blood collection. Blood samples were immediately chilled on ice in the field until return to the laboratory (<8 h). An aliquot (~100 µL) of well-mixed whole blood was placed into a separate cryovial and stored in an ultralow freezer (−80 °C) for up to six months prior to DNA extraction for PCR analysis. The remaining whole blood in the sodium heparin vacutainers was shipped overnight on ice packs to the University of Connecticut. Serum from the serum separator tubes was harvested after centrifugation at 1318 × g (3400 rpm) for 10 min and stored in an ultralow freezer for up to 6 months prior to serological analysis.
All turtles were tagged with metal flipper (Inconel®) and passive integrated transponder (PIT) tags for identification. Tagging sites were swabbed with povidone iodine and 70% isopropyl alcohol prior to and after tag application. After completion of sampling and tagging, turtles were released back into the water at or near the site of capture.

2.3. Molecular Diagnostics for ChHV5 and ChHV6 DNA

Quantitative PCR (qPCR) for ChHV5 DNA was performed at Florida Atlantic University’s Harbor Branch Oceanographic Institute in Fort Pierce, Florida. First, genomic DNA (gDNA) was extracted from thawed whole blood samples using the DNeasy Blood and Tissue kit according to the manufacturer’s instructions (Qiagen, Hilden, Germany). Concentrations of extracted gDNA samples were quantified using a NanoDrop 2000c (Thermo Fisher Scientific, Waltham, MA, USA) (units: µg/µL) spectrophotometer, and the ratio of absorbance at 260 and 280 nm was used to assess gDNA purity. Extracted gDNA samples were assessed for the presence of a ChHV5 UL30 gene segment using a singleplex, hydrolysis probe-based qPCR and the methodologies described in detail by Page-Karjian et al. [40]. Quantitative PCR reactions were conducted using an AriaMx Real-Time PCR System (Agilent, Santa Clara, CA, USA), and qPCR data were analyzed with AriaMx software (Agilent, Version 1.3). All ChHV5-positive qPCR products were purified using the QIAquick PCR Purification Kit (Qiagen) and subjected to Sanger sequencing (Genewiz) using 5 µM of ChHV5 UL30 forward primer. Sequences obtained were compared to those deposited in the National Center for Biotechnology Information GenBank database using BLAST software [71]. Aligned sequences with ≥97% identity to the sample sequence were considered a match.

Aliquots of frozen whole blood were shipped overnight on dry ice to University of Georgia’s Infectious Diseases Laboratory (UGA IDL; Athens, GA, USA). Genomic DNA was extracted from these samples using the methodology outlined above. Extracted gDNA samples were analyzed using a singleplex, hydrolysis probe-based qPCR assay that targets a unique 112 bp ChHV6 DNA amplicon, using standard operating procedures and controls for the assay. Detailed methodology is provided in Page-Karjian et al. [4].

2.4. ELISA Detection of Antibodies to ChHV5 and ChHV6 Peptides

Separated serum samples were analyzed for antibodies to ChHV5 and ChHV6 peptides at the UGA IDL. To evaluate for infection by ChHV5 and ChHV6 in a turtle’s immunologically detectable past, samples were analyzed in triplicate using ELISAs that test for antibodies to ChHV5 and ChHV6 purified synthetic peptide antigens. Both ELISA assays were developed and validated based on modifications of previously published protocols for ChHV5 and ChHV6 assays [54,72] and were performed using the laboratory’s standard operating procedures with negative and positive control sera. Detailed methodology is provided in Page-Karjian et al. [4].

2.5. Isolation of Peripheral Blood Mononuclear Cells

At the University of Connecticut (Storrs, CT, USA), whole blood from the sodium heparin tubes was mixed 1:1 with an equal volume of Hank’s Balanced Salt Solution (HBSS; Thermo Fisher Scientific). Peripheral blood mononuclear cells (PBMCs) were isolated by density gradient centrifugation using Ficoll-Paque plus (Amersham Biosciences, Uppsala, Sweden; 1.077 g/mL) for 30 min at 600 × g. The interface containing the PBMCs was re-suspended in complete RPMI and washed twice in complete RPMI. PBMCs were enumerated and their viability was assessed using the exclusion dye trypan blue and light microscopy. Complete RPMI consisted of RPMI 1640 (with 2 mM L-glutamine; Thermo Fisher Scientific) supplemented with 1 mM sodium pyruvate, 100 mM nonessential amino acids, 10 mM HEPES, 50 U/mL penicillin, 50 µg/mL streptomycin, 0.25 µg/mL fungizone, and 10% fetal bovine serum (all from Thermo Fisher Scientific).
2.6. Lymphocyte Proliferation

Mitogen-induced lymphocyte proliferation was evaluated in vitro as previously described [73], with some modifications. Briefly, PBMCs in complete RPMI were plated (2 × 10^5 cells/well) in a 96-well flat-bottom plate (Falcon; Becton, Dickinson, and Co.) in triplicate for each mitogen concentration. Cells were maintained for 96 h in an incubator at 28 °C and humidified atmosphere with 5% CO₂. The T-cell mitogen concanavalin A (ConA, C5275, Millipore Sigma, St. Louis, MO, USA) at 1 (suboptimal) and 10 µg/mL (optimal) was used. Lymphocyte proliferation was evaluated as the incorporation of 5-bromo-20-deoxyuridine (BrdU), a thymidine analogue, added for the last 18 h of incubation, and further detected with a monoclonal antibody and a colorimetric enzymatic reaction (Cell Proliferation colorimetric ELISA BrdU, RocheDiagnostics GmbH, Mannheim, Germany) as per manufacturer’s instructions using an ELISA plate reader (Multiskan EX v.1.0) at 450 nm with a reference wavelength of 690 nm. Data are reported as:

\[
\text{Stimulation index} = \left( \frac{\text{Mitogen optical density}}{\text{Unstimulated optical density}} \right) \times 100
\]

Results with cell viability percentages <80% were excluded from statistical analyses.

2.7. Natural Killer Cell Activity

For green turtles captured in Big Bend, natural killer cell activity was evaluated in vitro as previously described [73,74] with some modifications. Briefly, 1 mL of YAC-1 (TIB-160™, ATCC, Manassas, VA, USA) target cells was incubated with 10 µL of 3 mM 3,3′-dioctadecyloxacyanine perchlorate (DiO, Molecular Probes, Grand Island, NY, USA) dissolved in DMSO, and incubated for 20 min at 37 °C in 5% CO₂, followed by two washes in complete RPMI. The target cells were then re-suspended in complete RPMI. PBMCs (effector cells) were adjusted to 1 × 10^6 cells/mL and target cells were added to achieve effector:target (E:T) ratio of 50:1 [74]. The E:T mixtures were centrifuged for 30 s at 220 × g and further incubated for 150 min at 28 °C in 5% CO₂. All tests were performed in duplicate.

Following centrifugation at 220 × g for 10 min at 4 °C, the supernatant was discarded and the cells were re-suspended in 200 µL of phosphate buffered saline (PBS, Thermo Fisher Scientific) and placed on wet ice before immediate analysis. Cells were re-suspended in a solution of 50 µg/mL of propidium iodide (PI; Thermo Fisher Scientific) to evaluate mortality of the target cells immediately prior to acquisition using two-color (DiO vs. PI) flow cytometry. The fluorescence of at least 1000 target cells was read using a FACScan flow cytometer (Becton, Dickinson, and Co.) and the automated CellQuest software (Becton Dickinson Immunocytometry System, San Jose, CA, USA). Effector cells were identified by their relative size (forward-scattered light) and their complexity (side-scattered light) and distinguished from DiO-labeled target cells, which show higher fluorescence at 530 nm (FL-1). Dead or dying cells incorporate PI due to membrane instability and show higher fluorescence at 630 nm (FL-3). Results were calculated as:

\[
\text{Percent target cell mortality} = \left( \frac{\# \text{ dead target cells}}{\# \text{ dead target cells} + \# \text{ live target cells}} \right) \times 100
\]

The percent of spontaneous target cell mortality was then subtracted from the percent target cell mortality to calculate specific target cell mortality.

2.8. Statistical Analyses

Statistical analyses were conducted using SPSS (v27, SPSS Inc., Chicago, IL, USA). One-way analysis of variance (ANOVA) was used to compare turtle body mass, SCL, and BCI by sampling site and tumor score using transformations as necessary to meet test assumptions, followed by Tukey’s post hoc tests with Bonferroni correction. The relationship between log-transformed SCL and body mass was assessed using linear regression. For binary
data (e.g., presence/absence of FP, leeches, barnacles, injuries, and presence/absence of ChHV5 DNA or antibodies), chi-square analysis or Fisher’s exact tests (when \( N \leq 5 \)) were employed using Bonferroni correction for multiple comparisons. Independent samples t-tests were used to compare (1) calculated BCI between two subjective body condition scores (good versus robust) and (2) natural killer cell activity between turtles with and without FP. Cohen’s Kappa (\( \kappa \)) coefficient was calculated to determine the qualitative level of agreement between FP presence/absence and qPCR and ELISA results for ChHV5, and to compare qPCR to ELISA results for ChHV5 [75]. Logistic regression was used to determine if turtle size (i.e., SCL) was related to seroprevalence (i.e., positive and negative). Due to a lack of normality in lymphocyte proliferation assay data, Kruskal–Wallis tests with post hoc Dunn’s tests were used to calculate differences in lymphocyte proliferation by sampling location. Because there were no differences in lymphocyte proliferation by study site for non-tumored turtles and for all turtles combined, all sites were combined for comparisons of lymphocyte proliferation activity with regards to the presence/absence of FP, FP tumor severity, and ChHV5 seroprevalence.

3. Results
3.1. Physical Examination and Morphometrics

A total of 45 green turtles were captured from three study sites including Big Bend (\( N = 16 \)), Jennings Cove (\( N = 18 \)), and Jupiter Inlet (\( N = 11 \)). All turtles were considered to be immature based on SCL measurements (range: 26.0–70.1 cm) [7,76]. Straight carapace length (\( F(2, 42) = 13.840; p < 0.001 \)) and body mass (\( F(2,40) = 13.623; p < 0.001 \)) differed by capture site, as turtles captured at Jennings Cove were significantly larger and heavier than turtles captured at Big Bend (\( p < 0.001 \)) and the Jupiter Inlet (\( p < 0.001 \)). Log-transformed body mass and SCL were very strongly correlated (\( y = 3.07x - 3.99; r^2 = 0.99; p < 0.001 \)). Neither SCL nor body mass differed by tumor score category (\( p > 0.05 \) in both cases). Calculated BCI did not differ by capture site (\( p > 0.05 \)), but BCI was significantly higher (\( t(42) = 2.243; p = 0.030 \)) in turtles assigned a subjective body condition score of “robust” (mean \( \pm \) standard deviation [SD] = 1.34 \( \pm \) 0.11) compared to those given a score of “good” (mean \( \pm \) SD = 1.27 \( \pm \) 0.10). All captured turtles were considered to be of “good” (\( N = 30 \)) or “robust” (\( N = 15 \)) body condition, with no significant differences by site after Bonferroni correction. No sampled animals were score as “thin” or “emaciated” based on subjective observation.

Ozobranchus leeches, leech cocoons, and barnacles (Chelonibia spp.) were present on the skin and/or carapace of 22% (10/45), 20% (9/45), and 64% (29/45) of turtles, respectively. Leech (\( X^2(2) = 13.705; p = 0.001 \)) and leech cocoon (\( X^2(2) = 11.540; p = 0.003 \)) prevalence differed by site, wherein Jennings Cove turtles had a higher prevalence of leeches (\( p = 0.001 \)) and leech cocoons (\( p = 0.003 \)) in comparison to turtles from Big Bend. No differences in leech prevalence were observed between turtles from Jupiter Inlet and turtles from Big Bend or Jennings Cove (\( p > 0.05 \) in both cases).

Flipper damage was observed on 16% (7/45) of turtles, including small and large notches of unidentified cause (\( N = 4 \)), as well as partial (\( N = 2 \)) and complete amputations (\( N = 1 \)) of at least one limb. Carapace damage was observed on 11% (5/45) of turtles and included a small notch of unknown origin to the left posterior carapace (\( N = 1 \)), a possible healed shark predation injury (\( N = 1 \)), and healed caudal vessel strikes (\( N = 3 \)). Prevalence of flipper and carapace damage did not differ by site or between turtles with and without FP (\( p > 0.05 \) in both cases). A complete description of morphometrics, BCI, epibiota prevalence, and flipper and carapace damage prevalence is reported in Table 1.
Table 1. Morphometrics and results of external physical exam for green turtles (*Chelonia mydas*) from three locations in Florida, USA. For standard straight carapace length (SCL), body mass, body condition index (BCI), tumor score, and tumor number, mean ± standard deviation are given with the median and range in parentheses. Sample sizes differed in some categories as not all analyses were conducted on every sample; this is indicated parenthetically where relevant. Different superscript letters next to each category represent significant differences between sites, when present. Abbreviations: ChHV5, chelonid alphaherpesvirus 5; ChHV6, chelonid alphaherpesvirus 6; qPCR, quantitative polymerase chain reaction.

| Physical Examination | Big Bend | Jennings Cove | Jupiter Inlet | Total |
|----------------------|----------|---------------|--------------|-------|
| N                    | 16       | 18            | 11           | 45    |
| Sampling dates       | Aug 2020 | Sep 2018–Oct 2020 | Sep 2020 | Sep 2019–Oct 2020 |
| SCL (cm)             | 34.4 ± 4.6 A (33.8) (27.9–43.4) | 48.2 ± 10.3 B (49.8) (32.5–70.6) | 35.4 ± 9.1 A (32.1) (27.2–53.0) | 40.2 ± 10.6 (36.9) (27.2–70.6) |
| Body mass (kg)       | 5.6 ± 2.7 A (5.2) (2.8–12.3) | 16.8 ± 11.6 B (14.4) (4.5–46.8) | 6.7 ± 6.4 A (3.8) (2.8–19.5) | 10.2 ± 9.4 (6.6) (2.8–46.8) |
| Body condition index | 1.29 ± 0.11 (1.10–1.50) | 1.32 ± 0.12 (1.31) (1.07–1.52) | 1.26 ± 0.09 (1.27) (1.10–1.39) | 1.29 ± 0.11 (1.31) (1.07–1.52) |
| Subjective body condition | Good: 44% (7/16) (1.31) | Robust: 56% (9/16) (1.31) | Good: 72% (13/18) (1.31) (1.07–1.52) | Good: 91% (10/11) (1.31) (1.07–1.52) |
| Leech prevalence     | 0% (0/16) A (0–7) | 50% (9/18) B (0–7) | 9% (1/11) AB (0–7) | 22% (10/45) (0–7) |
| Leech cocoon prevalence | 0% (0/16) A (0–7) | 44% (8/18) B (0–7) | 9% (1/11) AB (0–7) | 20% (9/45) (0–7) |
| Barnacle prevalence  | 63% (10/16) (0–7) | 61% (11/18) (0–7) | 73% (8/11) (0–7) | 64% (29/45) (0–7) |
| Flipper damage       | 6% (1/16) (0–7) | 22% (4/18) (0–7) | 18% (2/11) (0–7) | 16% (7/45) (0–7) |
| Carapace damage      | 19% (3/16) (0–7) | 11% (2/18) (0–7) | 0% (0/11) (0–7) | 11% (5/45) (0–7) |
| FP and diagnostic assays | Big Bend | Jennings Cove | Jupiter Inlet | Total |
| FP prevalence        | 69% (11/16) (1.0 ± 0.8) (1–0) | 56% (10/18) (1.3 ± 1.2) (2–0) | 36% (4/11) (0.5 ± 0.8) (0–2) | 56% (25/45) (1.0 ± 1.0) (1–0) |
| Tumor number         | 10 ± 15 (0–44) (0–79) | 21 ± 25 (0–44) (0–79) | 4 ± 8 (0–26) (0–79) | 13 ± 20 (1–0) (2–0) |
| ChHV5 qPCR+          | 6% (1/16) (0–7) | 0% (0/13) (0–7) | 0% (0/11) (0–7) | 3% (1/40) (0–7) |
| ChHV5 seropositive   | 13% (2/15) (0–7) | 46% (6/16) (0–7) | 0% (0/9) (0–7) | 20% (8/40) (0–7) |
| ChHV6 qPCR+          | 0% (0/16) (0–7) | 0% (0/13) (0–7) | 0% (0/11) (0–7) | 0% (0/40) (0–7) |
| ChHV6 seropositive   | 0% (0/15) (0–7) | 8% (1/13) (0–7) | 0% (0/9) (0–7) | 3% (1/37) (0–7) |

3.2. Fibropapilloma Tumor Score and Severity

The overall prevalence of FP tumors across all three study sites was 56% (25/45), with the highest prevalence observed in turtles from Big Bend (69%; 11/16), followed by Jennings Cove (56%, 10/18), and then Jupiter Inlet (36%, 4/11); however, these differences were not statistically significant ($p > 0.05$). Balazs–Work tumor scores ranged from 0 to 3 across all sites; median tumor scores were 0 (Jupiter Inlet), 1 (Big Bend), and 2 (Jennings Cove). These results failed to achieve statistical significance, but it is worth noting that only Jennings Cove had turtles in the tumor score 3 category ($N = 3$) and also had no turtles of tumor score 1. Lastly, total tumor number per turtle ranged from 0 to 70; turtles from Jennings Cove had the most tumors (median = 13), followed by Big Bend (median = 1), and then Jupiter Inlet (median = 0). Again, these results did not significantly differ by site ($p > 0.05$).

Tumor score, severity, and total tumor number did not differ among turtles of different body sizes ($p > 0.05$), likely due to low sample sizes. Turtles in the 20–29.9 cm size range ($N = 7$) had the lowest FP prevalence, severity, and tumor number, which increased in turtles of 30–39.9 cm ($N = 21$) and 40–49.9 cm ($N = 6$), then decreased in turtles in the 50–59.9 cm size range ($N = 8$). Turtles in the >60 cm size range ($N = 3$) showed no external evidence of tumors. A complete description of FP prevalence, severity, and tumor number...
is reported in Table 1. Neither BCI nor subjective body condition significantly differed between turtles with and without FP ($p > 0.05$ in both cases). More specifically, BCI did not differ by Balazs–Work tumor score category ($N = 19$ for tumor score 0; $N = 8$ for tumor score 1; $N = 14$ for tumor score 2; $N = 3$ for tumor score 3; $p > 0.05$) (Figure 2). Prevalence of leeches, cocoons, and barnacles did not differ between turtles with and without FP ($p > 0.05$).

### 3.3. Molecular Detection of ChHV5 and ChHV6 DNA and Antibodies

Blood samples from $1/40$ (3%) and $0/40$ (0%) turtles tested positive for ChHV5 and ChHV6 DNA via qPCR, respectively. The amplified DNA sequence from the single sample (a turtle from Big Bend with a tumor score of 2) that tested positive for ChHV5 DNA was subjected to Sanger sequencing, and the resulting trace file matched to the ChHV5 partial genome (GenBank accession number HQ878327.2) with $\geq 99\%$ identity. This sample had 130 viral copies/µg DNA.

In total, $8/40$ (20%) of turtles tested positive for antibodies to ChHV5, with seroprevalence highest in turtles from Jennings Cove (46%, 6/16), followed by Big Bend (13%, 2/15), and Jupiter Inlet (0%, 0/9). These results were not statistically significant after Bonferroni correction. The single turtle that tested positive for ChHV5 DNA by qPCR tested negative for antibodies to the ChHV5 peptide. One turtle from Jennings Cove (3%; 1/37) tested positive for antibodies to ChHV6; this individual had no external tumors, but also tested positive for antibodies to ChHV5. Fibropapilloma status (i.e., tumored or nontumor) and FP tumor score were not significantly related to ChHV5 antibody prevalence ($p > 0.05$); however, the seroprevalence rate was 43% (6/14) in turtles with tumor scores 2 and 3, compared to 13% (3/21) seroprevalence in turtles with tumor scores 0 and 1. Using logistic regression, we observed a significantly positive association ($p = 0.034$) between ChHV5 seropositivity and turtle size (i.e., SCL). For each 1 cm increase in SCL, turtles were 1.098-fold (95% CI = 1.007–1.197) more likely to test positive for antibodies to ChHV5.

Cohen’s $k$ coefficient revealed poor agreement between: (1) FP status and ChHV5 qPCR results ($k = 0.041$; 95% CI = −0.039–0.121; $p > 0.05$); (2) FP status and ChHV5 ELISA re-
sults ($\kappa = 0.070; 95\% CI = -0.180–0.320; p > 0.05$); and (3) ChHV5 PCR and ELISA results ($\kappa = -0.050; 95\% CI = -0.141–0.040; p > 0.05$). Cohen’s $\kappa$ coefficient was not calculated for ChHV6 qPCR and ELISA results, as all qPCR results were negative for ChHV6.

3.4. Relationships of Fibropapillomatosis and ChHV5 Seroprevalence to Lymphocyte Proliferation Assays

Lymphocyte proliferative responses by capture location were compared in two ways: (1) in turtles with no external tumors (i.e., tumor score 0) and (2) in all turtles (i.e., tumor scores 0–3). In non-tumored turtles, no differences between sites were observed for the T-cell mitogen ConA ($p > 0.05$). No differences between sites were observed for ConA when all turtles, regardless of tumor score, were included ($p > 0.05$ in all cases). All sites were therefore combined for statistical analysis of lymphocyte proliferation assays in relation to tumor score (tumor score 0, $N = 17$; tumor score 1, $N = 8$; tumor score 2, $N = 11$; tumor score 3, $N = 3$). Proliferation results for both suboptimal and optimal stimulation concentrations of ConA showed no differences between turtles with or without FP, by tumor score category (Figure 3a), or between ChHV5 seronegative ($N = 24$) and seropositive turtles ($N = 8$; Figure 3b) ($p > 0.05$ in all cases).

Figure 3. In vitro mitogen-induced lymphocyte proliferation in green turtles (Chelonia mydas) from Florida, USA. T-cell mitogen concanavalin A (ConA) results are shown for suboptimal (ConA: 1 $\mu$g/mL) and optimal (ConA: 10 $\mu$g/mL) stimulation concentrations (mean ± standard error) for comparisons to (a) fibropapilloma tumor score and (b) ChHV5 serology. No significant differences were observed between the different tumor score groups or between seronegative and seropositive turtles at $p < 0.05$. 
3.5. Relationship of Fibropapillomatosis to Natural Killer Cell Activity

Comparisons of natural killer cell activity were compared in two ways in turtles from Big Bend: (1) between tumored (N = 9) and non-tumored (N = 5) turtles and (2) by tumor score (N = 5 for tumor score 0; N = 6 for tumor score 1; N = 3 for tumor score 2). Natural killer cell activity was significantly lower in turtles with FP compared to turtles without FP at an effector to target cell ratio of 50:1 (no FP mean ± SD: 11.2 ± 4.2%; FP mean ± SD: 5.6 ± 3.8%; \(t\)(12) = −2.557; \(p\) = 0.025) (Figure 4a). No significant differences by tumor score were observed for natural killer cell activity (\(H(2) = 4.467; \ p = 0.107\)) at an effector to target cell ratio of 50:1 (Figure 4b).

![Figure 4. In vitro natural killer cell activity in green turtles (Chelonia mydas) with and without fibropapillomatosis (FP) from Florida’s Big Bend, USA measured as a percentage of specific mortality (mean ± standard error). A ratio of effector cells (i.e., natural killer cells) to target cells (i.e., YAC-1 tumor cell lines) of 50:1 was chosen. (a) Natural killer cell activity was significantly higher in turtles (at \(p < 0.05\)) without fibropapilloma (FP) tumors, designated by the asterisk. (b) No statistically significant differences in natural killer cell activity by tumor score were observed.](image-url)

4. Discussion

4.1. Fibropapillomatosis in Green Turtles from Florida

Fibropapillomatosis is considered a panzootic in green turtles [77] that is likely transmitted horizontally once juvenile turtles recruit back to neritic habitats [19,36,78]. Overall prevalence of FP in green turtles in this study was 56% (25/45) and was highest in turtles from Big Bend (69%; 11/16), followed by Jennings Cove (56%; 10/18), and Jupiter Inlet (36%; 4/11). Although not statistically significant, results of FP prevalence and severity based on 10 cm size-class distributions follow what is previously known for green turtles, wherein FP prevalence, severity, and total tumor number are highest in green turtles ranging from 30 to 59.9 cm SCL [13]. Despite the low sample sizes of turtles captured in each foraging aggregation, our results are similar to prior studies. High prevalence of FP in Florida’s green turtles has been previously reported in Big Bend (66%), within the IRL near Sebastian Inlet (up to 72%) and Jennings Cove (up to 90%), and in Lake Worth Lagoon (up to 79%) [14,16,18,65,79,80]. Despite this high occurrence in certain areas of
Florida, several locations in the state host green turtle aggregations with few to no tumors including the Trident Submarine Basin (0–18%), the Key West National Wildlife Refuge (6%), turtles entrained in the St. Lucie Power Plant (up to 2–13%), and nearshore reef sites in central Florida (8–34%) [14,65,79–81]. Typically, high prevalence of FP is reported in poorly circulated lagoon systems that occur near heavily urbanized areas with a number of point- and nonpoint-sources of agricultural and urban effluent [13,14,16,18,19,36]. Therefore, it is expected that FP prevalence rates would be high in certain areas of Florida (e.g., IRL and Lake Worth Lagoon); however, Florida’s Big Bend has one of the highest documented FP prevalence in the state, with some studies reporting rates >80% (albeit with low sample sizes) [17,26]. This finding is unexpected as this area has natural land coverage, numerous conservation protections of its coastal and inland areas, a low human population density, and infrequent occurrences of harmful algal blooms that generate potential tumor-promoting biotoxins [17,22,23,26,58,59]. Four rivers empty into the Big Bend study site both locally (Crystal, Homosassa River, and Chassahowitzka Rivers) and regionally (Fenholloway River), and may carry contaminants from anthropogenic sources that act as FP cofactors [18,82]. Green turtles in Puerto Rico have also shown a higher prevalence of FP in more pristine sites in comparison to areas with increased human development [83]; however, reasons for this remain unknown. Turtles from more pristine sites are therefore not necessarily at a decreased risk of developing FP, and other cofactors besides habitat quality may play a role in FP transmission and pathogenesis. Lastly, presence of leeches or cocoons was not significantly related to FP prevalence, which does not support the hypothesis that Ozobranchus leeches are vectors of ChHV5 [84,85].

Despite a high prevalence of FP in turtles captured in Big Bend, tumor severity and number were highest in turtles captured in Jennings Cove (although not significantly so, likely due to small sample sizes). These findings are similar to other studies in Florida, whereby tumor severity was highest in the IRL compared to a nearshore Sabellariid worm reef and the Trident Submarine Basin. More severe FP in the IRL may be explained by size, environmental, dietary, or genetic differences between individuals in these areas [86]. Another potential explanation related to dissimilarities in FP severity may be due to ChHV5 viral variants that are present in these areas, as multiple studies have shown that different variants can lead to different presentations/severity of disease [35,36,78,87–90]; however, viral variants were not assessed in this study.

Body condition index of green turtles did not differ by tumor score category, and all animals sampled for this study were given subjective body condition scores of “good” or “robust.” In green turtles from Hawaii, Brazil, and Taiwan, no differences in BCI were observed between FP tumor score categories [24,49,91–93]. Interestingly, BCI in Brazilian green turtles was significantly higher in those with FP (all tumor scores combined) compared to FP-free turtles. This was attributed to tumors’ mass contributing to increased body mass, and therefore higher BCI [92]; however, a more likely explanation for our results is that adequate forage items are available in the locations examined here, allowing for the maintenance of good body condition [24]. This finding is corroborated by observations of food present during esophageal lavage (conducted for a separate study) in all turtles sampled during this study. It is also possible that Fulton’s BCI is not an adequate method for detecting differences in turtles with and without FP [47,91] and that subjective estimates (e.g., emaciated, thin, good, and robust) may be better for assessing body condition declines with increasing tumor score [49]. It is noteworthy that only three turtles of tumor score 3 were captured during this study, which prevented rigorous comparisons of BCI to the other tumor score groups.

4.2. Diagnostic Assays for ChHV5 and ChHV6

Detection of ChHV5 DNA in blood (i.e., DNAemia) indicates the presence of circulating ChHV5 DNA, suggesting active or recrudescence infection [40]. Despite an overall FP prevalence of 56% in green turtles captured during this study, only one individual from Florida’s Big Bend tested positive for ChHV5 DNA in whole blood using qPCR. Low
prevalence of ChHV5 in blood samples of turtles with FP is not uncommon as has been observed in previous studies of wild green turtles from the IRL (17%; 33/196) [93] and rehabilitating green turtles at three sea turtle hospitals across the southeastern United States (33%; 7/21) [40]. Low ChHV5 DNA prevalence values were also reported in sea turtles without external evidence of FP, including wild (0%, 0/20), and rehabilitating (14%, 7/52) green turtles from North Carolina, nesting green turtles from Juno Beach, Florida (8%, 5/60), and wild-caught green turtles from Grenada (0%, 0/55) [4,44,94]. Therefore, we expected only a few turtles to test positive for ChHV5 via qPCR, as many herpesviruses become latent within healthy hosts and therefore do not necessarily show relationships with herpesviral DNAemia [4,40]. It is also possible that FP tumors represent niduses for the virus, resulting in low circulating numbers of ChHV5 in the blood [40].

An additional and likely more sensitive method to determine past infection with ChHV5 is through serology, as this method allows for the detection of circulating antibodies to ChHV5 [95]. Once turtles recruit back to neritic habitats, they are more likely to be exposed to ChHV5 [78]. Previous studies of juvenile and subadult green turtles with and without FP captured in three locations across Florida’s east coast demonstrated a seroprevalence for antibodies to ChHV5 of 84% (143/171) [72]. In the present study, seroprevalence was 20% (8/40) for all green turtles, which did not significantly differ by capture location; however, green turtles captured at Jennings Cove had the highest seroprevalence rate (46%; 6/16). This supports our findings of both higher tumor numbers and increased seroprevalence with turtle size, as turtles from Jennings Cove were significantly larger than turtles from Big Bend and Jupiter Inlet (Table 1). Green turtles with experimentally transmitted FP developed antibodies to ChHV5 one year after inoculation; thus, turtles exposed to ChHV5 in near-shore habitats may not immediately test positive for antibodies to ChHV5 [21]. Seronegativity in turtles with external FP tumors could occur if (1) the tumor is antigenically restricted, (2) the virus is “trapped” in but not replicating in tumors (i.e., no immunologic stimulation), (3) immunosuppression is preventing a detectable antibody response, and/or (4) FP-positive, serologically negative turtles are infected with a different serotype of ChHV5 that is not detected with the ELISA used in this study [96,97]. A lower seroprevalence rate of 29% (12/41) was observed in nesting green turtles from Juno Beach, Florida, suggesting that antibody titers may decrease over time post-infection as turtles grow and age, and the virus goes into latency [4]. Virus–tumor interactions may also vary geographically, as green turtles from Florida will often show seropositivity independent of tumor status, whereas most seropositive turtles from Hawaii have tumors. This was attributed to dissimilar behaviors between the two populations, which may have led to the formation of different viral strains with differing pathogenicities [98].

Prevalence of antibodies to ChHV5 was higher in turtles with tumor scores 2 and 3 (43%) compared to 13% in turtles with tumor scores 0 and 1, although this finding was not significant. Increased sample sizes would have likely revealed a significant trend, since turtles in more advanced stages of FP have had adequate time to develop antibodies to ChHV5 [21,55,98]. We also observed low diagnostic agreement between turtles that tested positive for ChHV5 DNA using qPCR and those that tested positive for antibodies to ChHV5, similar to that previously shown in nesting green turtles from Juno Beach, Florida, indicating that these individuals were in different stages of infection (i.e., viral DNAemia as determined by PCR and previous infection as determined by ELISA) [4]. Therefore, we recommend that PCR and serology be utilized simultaneously to best understand ChHV5 infection and exposure [4].

No turtles in this study tested positive for ChHV6 DNA, and just one individual from Jennings Cove tested positive for antibodies to ChHV6. This individual was larger than average compared to other turtles captured for this study (SCL = 49.0 cm), had no external evidence of FP, and also tested positive for antibodies to ChHV5, indicating exposure to both viruses at some point in the turtle’s life [4,54,72]. Lung-eye-trachea disease associated with ChHV6 infection has never been identified in free-ranging green turtles since it was first identified in maricultured green turtles from the Cayman Turtle Farm in 1975 [53]. Indi-
viduals can be exposed and develop antibodies to viruses independently of clinical disease, however, and previous studies have shown that wild green and loggerhead turtles from Florida carry antibodies to ChHV6 at prevalence rates of 10–22% [4,55,95]. No differences in population-level ChHV6 seroprevalence rates were observed over a three-year period in green turtles from Florida, suggesting that ChHV6 exposure was not increasing over time [55]. Our results corroborate these findings. Viral seroconversion and seroreversion within individual turtles have also been observed over relatively short time periods of 5–42 days [44,55]; therefore, some turtles in this study that were previously exposed to ChHV6 (and also ChHV5) may have gone undetected. The suggested transmission mode of ChHV6 is horizontally through seawater, as the virus has been shown to remain infectious for up to five days in this medium with an inverse relationship with water temperature. Horizontal viral transmission via direct contact between turtles and with sediments is also probable [95,99]. Further studies on seroprevalence of ChHV5 and ChHV6 in other sea turtle species from Florida are warranted, as 100% (7/7) of loggerheads in Big Bend tested positive for antibodies to both viruses [100] and foraging grounds serve as areas that facilitate viral transmission [20].

4.3. Lymphocyte Proliferation

Lymphocyte proliferation assays have been utilized in previous studies in an effort to better understand immune function in relation to FP prevalence and/or severity. These studies found reduced T-cell (i.e., cell-mediated immunity) and B-cell (i.e., humoral immunity) proliferative responses in turtles with FP [46], with significant declines in moderately (i.e., tumor score 2) and severely afflicted (i.e., tumor score 3) turtles [47]. Additional changes in hematologic and biochemical parameters include anemia, increased corticosterone, higher heterophil:lymphocyte ratios, heterophilia, hypoproteinemia, and bacteremia, suggesting chronic antigenic stimulation, immunosuppression, and/or chronic stress in relation to FP [47,48,101,102]. No differences were observed, however, between turtles with and without FP in regards to phagocytosis and oxidative burst, which are indicators of innate immune function and antimicrobial defense, respectively [103]. Here, we found no differences in T-cell proliferation between turtles with or without FP, by tumor score (Figure 3a), or between ChHV5 seropositive and seronegative turtles (Figure 3b); yet, severely-afflicted turtles (N = 3) did show reduced T-cell proliferation by 159%, on average, in comparison to the other three groups. Larger sample sizes of turtles in this group could have likely revealed a significant trend. In a recent study, high concentrations of CD3+ lymphocytes and upregulation of leukocyte and lymphatic processes were observed in early-stage tumors, indicating a measurable immune response during earlier stages of tumor growth. Conversely, in turtles that did not survive rehabilitation with late-stage tumors, CD3+ lymphocytes were less concentrated with an observed downregulation of immune- and apoptotic-related genes [104]. Taken together, the evidence suggests that turtles with FP mount an immune response to infection at least until the moderate stages of the disease, followed by a decrease in immune function. Because moderately-afflicted turtles (i.e., tumor score 2) in this study showed the highest proliferative responses for the optimal concentration of the T-cell mitogen (ConA) and all were in good or robust body condition, it is likely that immunosuppression as a result of advanced stages of the disease occurs after late-stage disease as tumor burdens become severe [47]. This argument is especially convincing as immunosuppression is not a prerequisite for the formation of viral-related diseases [20,95] and sufficient and appropriate nutritional input diminishes the immunosuppressive effects of FP [105,106]. Lastly, these results can lend insight into treatment options for turtles with FP/ChHV5 infection through preventative or therapeutic cancer vaccines using tumor antigen specific T-cells [27].

4.4. Natural Killer Cell Activity

Natural killer cells are innate, cytotoxic lymphocytes that function to recognize and clear tumor and virus-infected cells, including a well-characterized response against her-
pesviruses [73,107–109]. Their protective mechanisms lie within the production of cytokines that work towards eliminating pathogens and generating antigen-specific immune responses [110]. Many herpesviruses have developed mechanisms to evade natural killer cell elimination and are known to suppress natural killer cell activity [111]. Only a handful of studies to date have examined natural killer cell activities in turtles, which suggest differences by sex and season, and that activity can decrease upon exposure to certain contaminants [74,112,113]; however, no studies have examined natural killer cell activity in response to disease in sea turtles. Interestingly, several natural killer cell-related cytotoxicity genes were found to be upregulated in FP tumors from juvenile green turtles undergoing rehabilitative care [114]. Additionally, several putative proteins observed in FP tumors that are considered atypical of alphaherpesviruses (e.g., F-lec1 and F-lec2) are associated with activation or inhibition of natural killer cells [115]. Lastly, lymphocytic inflammatory infiltrate, including CD3+ T-cells, is associated with FP tumor tissues [104]. These findings suggest that active inflammatory processes occur within and around FP tumors in response to the high viral load found in tumor tissues compared to healthy, non-tumored tissue [40]. In this study, natural killer cell activity was significantly lower in green turtles with FP compared to turtles without FP (Figure 4a). Collectively, these results indicate that FP tumor development is associated with first an increase, then a subsequent exhaustion of innate immune responses including natural killer cell activity. This depletion of innate immunity may lead to susceptible and/or infected cells becoming more permissive to ChHV5 infection and neoplasia development, respectively. More specifically, ChHV5 infection and FP development may lead to suppression of natural killer cell activity, as has been described with human cancers and infectious and autoimmune diseases. Additionally, depletion of natural killer cells led to enhanced tumor formation in in vivo mouse tumor models [116]. More research is needed to better understand how natural killer cells and other immune responses influence FP disease pathogenesis and prognosis.

5. Conclusions

Although there is no clear trend in the occurrence or severity of fibropapillomatosis in Florida, and there is no evidence that the disease is a major cause of mortality for afflicted animals in this area, >20% of green turtles stranding in southern Florida have FP [13,14,98]. Therefore, advancing our understanding of FP and its impacts on sea turtle health and survival remains an important research priority for sea turtle conservation [36,117]. Lymphocyte proliferative responses of green turtles from this study suggest that turtles in Florida are able to mount sufficient immune responses after the development of tumors, at least until the disease becomes severe. These results support the hypothesis of Work et al. [47] that immunosuppression occurs after disease onset. We propose that adequate forage availability and good-to-robust nutritional status may help prevent severe immunosuppression, at least until the moderate stages of the disease, as the results of our lymphocyte proliferation assays differed from previous studies showing immunosuppression in the more advanced stages of disease [46,47]. Lastly, we observed that natural killer cell activity was significantly reduced in turtles with FP in comparison to tumor-free turtles, lending further insight on immune system function in relation to FP. It is likely that FP tumor development involves factors related to host and viral biology and physiology, in addition to environmental cofactors [20,21,48].

Author Contributions: Conceptualization, J.R.P., M.L. and A.P.-K.; Methodology, J.R.P., M.L., C.R.M., C.M.B., M.J.B., R.M.C., C.R.G., J.R.G., S.E.H., B.W.R., S.T.W., R.C.W., B.E.W. and A.P.-K.; Validation, M.L. and C.R.G.; Formal Analysis, J.R.P., M.L., C.R.G. and A.P.-K.; Data Curation: J.R.P., M.L., C.R.M. and A.P.-K.; Writing—Original Draft Preparation, J.R.P., M.L. and A.P.-K.; Writing—Review and Editing, J.R.P., M.L., C.R.M., C.M.B., M.J.B., R.M.C., C.R.G., J.R.G., S.E.H., B.W.R., S.T.W., R.C.W., B.E.W. and A.P.-K.; Supervision, J.R.P., C.R.M. and A.P.-K.; Project Administration, J.R.P. and A.P.-K.; Funding Acquisition, J.R.P. and A.P.-K. All authors have read and agreed to the published version of the manuscript.
Animals 2021, 11, 861

Funding: Sample analysis for this project was supported by funds provided by a grant to the Florida Center for Coastal and Human Health (FLCCHH). Ongoing funding for this study and for the FLCCHH is provided by the Harbor Branch Oceanographic Institute Foundation. Additional funding for sample analysis was provided by the Florida Department of Environmental Protection (FDEP, #20PBL).

Institutional Review Board Statement: All sampling was conducted under Florida Fish and Wildlife Conservation Commission Marine Turtle Permits 125 and 139 and National Marine Fisheries Service Permit 19528 and 16598. Sampling protocols and collection were reviewed and approved by the Florida Atlantic University Institutional Animal Care and Use Committee under Animal Product Use protocol #A18-14.

Data Availability Statement: The data sets utilized for this study can be made available by the corresponding author upon reasonable request.

Acknowledgments: We thank the staff of East Coast Biologists, Inwater Research Group, Loggerhead Marinelife Center, and Florida Atlantic University’s Harbor Branch Oceanographic Institute including Matt Ajemian, Derek Aoki, Hunter Brown, Karen Holloway-Adkins, Skyler Klingshirn, Ashley Morgan, Marisa Pico, Max Polyak, Kait Toebe, and Steve Traylor for assistance with sample collection and analysis and Meghan Koperski for assistance with the permitting process. Lastly, we thank Amy Wright and Claire Wilson for support with grant administration.

Conflicts of Interest: The authors declare no conflict of interest.

References

1. Jackson, J.B.C.; Kirby, M.X.; Berger, W.H.; Bjorndal, K.A.; Botsford, L.W.; Bourque, B.J.; Bradbury, R.H.; Cooke, R.; Erlandson, J.; Estes, J.A.; et al. Historical overfishing and the recent collapse of coastal ecosystems. Science 2001, 293, 629–638. [CrossRef]

2. Aguirre, A.A.; Lutz, P.L. Marine turtles as sentinels of ecosystem health: Is fibropapillomatosis an indicator? EcoHealth 2004, 1, 275–283. [CrossRef]

3. Chaloupka, M.; Bjorndal, K.A.; Balazs, G.H.; Bolten, A.B.; Ehrhart, L.M.; Limpus, C.J.; Suganuma, H.; Troëng, S.; Yamaguchi, M. Encouraging outlook for recovery of a once severely exploited marine megarherbivore. Glob. Ecol. Biogeogr. 2008, 17, 297–304. [CrossRef]

4. Page-Karjian, A.; Chabot, R.; Stacy, N.I.; Morgan, A.S.; Valverde, R.A.; Stewart, S.; Coppenrath, C.M.; Manire, C.A.; Herbst, L.H.; Gregory, C.R.; et al. Comprehensive health assessment of green turtles Chelonia mydas nesting in southeastern Florida, USA. Endang. Spec. Res. 2020, 42, 21–35. [CrossRef]

5. Thayer, G.W.; Bjorndal, K.A.; Ogden, J.C.; Williams, S.L.; Zieman, J.C. Role of larger herbivores in seagrass communities. Estuaries 1984, 7, 351–376. [CrossRef]

6. Bjorndal, K.A.; Bolten, A.B.; Chaloupka, M.Y. Green turtle somatic growth model: Evidence for density dependence. Ecol. Appl. 2000, 10, 269–282.

7. Witherington, B.E.; Bresette, M.J.; Herren, R. Chelonia mydas—Green turtle. In Biology and Conservation of Florida Turtles; Meylan, P.A., Ed. Chelonian Res. Monogr. 2006, 3, 90–104. [CrossRef]

8. Valdivia, A.; Wolf, S.; Suckling, K. Marine mammals and sea turtles listed under the U.S. Endangered Species Act are recovering. PLoS ONE 2019, 14, e0210164. [CrossRef] [PubMed]

9. Ehrhart, L.M.; Redfoot, W.E.; Bagley, D.A. Marine turtles of the central region of the Indian River Lagoon System, Florida. Fla. Sci. 2007, 70, 415–434.

10. Seminoff, J.A.; Allen, C.D.; Balazs, G.H.; Dutton, P.H.; Eguchi, T.; Haas, H.L.; Hargrove, S.A.; Jensen, M.; Klemm, D.L.; Lauritsen, A.M.; et al. Status Review of the Green Turtle (Chelonia mydas) under the Endangered Species Act. NOAA Tech Memo NOAA-NMFS-SWFC-59; NOAA Tech Memo: La Jolla, CA, USA, 2015; p. 571.

11. Brost, B.; Witherington, B.; Meylan, A.; Leone, E.; Ehrhart, L.; Bagley, D. Sea turtle hatching production from Florida (USA) beaches, 2002–2012, with recommendations for analyzing hatching success. Endang. Spec. Res. 2015, 27, 53–68. [CrossRef]

12. Florida Fish and Wildlife Conservation Commission (FWC). Index Nesting Beach Survey Totals (1989–2019). 2020. Available online: https://myfwc.com/research/wildlife/sea-turtles/nesting/beach-survey-totals/ (accessed on 30 October 2020).

13. Foley, A.M.; Schroeder, B.A.; Redlow, A.E.; Fick-Child, K.J.; Teas, W.G. Fibropapillomatosis in stranded green turtles (Chelonia mydas) from the eastern United States (1980–1998): Trends and associations with environmental factors. J. Wildl. Dis. 2005, 41, 29–41. [CrossRef] [PubMed]

14. Hiramia, S.; Ehrhart, L.M. Description, prevalence and severity of green turtle fibropapillomatosis in three developmental habitats on the east coast of Florida. Fla. Sci. 2007, 70, 435–448.

15. Hiramia, S.; Ehrhart, L.M.; Rea, L.D.; Kittie, R.A. Relating fibropapilloma tumor severity to blood parameters in green turtles Chelonia mydas. Dis. Aquat. Org. 2014, 111, 61–68. [CrossRef]

16. Gorham, J.C.; Bresette, M.J.; Guertin, J.R.; Shamblin, B.M.; Nairn, C.J. Green turtles (Chelonia mydas) in an urban estuary system: Lake Worth Lagoon, Florida. Fla. Sci. 2016, 79, 14–27.

Acknowledgments: We thank the staff of East Coast Biologists, Inwater Research Group, Loggerhead Marinelife Center, and Florida Atlantic University’s Harbor Branch Oceanographic Institute including Matt Ajemian, Derek Aoki, Hunter Brown, Karen Holloway-Adkins, Skyler Klingshirn, Ashley Morgan, Marisa Pico, Max Polyak, Kait Toebe, and Steve Traylor for assistance with sample collection and analysis and Meghan Koperski for assistance with the permitting process. Lastly, we thank Amy Wright and Claire Wilson for support with grant administration.
17. Perrault, J.R.; Stacy, N.I.; Lehner, A.F.; Mott, C.R.; Hirsch, S.; Gorham, J.C.; Buchweitz, J.P.; Bresette, M.J.; Walsh, C.J. Potential effects of brevotoxins and toxic elements on various health variables in Kemp's ridley (Lepidochelys kempi) and green (Chelonia mydas) sea turtles after a red tide bloom event. *Sci. Total Environ.* 2017, 605–606, 967–979. [CrossRef]

18. Chabot, R.M.; Welsh, R.C.; Mott, C.R.; Guertin, J.R.; Shamblin, B.M.; Witherington, B.E. A sea turtle population assessment for Florida's Big Bend, northeastern Gulf of Mexico. *Gulf Caribb. Res.* 2021, in press.

19. Herbst, L.H. Fibropapillomatosis of marine turtles. *Ann. Rev. Fish. Dis.* 1994, 4, 389–425. [CrossRef]

20. Herbst, L.H.; Klein, P.A. Green turtle fibropapillomatosis: Challenges to assessing the role of environmental cofactors. *Environ. Health Perspect.* 1995, 103, 27–30.

21. Herbst, L.H.; Jacobson, E.R.; Moretti, R.; Brown, T.; Sundberg, J.P.; Klein, P.A. Experimental transmission of green turtle fibropapillomatosis using cell-free tumor extracts. *Dis. Aquat. Org.* 1995, 22, 1–12. [CrossRef]

22. Landsberg, J.H.; Balazs, G.H.; Steidinger, K.A.; Baden, D.G.; Work, T.M.; Russell, D.J. The potential role of natural tumor promoters in marine turtle fibropapillomatosis. *J. Aquat. Anim. Health* 1999, 11, 199–210. [CrossRef]

23. Arthur, K.; Limpus, C.; Balazs, G.; Capper, A.; Udy, J.; Shaw, G.; Keuper-Bennett, U.; Bennett, P. The exposure of green turtles (*Chelonia mydas*) to tumour promoting compounds produced by the cyanobacterium *Lyngbya majuscula* and their potential role in the aetiology of fibropapillomatosis. *Harmful Algae* 2008, 7, 114–125. [CrossRef]

24. dos Santos, R.G.; Martins, A.S.; Torezani, E.; Baptistotte, C.; Farias, J.d.N.; Horta, P.A.; Work, T.M.; Balazs, G.H. Relationship between fibropapillomatosis and environmental quality: A case study with *Chelonia mydas* off Brazil. *Dis. Aquat. Org.* 2010, 89, 87–95. [CrossRef] [PubMed]

25. da Silva, C.C.; Klein, R.D.; Barcarolli, I.F.; Bianchini, A. Metal contamination as a possible etiology of fibropapillomatosis in juvenile female green sea turtles *Chelonia mydas* from the southern Atlantic Ocean. *Aquat. Toxicol.* 2016, 170, 42–51. [CrossRef] [PubMed]

26. Perrault, J.R.; Perkins, C.R.; Ajemian, M.J.; Bresette, M.J.; Mott, C.R.; Page-Karjian, A. Harmful algal and cyanobacterial toxins in foraging green turtles (*Chelonia mydas*) in Florida’s Big Bend. *Toxicon X* 2020, 5, 100020. [CrossRef]

27. Duffy, D.J.; Schnitzler, C.; Karpinski, L.; Thomas, R.; Whilde, J.; Eastman, C.; Yang, C.; Krstic, A.; Rollinson, D.; Zirkelbach, B.; et al. Sea turtle fibropapilloma tumors share genomic drivers and therapeutic vulnerabilities with human cancers. *Commun. Biol.* 2018, 1, 63. [CrossRef] [PubMed]

28. Smith, G.M.; Coates, C.W. Fibro-epithelial growths of the skin in large marine turtles, *Chelonia mydas* (Linnaeus). *Zoologica* 1938, 23, 93–98.

29. Work, T.M.; Balazs, G.H. Relating tumor score to hematology in green turtles with fibropapillomatosis in Hawaii. *J. Wildl. Dis.* 1999, 35, 804–807. [CrossRef]

30. Limpus, C.J.; Couper, P.J.; Couper, K.L.D. Crab Island revisited: Reassessment of the world’s largest flatback turtle rookery after twelve years. *Mem. Qld. Mus.* 1993, 33, 227–289.

31. Barragan, A.R.; Sarti, M.L. A possible case of fibropapilloma in Kemp’s ridley turtle (*Lepidochelys kempi*). *Mar. Turtle Newsl.* 1994, 67, 27.

32. Quackenbush, S.L.; Work, T.M.; Balazs, G.H.; Casey, R.N.; Rovnak, A.; Chaves, A.; duToit, L.; Baines, C.R.; Parrish, C.R.; Bowser, P.R.; et al. Three closely related herpesviruses are associated with fibropapillomatosis in marine turtles. *Virology* 1998, 246, 392–399. [CrossRef]

33. Aguirre, A.A.; Spraker, T.R.; Chaves, A.; Toit, L.; Eure, W.; Balazs, G.H. Pathology of fibropapillomatosis in olive ridley turtles *Lepidochelys olivacea* nesting in Costa Rica. *J. Aquat. Health* 1999, 11, 283–289. [CrossRef]

34. D’Amato, A.F.; Moraes-Neto, M. First documentation of fibropapillomas verified by histopathology in *Eretmochelys imbricata*. *Mar. Turt. Newsl.* 2000, 89, 12–13.

35. Page-Karjian, A.; Norton, T.M.; Harms, C.; Mader, D.; Herbst, L.H.; Stedman, N.; Gottdenker, N.L. Case descriptions of fibropapillomatosis in rehabilitating loggerhead sea turtles *Caretta caretta* in the southeastern USA. *Dis. Aquat. Org.* 2015, 115, 185–191. [CrossRef]

36. Jones, K.; Ariel, E.; Burgess, G.; Read, M. A review of fibropapillomatosis in green turtles (*Chelonia mydas*). *Vet. J.* 2016, 212, 48–57. [CrossRef]

37. Diaz-Delgado, J.; Gomes-Borges, J.C.; Silveira, A.M.; Einhardt-Vergara, J.; Groch, K.R.; Cirqueira, C.S.; Sansone, M.; Gattamorta, M.A.; Matsushima, E.R.; Cataro-Dias, J.L. Primary multicentric pulmonary low-grade fibromyxoid sarcoma and cheloid alapherpesvirus 5 in a leatherback sea turtle (*Dermochelys coriacea*). *Comp. Pathol. 2019, 168, 1–7. [CrossRef]

38. Herbst, L.H.; Jacobson, E.R.; Klein, P.A.; Balazs, G.H.; Moretti, R.; Brown, T.; Sundberg, J.P. Comparative pathology and pathogenesis of spontaneous and experimentally induced fibropapillomas of green turtles (*Chelonia mydas*). *Vet. Pathol.* 1999, 36, 551–564. [CrossRef]

39. Monezi, T.A.; Mehnert, D.U.; de Moura, E.M.M.; Muller, N.M.G.; Garrafa, P.; Matsushima, E.R.; Werneck, M.R.; Borella, M.I. Cheloid herpesvirus 5 in secretions and tissue samples from green turtles (*Chelonia mydas*) from Southeastern Brazil: A ten-year study. *Vet. Microbiol.* 2016, 186, 150–156. [CrossRef]

40. Page-Karjian, A.; Norton, T.M.; Ritchie, B.; Brown, C.C.; Mancia, C.; Jackwood, M.; Gottdenker, N.L. Quantifying cheloid herpesvirus 5 in symptomatic and asymptomatic rehabilitating green sea turtles. *Endanger. Species Res.* 2015, 28, 135–146. [CrossRef]
41. Page-Karjian, A.; Norton, T.M.; Ritchie, B.; Whitfield, J.; Gottdenker, N.L. Potential non-cutaneous sites of chelonid herpesvirus 5 persistence and shedding in green sea turtles (*Chelonia mydas*). *J. Aquat. Anim. Health* 2017, 29, 136–142. [CrossRef]

42. Yetsko, K.; Farrell, J.A.; Blackburn, N.B.; Whitmore, L.; Stamm, M.R.; Whilde, J.; Eastman, C.B.; Ramia, D.R.; Thomas, R.; Kristic, A.; et al. Molecular characterization of a marine turtle tumor epizootic, profiling external, internal and post-surgical regrowth tumors. *Commun. Biol.* 2021, 4, 152. [CrossRef]

43. Milton, S.L.; Lutz, P.L. Physiological and genetic responses to environmental stress. In *The Biology of Sea Turtles Volume II*; Lutz, P.L., Musick, J.A., Wynken, J., Eds.; CRC Press: Boca Raton, FL, USA, 2003; pp. 163–197.

44. Page-Karjian, A.; Barile, P.J. Widespread sewage pollution of the Indian River Lagoon system, Florida (USA) resolved by spatial analyses of occurrences of *Chelonia mydas* and *Caretta caretta* sea turtles in North Carolina, USA, 2015–2019. *Animals* 2020, 10, 1964. [CrossRef] [PubMed]

45. Aguirre, A.A.; Balazs, G.H. Blood biochemistry values of green turtles, *Chelonia mydas*, with and without fibropapillomatosis. *Comp. Haematol. Int.* 2000, 10, 132–137. [CrossRef]

46. Clay, C.; Varella, R.; Bossart, G.D.; Lutz, P. Altered in vitro immune responses in green turtles (*Chelonia mydas*) with fibropapillomatosis. *J. Zoo Wildl. Med.* 2001, 32, 436–440. [PubMed]

47. Work, T.M.; Ramey, R.A.; Balazs, G.H.; Clay, C.; Chang, S.P. Immune status of free-ranging green turtles with fibropapillomatosis from Hawaii. *J. Wildl. Dis.* 2001, 37, 574–581. [CrossRef] [PubMed]

48. Page-Karjian, A.; Barile, P.J. Prevalence and Trends in Fibropapillomatosis in southeast Brazil. *Dis. Aquat. Org.* 2015, 115, 193–201. [CrossRef]

49. Deus Santos, M.R.d.; Martins, A.S.; Baptистotte, C.; Work, T.M. Health condition of juvenile *Chelonia mydas* related to fibropapillomatosis in southeast Brazil. *Dis. Aquat. Org.* 2015, 115, 193–201. [CrossRef]

50. da Silva, C.C.; Bianchini, A. Blood cholesterol as a biomarker of fibropapillomatosis in green turtles. *Mar. Turt. Newsl.* 2019, 158, 16–21.

51. Li, T.-H.; Chang, C.-C. The impact of fibropapillomatosis on clinical characteristics, blood gas, plasma biochemistry, and hematological profiles in juvenile green turtles (*Chelonia mydas*). *Bull. Mar. Sci.* 2020, 96, 723–734. [CrossRef]

52. Davison, A.J. Herpesvirus systematics. *Vet. Microbiol.* 2010, 143, 52–69. [CrossRef]

53. Jacobson, E.R.; Gaskin, J.M.; Roelke, M.; Greiner, E.C.; Allen, J. Conjunctivitis, tracheitis, and pneumonia associated with herpesvirus infection in green sea turtles. *J. Zoo Wildl. Med.* 2014, 45, 507–519. [CrossRef]

54. Coberley, S.S.; Herbst, L.H.; Brown, D.R.; Ehrhart, L.M.; Bagley, D.A.; Schaf, S.A.; Moretti, R.H.; Jacobson, E.R.; Klein, P.A. Detection of antibodies to a disease-associated herpesvirus of the green turtle, *Chelonia mydas*. *J. Clin. Microbiol.* 2001, 39, 3572–3577. [CrossRef]

55. Coberley, S.S.; Herbst, L.H.; Ehrhart, L.M.; Bagley, D.A.; Hiramia, S.; Jacobson, E.R.; Klein, P.A. Survey of Florida green turtles for exposure to a disease-associated herpesvirus. *Dis. Aquat. Org.* 2001, 47, 159–167. [CrossRef]

56. Iverson, R.L.; Bittaker, H.F. Seagrass distribution and abundance in eastern Gulf of Mexico coastal waters. *Estuar. Coast. Shelf Sci.* 1986, 22, 577–602. [CrossRef]

57. Hale, J.A.; Frazer, T.K.; Tomasko, D.A.; Hall, M.O. Changes in the distribution of seagrass species along Florida’s Central Gulf Coast: Iverson and Bittaker revisited. *Estuaries 2004*, 27, 36–43. [CrossRef]

58. Livingston, R.J. Ontogenetic trophic relationships and stress in a coastal seagrass system in Florida. In *Estuarine Perspectives*; Kennedy, V., Ed.; Academic Press: New York, NY, USA, 1980; pp. 423–435.

59. Seavy, J.R.; Pine III, W.E.; Frederick, P.; Sturmer, L.; Berrigan, M. Decadal changes in oyster reefs in the Big Bend of Florida’s Gulf Coast. *Ecosphere* 2011, 2, 1–14. [CrossRef]

60. Swain, H.M.; Breininnger, D.R.; Busby, D.S.; Clark, K.B.; Cook, S.B.; Day, R.A.; Defreece, D.E.; Gilmore, R.G.; Hart, A.W.; Hinkle, C.R.; et al. Indian River Lagoon biodiversity conference-Introduction. *Bull. Mar. Sci.* 1995, 57, 1–7.

61. Indian River Lagoon National Estuary Program (IRLNEP). *IRL Comprehensive Conservation and Management Plan Update; IRL National Estuary Program: Palm Bay, FL, USA, 2008;* p. 120. Available online: https://www.epa.gov/sites/production/files/2015-09/documents/ccmp_update_2008_final.pdf (accessed on 6 November 2020).

62. Sigua, G.C.; Tweedale, W.A. Watershed scale assessment of nitrogen and phosphorus loadings in the Indian River Lagoon basin, Florida. *J. Environ. Manag.* 2003, 67, 363–372. [CrossRef]

63. Barile, P.J. Evidence of anthropogenic nitrogen enrichment of the littoral waters of east central Florida. *J. Coast. Res.* 2004, 20, 1237–1245. [CrossRef]

64. Barile, P.J. Widespread sewage pollution of the Indian River Lagoon system, Florida (USA) resolved by spatial analyses of macroalgal biochemistry. *Mar. Pollut. Bull.* 2018, 128, 557–574. [CrossRef]

65. Ehrhart, L.; Redfoot, W.; Mansfield, K.; Gorham, J.; Weege, S.; Provancha, J. Prevalence and Trends in Fibropapillomatosis in Green Turtles on Florida’s Atlantic Coast. In *Proceedings of the 2015 International Summit on Fibropapillomatosis: Global Status, Trends, and Population Impacts, Honolulu, HI, USA, 11–14 June 2015*; Hargrove, S., Work, T., Brunson, S., Foley, A.M., Balazs, G., Eds.; NOAA Technical Memorandum, NOAA-TM-NMFS-PIFSC-54. U.S. Department of Commerce: Honolulu, HI, USA, 2016; pp. 15–21.

66. Crigger, D.K.; Graves, G.A.; Fike, D.L. Lake Worth Lagoon conceptual ecological model. *Wetlands* 2005, 25, 943–954.

67. Bolter, K. An Ecological Study of Photoautotrophs in Lake Worth. Master’s Thesis, Florida Atlantic University, Boca Raton, FL, USA, 2010.
Animals 2021, 11, 861

68. Eaton, C.; McMichael, E.; Witherington, B.; Foley, A.; Hardy, R.; Meylan, A. In-Water Sea Turtle Monitoring and Research in Florida: Review and Recommendations; NOAA Technical Memorandum NMFS-OPR-38; U.S. Department of Commerce: St. Petersburg, FL, USA, 2008; p. 233.

69. Deem, S.L.; Harris, H.S. Health assessments. In Sea Turtle Health and Rehabilitation; Manire, C.A., Norton, T.M., Stacy, B.A., Harms, C.A., Innis, C.J., Eds.; J. Ross Publishing: Plantation, FL, USA, 2017; pp. 945–957.

70. Page-Karjian, A.; Perrault, J.R. Sea turtle health assessments: Maximizing turtle encounters to better understand health. In Sea Turtle Research and Conservation: Lessons from Working in the Field; Nahill, B., Ed.; Academic Press: San Diego, CA, USA, 2021; pp. 31–44.

71. Altschul, S.F.; Gish, W.; Miller, W.; Myers, E.W.; Lipman, D.J. Basic local alignment search tool. J. Mol. Biol. 1990, 215, 403–410. [CrossRef]

72. Herbst, L.H.; Lemaire, S.; Ene, A.R.; Haslin, D.J.; Ehrhart, L.M.; Bagley, D.A.; Klein, P.A.; Lenz, J. Use of baculovirus-expressed glycoprotein H in an enzyme-linked immunosorbent assay developed to assess exposure to chelonid fibropapillomatosis-associated herpesvirus and its relationship to the prevalence of fibropapillomatosis in sea turtles. Clin. Vaccine Immunol. 2008, 15, 843–851. [CrossRef] [PubMed]

73. Rousselet, E.; Levin, M.; Gebhard, E.; Higgins, B.M.; De Guise, S.; Godard-Coddin, C.A. Evaluation of immune functions in captive immature loggerhead sea turtles (Caretta caretta). Vet. Immunol. Immunopathol. 2013, 156, 43–53. [CrossRef]

74. Rousselet, E.; Levin, M.; Gebhard, E.; Higgins, B.M.; De Guise, S.; Godard-Coddin, C. Polychlorinated biphenyls modulate both phagocytosis and NK cell activity in vitro in juvenile loggerhead sea turtles (Caretta caretta). J. Toxicol. Environ. Health A 2017, 80, 556–561. [CrossRef]

75. Landis, J.R.; Koch, G.G. The measurement of observer agreement for categorical data. Biometrics 1977, 33, 159–174. [CrossRef] [PubMed]

76. FWC. Marine Turtle Conservation Handbook. 2016. Available online: https://myfwc.com/media/3133/fwc-mtconservationhandbook.pdf (accessed on 10 December 2020).

77. Williams, E.H.; Bunkley-Williams, L.; Peters, E.C.; Pinto-Rodriguez, B.; Matos-Morales, R.; Mignucci-Giannoni, A.A.; Hall, K.V.; Rueda-Almonacid, J.V.; Sybesma, J.; de Calventi, I.B.; et al. An epizootic of cutaneous fibropapillomas in green turtles Chelonia mydas of the Caribbean: Part of a panzootic? J. Aquat. Anim. Health 1994, 6, 70–78. [CrossRef]

78. Ene, A.; Su, M.; Lemaire, S.; Rose, C.; Schaff, S.; Moretti, R.; Lenz, J.; Herbst, L.H. Distribution of chelonid fibropapillomatosis-associated herpesvirus variants in Florida: Molecular genetic evidence for infection of turtles following recruitment to neritic developmental habitats. J. Wildl. Dis. 2005, 41, 489–497. [CrossRef] [PubMed]

79. Holloway-Adkins, K.G. A comparative study of the feeding ecology of Chelonia mydas (green turtle) and the incidental ingestion of Prorocentrum ssp. Master’s Thesis, University of Central Florida, Orlando, FL, USA, 2001.

80. Bresette, M.J.; Herren, R.M.; Singewald, D.A. Comparison of Fibropapilloma Rates of Green Turtles (Chelonia mydas) from Two Different Sites in St. Lucie County, Florida. In Proceedings of the Twenty-First Annual Symposium on Sea Turtle Biology and Conservation, Philadelphia, PA, USA, 24–28 February 2001; NOAA Technical Memorandum, NOAA-NMFS-SEFSC-528. Coyne, M.S., Clark, R.D., Eds.; U.S. Department of Commerce: Miami, FL, USA, 2004; pp. 125–126.

81. Herren, R.M.; Bagley, D.A.; Bresette, M.J.; Holloway-Adkins, K.G.; Clark, R.D.; Witherington, B.E. Sea turtle abundance and demographic measurements in a marine protected area in the Florida Keys, USA. Herpetol. Conserv. Biol. 2018, 13, 224–239.

82. Mattson, R.A. Seagrass ecosystem characteristics and research and management needs in the Florida Big Bend. In Seagrasses: Monitoring, Ecology, Physiology, and Management; Borlone, S.A., Ed.; CRC Press: Boca Raton, FL, USA, 2000; pp. 259–277.

83. Patricio, A.R.; Velez-Zuazo, X.; Diez, C.E.; Van Dam, R.; Sabat, A.M. Survival probability of immature green turtles in two foraging grounds at Culebra, Puerto Rico. Mar. Ecol. Prog. Ser. 2011, 440, 217–227. [CrossRef]

84. Greenblatt, R.J.; Work, T.M.; Balazs, G.H.; Sutton, C.A.; Casey, R.N.; Casey, J.W. The Ozobranchus leech is a candidate mechanical vector for the fibropapilloma-associated turtle herpesvirus found latently infecting skin tumors on Hawaiian green turtles (Chelonia mydas). Virology 2004, 321, 101–110. [CrossRef]

85. Rittenburg, L.T.; Kelley, J.R.; Mansfield, K.L.; Savage, A.E. Marine leech parasitism of sea turtles varies across host species, seasons, and the tumor disease fibropapillomatosis. Dis. Aquat. Org. 2021, 143, 1–12. [CrossRef] [PubMed]

86. Alfaro-Núñez, A.; Bertelsen, M.F.; Bojesen, A.M.; Rasmussen, I.; Zepeda-Mendoza, L.; Olsen, M.T.; Gilbert, M.T.P. Global distribution of Chelonid fibropapilloma-associated herpesvirus among clinically healthy sea turtles. BMC Evol. Biol. 2014, 14, 206. [CrossRef]

87. Laegreid, W.W.; Skowronek, A.; Stone-Marschat, M.; Burrage, T. Characterization of virulence of African horsesickness virus. Virology 1993, 195, 836–839. [CrossRef]

88. Herbst, L.; Ene, A.; Su, M.; Sesalle, R.; Lenz, J. Tumor outbreaks in marine turtles are not due to recent herpesvirus mutations. Curr. Biol. 2004, 14, R697–R699. [CrossRef] [PubMed]

89. Yunis, R.; Jarosinski, K.W.; Schat, K.A. Association between rate of viral genome replication and virulence of Marek’s disease herpesvirus strains. Virology 2004, 328, 142–150. [CrossRef] [PubMed]

90. Jones, K.; Burgess, G.; Budd, A.M.; Huerlimann, R.; Mashkour, N.; Ariel, E. Molecular evidence for horizontal transmission of chelonid alphaherpesvirus 5 at green turtle (Chelonia mydas) foraging grounds in Queensland, Australia. PLoS ONE 2020, 15, e0227268. [CrossRef] [PubMed]
Animals 2021, 11, 861

91. Work, T.M.; Balazs, G.H.; Rameye, R.A.; Morris, R.A. Retrospective pathology survey of green turtles Chelonia mydas with fibropapillomatosis in the Hawaiian Islands, 1993–2003. *Dis. Aquat. Org.* 2004, 62, 163–176. [CrossRef][PubMed]

92. Rossi, S.; Sánchez-Sarmiento, A.M.; dos Santos, R.G.; Zamana, R.R.; Prioste, F.E.S.; Gattamorta, M.A.; Ochoa, P.F.C.; Grisibili, J.H.H.; Matushima, E.R. Monitoring green sea turtles in Brazilian feeding areas: Relating body condition index to fibropapillomatosis prevalence. *J. Mar. Biol. Assoc. UK* 2019, 59, 1879–1887. [CrossRef]

93. Lawrance, M.F.; Mansfield, K.L.; Sutton, E.; Savage, A.E. Molecular evolution of fibropapilloma-associated herpesviruses infecting juvenile green and loggerhead sea turtles. *Virology* 2018, 521, 190–197. [CrossRef]

94. James, A. Prevalence of Chelonid Alphaherpesvirus 5 and Associated Disease in Sea Turtles in Grenada, West Indies. Master’s Thesis, St. George’s University, True Blue, Grenada, 2020.

95. Page-Karjian, A.; Herbst, L.H. Viruses. In *Sea Turtle Health and Rehabilitation*; Manire, C.A., Norton, T.M., Stacy, B.A., Harms, C.A., Innis, C.J., Eds.; J. Ross Publishing: Plantation, FL, USA, 2017; pp. 751–777.

96. Rouse, B.T.; Sehrawat, S. Immunity and immunopathology to viruses: What decides the outcome? *Nat. Rev. Immunol.* 2010, 10, 514–526. [CrossRef]

97. White, D.W.; Beard, R.S.; Barton, E.B. Immune modulation during latent herpesvirus infection. *Immunol. Rev.* 2012, 245, 189–208. [CrossRef][PubMed]

98. Work, T.M.; Dagenais, J.; Willimann, A.; Balazs, G.; Mansfield, K.; Ackermann, M. Differences in antibody response against chelonid alpha herpesvirus 5 (ChHV5) suggest differences in virus biology in ChHV5-seropositive green turtles from Hawaii and ChHV5-seropositive green turtles from Florida. *J. Virol.* 2020, 94, e01658-19. [PubMed]

99. Curry, S.S.; Brown, D.R.; Gaskin, J.M.; Jacobson, E.R.; Ehrhart, L.M.; Blahak, S.; Herbst, L.H.; Klein, P.A. Persistent infectivity of a disease-associated herpesvirus in green sea turtles after exposure to seawater. *J. Wildl. Dis.* 2000, 36, 792–797. [CrossRef][PubMed]

100. Perrault, J.R.; Page-Karjian, A. Disease survey of loggerhead sea turtles (Caretta caretta) in Florida’s Big Bend. Unpublished data.

101. Aguirre, A.A.; Balazs, G.H.; Spraker, T.R.; Gross, T.S. Adrenal and hematological responses to stress in juvenile green turtles (Chelonia mydas) with and without fibropapillomas. *Physiol. Zool.* 1995, 68, 831–854. [CrossRef]

102. Work, T.M.; Balazs, G.H.; Volcott, M.; Morris, R. Bacteraemia in free-ranging Hawaiian green turtles Chelonia mydas with fibropapillomatosis. *Dis. Aquat. Org.* 2003, 53, 41–46. [CrossRef][PubMed]

103. Rossi, S.; Hazarbashanov, N.G.T.d.Q.; Sánchez-Sarmiento, A.M.; Prioste, F.E.S.; Matushima, E.R. Immune response of green sea turtles with and without fibropapillomatosis: Evaluating oxidative burst and phagocytosis via flow cytometry. *Chelonian Conserv. Biol.* 2016, 15, 273–278. [CrossRef]

104. Yetsko, K.; Farrell, J.; Stammnitz, M.R.; Whitmore, L.; Whilde, J.; Eastman, C.B.; Ramia, D.R.; Thomas, R.; Krstic, A.; Linse, P.; et al. Mutational, transcriptional and viral shedding dynamics of the marine turtle fibropapillomatosis tumor epizootic. *bioRxiv* 2021, in press.

105. Borysenko, M.; Lewis, S. The effect of malnutrition on immunocompetence and whole body resistance to infection in Chelydra serpentina. *Dev. Comp. Immunol.* 1979, 3, 89–100. [CrossRef]

106. Acevedo-Whitehouse, K.; Duffus, A.L.J. Effects of environmental change on wildlife health. *Philos. Trans. R. Soc. B* 2009, 364, 3429–3438. [CrossRef][PubMed]

107. Cervenka, A.; Lanier, L.L. Natural killer cells, viruses and cancer. *Nat. Rev. Immunol.* 2001, 1, 41–49. [CrossRef][PubMed]

108. Keller, J.M.; Peden-Adams, M.M.; Aguirre, A.A. Immunotoxicology and implications for reptilian health. In *Toxicology of Reptiles*; Gardner, S.C., Oberdörster, E., Eds.; CRC Press: Boca Raton, FL, USA, 2006; pp. 199–240.

109. Ghorai, S.M.; Priyam, M. Reptilia: Cellular immunity in reptiles: Perspective on elements of evolution. In *Endanger. Spec. Res.* 2012, 2001, 171, 195–200.

110. Blackburn, N.B.; Leandro, A.C.; Nahvi, N.; Devlin, M.A.; Leandro, M.; Escobedo, I.M.; Peralta, J.M.; George, J.; Stacy, B.A.; deMaar, T.W.; et al. Transcriptomic profiling of fibropapillomatosis in green sea turtles (Chelonia mydas) from south Texas. *Front. Immunol.* 2021, 12, 630988. [CrossRef][PubMed]

111. Ackermann, M.; Koriabine, M.; Hartmann-Fritsch, F.; de Jong, P.J.; Lewis, T.D.; Schetle, N.; Work, T.M.; Dagenais, J.; Balazs, G.H.; Leong, J.-A.C. The genome of chelonid herpesvirus 5 harbors atypical genes. *Philos. Trans. R. Soc. B* 2019, 364, 3429–3438. [CrossRef][PubMed]

112. Muñoz, F.J.; De la Fuente, M. The immune response of thymic cells from the turtle Mauremys caspica. *Physiol. Biochem. Zool.* 2001, 74, 667–672. [CrossRef]

113. Muñoz, F.J.; De la Fuente, M. The immune response of thymic cells from the turtle Mauremys caspica. *J. Comp. Physiol. B* 2001, 171, 195–200.

114. Blackburn, N.B.; Leandro, A.C.; Nahvi, N.; Devlin, M.A.; Leandro, M.; Escobedo, I.M.; Peralta, J.M.; George, J.; Stacy, B.A.; deMaar, T.W.; et al. Transcriptomic profiling of fibropapillomatosis in green sea turtles (Chelonia mydas) from south Texas. *Front. Immunol.* 2021, 12, 630988. [CrossRef][PubMed]

115. Ackermann, M.; Koriabine, M.; Hartmann-Fritsch, F.; de Jong, P.J.; Lewis, T.D.; Schetle, N.; Work, T.M.; Dagenais, J.; Balazs, G.H.; Leong, J.-A.C. The genome of chelonid herpesvirus 5 harbors atypical genes. *PLOS ONE* 2012, 7, e46623. [CrossRef][PubMed]

116. Smyth, M.J.; Godfrey, D.I.; Trapani, J.A. A fresh look at tumor immunosurveillance and immunotherapy. *Nat. Immunol.* 2001, 2, 293–299. [CrossRef]

117. Hamann, M.; Godfrey, M.; Seminoff, J.; Arthur, K.; Barata, P.; Bjornsdal, K.; Bolten, A.; Broderick, A.; Campbell, L.; Carreras, C.; et al. Global research priorities for sea turtles: Informing management and conservation in the 21st century. *Endanger. Spec. Res.* 2010, 11, 245–269. [CrossRef]