Dead tired: evaluating the physiological status and survival of neonatal reef sharks under stress

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Marine protected areas (MPAs) can protect shark populations from targeted fisheries, but resident shark populations may remain exposed to stressors like capture as bycatch and environmental change. Populations of young sharks that rely on shallow coastal habitats, e.g. as nursery areas, may be at risk of experiencing these stressors. The purpose of this study was to characterize various components of the physiological stress response of neonatal reef sharks following exposure to an exhaustive challenge under relevant environmental conditions. To accomplish this, we monitored markers of the secondary stress response and measured oxygen uptake rates ($\dot{MO}_{2}$) to compare to laboratory-derived baseline values in neonatal blacktip reef ($Carcharhinus melanopterus$) and sicklefin lemon sharks ($Negaprion acutidens$). Measurements occurred over three hours following exposure to an exhaustive challenge (gill-net capture with air exposure). Blood lactate concentrations and pH deviated from baseline values at the 3-h sample, indicating that both species were still stressed 3 h after capture. Evidence of a temperature effect on physiological status of either species was equivocal over 28–31°C. However, aspects of the physiological response were species-specific; $N. acutidens$ exhibited a larger difference in blood pH relative to baseline values than $C. melanopterus$, possibly owing to higher minimum $MO_{2}$. Neither species experienced immediate mortality during the exhaustive challenge; although, single instances of delayed mortality were documented for each species. Energetic costs and recovery times could be extrapolated for $C. melanopterus$ via respirometry; sharks were estimated to expend 9.9 kJ kg$^{-1}$ (15% of energy expended on daily swimming) for a single challenge and could require 8.4 h to recover. These data suggest that neonatal $C. melanopterus$ and $N. acutidens$ are resilient to brief gill-net capture durations, but this was under a narrow temperature range. Defining species’ vulnerability to stressors is important for understanding the efficacy of shark conservation tools, including MPAs.

Key words: Bycatch, marine protected areas, oxygen uptake rates, physiological stress response, shark nursery areas, temperature

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Introduction

Marine protected areas (MPA), including shark sanctuaries, can be important conservation tools for protecting threatened shark populations. Indeed, some shark populations face declines worldwide, owing to overexploitation in fisheries (Dulvy et al., 2014). One strategy to potentially reduce the threat of fishing to shark populations is through the creation of MPAs with specific regulations that protect shark populations. For instance, “shark sanctuaries” ban targeted shark fisheries within a country’s exclusive economic zone (EEZ) (Cramp et al., 2018). A general concern regarding protected habitats for sharks and other top predators is that other significant threats, like bycatch or environmental change, are not adequately managed (Ward-Paige and Worm, 2017). Incidental capture, or bycatch, affects shark populations through fishing-induced mortality and negative sub-lethal outcomes (Skomal and Mandelman, 2012; Wilson et al., 2014; Ellis et al., 2017). Climate change is resulting in ocean warming and acidification and can affect shark populations through local extirpation as conditions become too extreme in addition to negative sub-lethal outcomes (Rosa et al., 2017; Payne et al., 2018). Protected shark populations may be inherently at risk of experiencing negative outcomes associated with these stressors because virtually all shark sanctuaries are in the tropics. Here, environmental conditions may border species’ limits to optimal physiological performance and therefore impede resolving stressors (Rummer et al., 2014). Furthermore, populations that rely on shallow coastal waters during key parts of their life histories (e.g. neonates in nursery areas) are already facing quite variable environmental conditions and can also be risk of fishing interactions (Knip et al., 2010). Therefore, developing an understanding of shark populations’ resilience to stressors that they are still expected to face within MPAs can provide valuable information for improving the efficacy of these conservation tools (Chin et al., 2010; Illing and Rummer, 2017).

Neonatal and juvenile shark populations that rely on nearshore habitats may be vulnerable to bycatch. Shallow coastal environments are important for young sharks as nursery areas (Heupel et al., 2007). Alternatively, non-nursery areas can provide stability to young shark populations that typically utilize a diversity of habitats (Yates et al., 2012). While shallow waters may offer young sharks protection from predators, proximity to the coastline increases the probability of fishing interactions (Knip et al., 2010). Specifically, young sharks can be caught as bycatch in artisanal and recreational fisheries. Depending on the type of fishery (e.g. hook-and-line or net fishing), different species have varying susceptibilities to lethal or sub-lethal outcomes (Dapp et al., 2016). Capture is generally associated with vigorous escape attempts that can drive a physiological stress response (Brooks et al., 2012; Guida et al., 2016; Gallagher et al., 2017). The stress response is generally characterized by a release of hormones (e.g. adrenaline and noradrenaline), the accumulation of by-products of anaerobic metabolism (e.g. lactate) that drive declines in tissue pH, and resultant osmotic and ion imbalances (Skomal and Mandelman, 2012). Capture is also associated with an increased rate of energy expenditure (Bouyoucos et al., 2017b). Physiological stress and depleted energy reserves following fisheries capture can even contribute to exhaustion-induced mortality or post-release predation (Danylchuk et al., 2014; Lennox et al., 2017). Additional lethal stressors can be problematic because young sharks may already experience high mortality rates during their first year of life (Gruber et al., 2001; Heupel and Simpford, 2002).

Young shark populations in shallow coastal habitats must also contend with stressors associated with variable environmental conditions. Shallow coastal environments can be prone to seasonal and tidal variations in environmental conditions, such as temperature, salinity and dissolved oxygen concentrations that affect the abundance and distribution of various species of sharks (Knip et al., 2010; Schlaff et al., 2014; Oh et al., 2017b). Changes in abundance and distribution may be partially attributed to physiological costs associated with variable environmental conditions. Increases in temperature decrease oxygen’s solubility in water. Oxygen uptake rates (a proxy for metabolic rate) also increase, along with concomitant decreases in haemoglobin–oxygen (Hb–O2) affinity (Bernal et al., 2012, 2018). In addition, parameters associated with sharks’ stress response to capture vary with temperature, such that capture at high temperatures can be fatal for some species (Hoffmayer et al., 2012; Danylchuk et al., 2014; Guida et al., 2016). While sharks may attempt to maintain a preferred body temperature or boundaries to their critical thermal limits, life history stages (e.g. neonates) that derive specific benefits from confined habitats (e.g. predator avoidance within nursery areas) must be able to tolerate local conditions (Knip et al., 2010; Payne et al., 2016, 2018). However, sharks in the tropics are expected to be adapted to a narrow range of temperatures and, therefore, to have a low tolerance for variable environmental temperature conditions (Rummer and Munday, 2017). While there is a paucity of data on thermal tolerance limits for sharks, it is likely that sharks within coastal habitats in tropical latitudes may already be living close to their thermal tolerance limits (Rummer et al., 2014).

The purpose of this study was to characterize various components of the stress response of neonatal reef sharks following an exhaustive challenge. Specifically, we sought to measure the physiological status of neonatal blacktip reef sharks (Carcharhinus melanopterus) and sicklefin lemon sharks (Negaprion acutidens) at multiple points in time following in situ gill-net capture. The objectives of this study were to (1) characterize physiological responses in neonatal reef sharks following capture, (2) predict the effect of changes in environmental temperatures on physiological status, (3) assess the differential vulnerability of co-occurring neonatal reef shark species to stress-induced physiological impairment and (4) estimate the energetic cost of an
exhaustive challenge in the context of routine energy requirements. Studies of this nature are necessary for understanding whether stressors hold lethal or sub-lethal consequences under predictable environmental conditions in important habitats like shark nursery areas. As such, these data will have management applications to better support conservation initiatives for reef sharks (Illing and Rummer, 2017).

Materials and methods

All experiments were approved by James Cook University Animal Ethics Committee protocol A2089. Research on sharks in French Polynesia was approved under Arrêté No 9524 issued by the Ministère de la Promotion des Langues, de la Culture, de la Communication et de l’Environnement of the French Polynesian government on 30 October 2015.

Study site, animal collection and husbandry

Fieldwork was conducted from shore around Moorea, French Polynesia (17°30’S, 149°51’W), where targeted shark fishing in the country’s EEZ has been banned since 2012 (Ward-Paige and Worm, 2017). Newborn *C. melanopterus* and *N. acutidens* are abundant during parturition months from September through February (Mourier and Planes, 2013; Mourier et al., 2013a, b). Sharks were collected during November and December 2016 using monofilament gill-nets (50.0 m × 1.5 m, 5.0 cm mesh) fished at dusk (17:00–20:00). Captured sharks were immediately identified and removed from the net in under five minutes. Prior to release, biological data (total length, mass, and sex) were collected from all individuals subjected to one of four treatments. One group of laboratory-acclimated sharks was phlebotomized in a quiescent state after 2–4 weeks in captivity and a 48-hour fasting period to generate minimally-stressed values (“baseline” treatment). A second group of sharks was phlebotomized immediately following the exhaustive challenge in the field (“immediate” treatment). The third group of sharks faced the same exhaustive challenge and was retained in flow-through mesh bags in the field for 3 h before phlebotomy (“three-hour” treatment). A final group of sharks was sampled after 3 h in a respirometry chamber that was used to estimate energetic costs and recovery times for the exhaustive challenge (“respirometry” treatment). All blood samples were processed immediately following phlebotomy.

Sharks were phlebotomized via caudal puncture using heparin-rinsed 23.0 gauge 3.8 cm needles. Five parameters were measured using point-of-care analytical devices: blood glucose concentration (mmol l⁻¹), blood lactate concentration (mmol l⁻¹), blood pH, haemoglobin concentration (Hb); g dl⁻¹), and haematocrit (Hct). Blood was first transferred from syringes directly to 70-μl microcapillary tubes that were run in parallel in a microhaematocrit centrifuge (ZIPocrit, LW Scientific, Lawrenceville, GA, USA) for 2 min at 4400 g (Danylchuk et al., 2014). Whole blood glucose and lactate concentrations were measured with 10 μl samples of whole blood using an Accutrend Plus (Roche Diagnostics Ltd, Rotkreuz, Switzerland), with ranges of 1.1–33.3 mmol l⁻¹ and 0.8–22.0 mmol l⁻¹, respectively (Butcher et al., 2015). Readings that were outside the measurement range were reported as the value of the upper or lower device limit for statistical analyses. Haemoglobin concentration was measured with a Hemocue Hb 201 System (Australia Pty Ltd, Victoria, Australia) using 10 μl of whole blood, and was corrected using a calibration equation generated for fish that has previously been applied to sharks (Clark et al., 2008; Heinrich et al., 2014). Haemoglobin concentration was then converted to tetramer Hb concentration (Hb₄, in mmol l⁻¹).
using conversions generated for tropical reef species in order to calculate mean cell haemoglobin concentration (MCHC; mmol L\(^{-1}\)), as Hb\(\text{blood}\) divided by Hct (Rummer et al., 2013; Heinrich et al., 2014). Blood pH was measured using a HI98165 pH meter (Hanna Instruments, Victoria, Australia), and raw pH values were converted to values derived from the conventional i-STAT system using a correction formula generated for juvenile lemon sharks (N. brevirostris) at 25.6–31.3°C (Talwar et al., 2017).

**Estimating energetic costs and recovery**

To estimate costs of an exhaustive challenge and recovery times, individuals from another subset of sharks (C. melanopterus) were transferred to individual field respirometry chambers immediately after capture and air exposure so that oxygen uptake rates (\(\text{MO}_2\), in mg O\(_2\) kg\(^{-1}\) h\(^{-1}\)) could be measured over 3 h. To do this, two respirometry chambers (24.0 cm diameter and 70.0 cm long, 32.0 l volume including tubing) were submerged in a 400.0 l circular pool positioned ~3 m from the shoreline. Water in the pool was continuously aerated, and was supplied at a rate of 4800.0 l h\(^{-1}\) from a pump approximately 5.0 m offshore in at least 0.3 m of water. Respirometry chambers were configured for intermittent-flow respirometry with 2500.0 l h\(^{-1}\) flush and recirculating pumps (Rummer et al., 2016; Svendsen et al., 2016). Dissolved oxygen concentration (DO, in mg L\(^{-1}\)) was measured every second with fibre optic probes that were mounted within chambers and connected to a Firesting Optical Oxygen Meter (Pyroscience, Aachen, Germany). Probes were calibrated to fully-aerated freshwater (100.0% saturation) before each use and to 0.0% saturation with sodium sulphite as needed. Flush pumps were manually operated to cycle flush (9.1 ± 6.5 min SD) and measurement periods (11.5 ± 6.1 min) such that DO remained above 80.0% air saturation. The timing of cycles was determined by watching DO in real-time on a laptop computer. Sharks were placed into the chambers immediately upon capture, and therefore the time from the onset of capture to the beginning of the first measurement was 4.4 ± 1.2 min (i.e. the length of the exhaustive challenge). Each field respirometry trial consisted of 6–12 measurement periods over 3 h. Then, immediately after removal from respirometry chambers, all sharks were phlebotomized to determine whether undergoing respirometry influenced the stress response (the “respirometry” treatment).

Oxygen uptake rates were estimated by first calculating rates of DO decline every 30 s during each measurement using LabChart (7.3.8, ADInstruments, Dunedin, New Zealand). Specifically, \(\text{MO}_2\) was calculated as \(\text{MO}_2 = \frac{V_{\text{Resp}} \cdot S}{M}\) where \(S\) is the slope of the linear decline in DO (in mg O\(_2\) L\(^{-1}\) s\(^{-1}\)), \(V_{\text{Resp}}\) is the volume of the respirometer minus the shark’s volume (in l), and \(M\) is the mass of the fish (in kg). Background respiration was accounted for by modelling the linear increase in background \(\text{MO}_2\), measured before and after each trial in chambers without fish, and subtracting proportional background \(\text{MO}_2\) from each \(\text{MO}_2\) measurement (Rodgers et al., 2016; Rummer et al., 2016). The highest \(\text{MO}_2\) during each measurement period was selected, and these values were fit with an exponential decay curve (recovery curve). The highest \(\text{MO}_2\) value for each shark was recorded as its maximum \(\text{MO}_2\) (\(\text{MO}_{2\text{Max}}\)).

Oxygen uptake rates of minimally-stressed, resting sharks (C. melanopterus and N. acutidens; the same animals used in for the “baseline” treatment) were measured in the laboratory. The same respirometry chambers described above were placed in holding tanks, and flush pumps were automated with a custom-built data acquisition system and software (National Instruments, Austin, Texas, USA). Flush pumps were automated to shut off for 5 min every 12 min for C. melanopterus, and 5 min every 15 min for N. acutidens, yielding at least 120 measurements for C. melanopterus and at least 96 measurements for N. acutidens over 24 h. Shorter measurement periods and longer flush periods were deemed necessary for N. acutidens because all individuals were larger than the C. melanopterus used for this study and had higher \(\text{MO}_2\). One slope (\(S\)) was calculated for each measurement. Minimum (\(\text{MO}_{2\text{Min}}\)) was calculated as the mean of the lowest 10% of \(\text{MO}_2\) values, excluding values outside of the mean ± 2 SD (Clark et al., 2013).

**Statistical and data analyses**

Underlying physiological responses were characterized by comparing values of physiological parameters over time after an exhaustive challenge, and against baseline values. The influence of temperature on physiological status (i.e. values of physiological and oxygen uptake parameters) was assessed by including temperature as a covariate in models. Physiological parameters (i.e. blood glucose and lactate concentrations, blood pH, [Hb], Hct and MCHC) were fit with linear models to observe variation in responses with treatment (fixed effect), temperature and mass (covariates) for both species. For C. melanopterus, the factor “treatment” had four levels (i.e. baseline, immediate, three-hour, and respirometry). It was not possible to catch comparable numbers of N. acutidens, and as a result the factor “treatment” only had three levels (i.e. baseline, immediate and 3-h). All possible interactions (two-way and three-way) were included in these models for C. melanopterus. Samples sizes were too small to include interactions for N. acutidens. Post hoc multiple comparisons were made with Tukey’s honest significant difference (HSD) tests. Models were validated with Q–Q plots of model residuals, and by plotting residuals against treatment and fitted values (Zuur et al., 2007). For all tests, the acceptable Type I error rate (\(\alpha\)) was 0.05, and all analyses were conducted using the R Stats Package (R Core Team, 2016).

Recovery times and costs were estimated for C. melanopterus using respirometry data. The mean value of \(\text{MO}_{2\text{Min}}\) that was derived from the laboratory was used as a baseline
for estimating the excess post-exercise oxygen consumption (EPOC, in mg O$_2$ kg$^{-1}$) of individual sharks from field respirometry. Recovery times were estimated for individual sharks as the time when the recovery curve intersected the upper 95% confidence interval limit of MO$_{2Min}$ (Bouyoucos et al., 2017a). Excess post-exercise oxygen consumption, which represents the cost of recovery from exhaustive activity (Gaesser and Brooks, 1984), was calculated as the area bound by individual sharks’ recovery curves, MO$_{2Min}$, the time of the first MO$_2$ measurement, and the time of recovery (Bouyoucos et al., 2017a). Oxygen uptake parameters (i.e. MO$_{2Min}$, MO$_{2Max}$, EPOC and recovery time) were fit with linear models to observe variation with temperature and mass, including interactions.

**Results**

**Quantifying physiological responses**

Morphometric data for *C. melanopterus* are presented in Table 1. Sharks exhibited significant changes in blood glucose and lactate concentrations as well as blood pH across treatments (Supplementary Table S1). Blood glucose concentrations at three hours were higher than baseline values (Tukey’s HSD, $t = 4.387, P < 0.001$) and values for immediately-sampled sharks (Tukey’s HSD, $t = 4.062, P = 0.002$) (Fig. 1a). Blood glucose concentrations also had a positive linear relationship with temperature (Linear regression, $R^2 = 0.27, F_{1, 25} = 10.82, P = 0.003$; 27.9–30.9 °C; Fig. 2) across treatments (Supplementary Table S1). Baseline and immediately-sampled values for blood lactate concentrations did not differ (Tukey’s HSD, $t = 1.436, P = 0.489$), and values after 3 h in recovery bags and respirometry chambers were not different (Tukey’s HSD, $t = -0.639, P = 0.918$). Blood lactate concentrations were at least 14-fold higher 3-h post-capture relative to baseline and immediately-sampled values (Tukey’s HSD, $P < 0.001$) (Fig. 1b). Lastly, blood pH was uniformly reduced across all treatments relative to baseline values (Tukey’s HSD, $P < 0.001$) (Fig. 1c). No significant differences in [Hb] (4.48 ± 0.77 g dl$^{-1}$), Hct (0.17 ± 0.03) or MCHC (4.26 ± 0.17) were detected.

Morphometric data for *N. acutidens* are presented in Table 1. Differences between treatments were only detected for blood lactate concentration and blood pH (Supplementary Table S2). Blood lactate concentrations were at least 6-fold higher for *N. acutidens* three hours after capture relative to baseline (Tukey’s HSD, $t = 9.128, P < 0.001$) and immediately-sampled values (Tukey’s HSD, $t = 8.407, P < 0.001$), which were not different (Tukey’s HSD, $t = -0.679, P = 0.269$) (Fig. 3b). In addition, blood pH was significantly reduced for sharks sampled immediately (Tukey’s HSD, $t = -3.153, P = 0.014$) and 3 h post-capture (Tukey’s HSD, $t = -2.940, P = 0.037$) relative to baseline pH (Fig. 3c). Blood pH values sampled immediately or three hours post-capture were not different (Tukey’s HSD, $t = 0.185, P = 0.981$).

There were no significant differences across treatments for blood glucose concentration $(5.22 ± 0.88$ mmol l$^{-1}$), [Hb] $(5.15 ± 0.83$ g dl$^{-1}$), Hct $(0.19 ± 0.03)$ or MCHC $(4.26 ± 0.77$ mmol l$^{-1}$) for *N. acutidens*. No physiological parameter varied with mass (Supplementary Table S2), and [Hb] had a positive linear relationship with temperature (Linear regression, $R^2 = 0.37, F_{1, 11} = 7.59, P = 0.016$; 29.5–30.9°C; Fig. 4).

**Estimating energetic costs and recovery**

Mean MO$_{2Max}$ was $322.91 ± 72.93$ mg O$_2$ kg$^{-1}$ h$^{-1}$, and EPOC was $703.72 ± 361.53$ mg O$_2$ kg$^{-1}$ at 30.06 ± 1.28°C (Fig. 5a). From laboratory measurement for *C. melanopterus*, MO$_{2Min}$ was $100.92 ± 11.30$ mg O$_2$ kg$^{-1}$ h$^{-1}$ at 29.66 ± 0.69°C, and estimated aerobic scope (AS = MO$_{2Max}$ − MO$_{2Min}$) was $221.98$ mg O$_2$ kg$^{-1}$ h$^{-1}$. No shark had recovery curves that intersected the upper 95% CI limit of MO$_{2Min}$ in under 3 h, and extrapolated recovery times ranged from 3.1 to 19.8 h (8.42 ± 5.78 h). None of the oxygen uptake parameters varied with temperature, mass, or their interaction (Supplementary Table S1). Lastly, only three *N. acutidens* were brought to the CRIOBE to generate baseline values for this species, where MO$_{2Min}$ was determined to be $139.95 ± 12.07$ mg O$_2$ kg$^{-1}$ h$^{-1}$.

**Observed mortality**

Immediate mortality was 0% for both species, but delayed mortality was observed for both *C. melanopterus* and *N. acutidens*. A single *C. melanopterus*, which was caught at 32.33°C, was moribund upon release from its field respirometry chamber. Oxygen uptake data suggest that this animal experienced aerobic failure at ~1.5 h following the exhaustive challenge (Fig. 5b). Including this animal, delayed mortality for *C. melanopterus* was 5.9% (1/17). The body of one *N. acutidens* was recovered the day after release from a recovery bag, suggesting that delayed mortality for this species was 25.0% (1/4); this animal was caught at 29.75°C.

**Discussion**

Neonatal *C. melanopterus* and *N. acutidens* were still stressed 3 h after facing an exhaustive challenge. Values for blood glucose, lactate and pH taken 3 h after the exhaustive challenge deviated from baseline values for both species (except blood glucose concentrations in *N. acutidens*). These physiological responses are characteristic of the esmolbranch secondary stress response (Skomal and Mandelman, 2012; Wilson et al., 2014). Vigorous attempts by sharks to escape fishing gear are generally supported by anaerobic metabolic pathways that are partially characterized by increases in blood glucose and lactate concentrations and a resultant drop in blood pH (Guida et al., 2016; Gallagher et al., 2017; Bouyoucos et al., 2017b). Furthermore, sharks entangled in gill nets may not be able to ventilate, thereby driving further declines in blood pH by restricting carbon dioxide offloading, and relying on anaerobic metabolic pathways while oxygen uptake is impeded (Dapp et al., 2016).
Even if a shark that is restrained in a net can actively ventilate, for example via buccal pumping, this strategy could be a far less efficient method for gas exchange and may actually exacerbate the stress response (Parsons and Carlson, 1998; Brooks et al., 2011). Many shark species also lack mechanisms to modulate haematological parameters related to...
improving oxygen delivery during a stress response (Brill and Lai, 2015). Previous studies have also documented that sharks facing brief exhaustive challenges can take over 3 h to recover (Brooks et al., 2011, 2012). While we documented neonatal sharks experiencing various aspects of the stress response, it was beyond the scope of this study to determine exactly how detrimental the levels of stress experienced were (i.e., changes in recovery times or risk of experiencing mortality). Interestingly, recapture rates for both species have been relatively high within a given parturition season (~15–30%), but low recapture rates from over five years of annual surveys around Moorea suggest that natural mortality (e.g., starvation or predation) is quite high among these populations (S.P. unpublished results). Size classes between neonates and adults are notably absent from gill-net and hook-and-line surveys; although, variable habitat use or size-selective gears may appear to suggest high juvenile mortality in the absence of natural mortality rate estimates for this population (Mourier et al., 2013b). Around Moorea, exhaustive challenges are expected in the form of artisanal and recreational fisheries bycatch and predator-prey interactions (Chin et al., 2015; Mourier et al., 2017; Thiault et al., 2017). Although French Polynesia is a shark sanctuary, implementing and enforcing management strategies to mitigate fishing pressure during parturition months could reduce neonatal sharks’ chance of facing exhaustive challenges (i.e., fishing capture).

Figure 2: Relationship between temperature and physiological status (blood glucose concentrations) for blacktip reef sharks (Carcharhinus melanopterus).

Evidence of an effect of temperature on the physiological status of C. melanopterus and N. acutidens was equivocal over a narrow, albeit ecologically relevant temperature range. Blood glucose concentrations doubled, on average, over a 3.0°C range for C. melanopterus (27.9–30.9°C) and [Hb] increased with temperature by ~23% over a 1.4°C range (29.5–30.9°C) in N. acutidens. Some markers of physiological status may respond to changing environmental temperatures for elasmobranchs because of temperature’s influence on the metabolic rates of ectothermic organisms (Hoffmayer et al., 2012; Guida et al., 2016). Conversely, temperature-associated changes in blood glucose concentrations of C. melanopterus could simply reflect increased activity levels of sharks in warmer water, as opposed to a temperature-mediated metabolic response (Whitney et al., 2016; Payne et al., 2016, 2018). Increases in [Hb] of N. acutidens with increasing temperature may be a compensatory mechanism as Hb–O2 affinity decreases (Bernal et al., 2018). Alternatively, the apparent correlation between [Hb] and temperature may have been spurious, as changes in [Hb] ultimately did not result in variation in MCHC or Hct. However, N. acutidens in warmer waters may have had smaller red blood cells (RBCs) or immature RBCs in greater circulation but with similar [Hb] to sharks at cooler temperatures that would appear as an increase in [Hb] without affecting other haematological variables. No other physiological or oxygen uptake parameters that were measured displayed variations with temperature. Metabolic compensation, where an organism maintains consistent MO2 with temperature acclimation, has not been documented for elasmobranchs (Tullis and Baillie, 2005). It is likely, however, that, even for seasonally-acclimated elasmobranchs, variations in temperature exceeding 3.0°C may be necessary to elicit an observable response (Carlson and Parsons, 1999; Neer et al., 2006). Moorea’s neonatal shark populations face summer temperatures that average 30°C during parturition months, daily variations of up to 8°C, and extreme temperatures ranging 26–36°C (J.L.R. unpublished results). For juvenile sharks, facing an exhaustive challenge in shallow coastal waters when temperatures are high can be lethal (Danychuk et al., 2014). The only C. melanopterus to die in this study was, coincidentally, captured at >32°C, but we could not confirm whether this single mortality was related to temperature. Both C. melanopterus and N. acutidens exhibit some degree of philopatry to natal areas around Moorea and elsewhere, such that extreme temperature events in these potential nursery areas could put neonates at risk of mortality after facing exhaustive challenges (Mourier and Planes, 2013; Mourier et al., 2013b; Oh et al., 2017a). Without controlled studies to investigate the effect of temperature on reef sharks’ resilience to stress, it is unclear whether thermal stressors like ocean warming brought on by climate change could be problematic for neonatal sharks in tropical nearshore habitats.

Physiological status before and after the exhaustive challenge was species-specific. Notably, N. acutidens exhibited a larger difference in blood pH relative to baseline values, did not exhibit variation in blood glucose concentrations across the samples, and had high baseline lactate concentrations compared to C. melanopterus. Overall trends in blood
lactate concentrations, blood pH and haematological parameters, however, were similar for both species and consistent with what has been reported for other elasmobranchs (Lowe et al., 1995; Richards et al., 2003; Brill et al., 2008). The higher $M\dot{O}_2$Min observed for $N$. acutidens could explain the larger drop in blood pH ($\Delta$ pH = 0.44) following the exhaustive challenge relative to $C$. melanopterus ($\Delta$ pH = 0.28). It is hypothesised that the magnitude and severity of a stress response is related to $M\dot{O}_2$ for elasmobranchs (Skomal and Mandelman, 2012). While it was not possible to calculate AS for both species, Carcharhinus melanopterus are generally regarded as stronger aerobic swimmers than $N$. acutidens, which are less active and known to rest (Baldwin and Wells, 1990; Wells et al., 1992). High blood glucose concentrations and resting blood lactate concentrations in $N$. acutidens could be a result of this species recruiting anaerobic metabolism to support bouts of swimming that are interspersed with periods of resting (Piiper et al., 1977). Blood-oxygen transport properties ([Hb], Hct, MCHC) were not affected by exercise, and were similar between the two species, as has been previously reported (Wells and Baldwin, 1990; Wells et al., 1992). In addition, juvenile $C$. melanopterus and $N$. acutidens from Heron Island (on the Great Barrier Reef) were reported to exhibit similarly pH-insensitive haemoglobins, suggesting that Hb–O₂ affinity and oxygen transport are not greatly affected by an acidosis for these species (Baldwin and Wells, 1990; Wells et al., 1992). Taken together, these data suggest that each species has a unique physiological response to stress in relation to their behaviour and aerobic capacity, where $N$. acutidens may have experienced a more intense stress response owing to a potential greater reliance on anaerobic metabolism to support activity. It would be informative to characterize the full physiological response from initiation to resolution (i.e. recovery or mortality) to determine how differently these two species respond to capture.

Figure 3: Indicators of the stress response in juvenile sicklefin lemon sharks (Negaprion acutidens) following an exhaustive challenge in situ. Baseline values were taken from quiescent, fasted sharks (“baseline”). Other sharks were phlebotomized immediately following exhaustive gill-net capture (“immediate”) or after 3 h in a recovery bag (“three-hour”). Differing letters denote statistically significant differences. Abbreviation: mean cell haemoglobin concentration (MCHC).
There was no observable immediate mortality for *C. melanopterus* and *N. acutidens*. However, delayed mortality rates were higher for *N. acutidens* (25%) than *C. melanopterus* (5.9%), although this study’s experimental design precluded quantification of robust mortality rates for either species. Adult *C. melanopterus* around Moorea appear to be quite resilient to hook-and-line capture (Mourier et al., 2017), and neonate and juvenile *C. melanopterus* and *N. acutidens* both exhibited near 0% mortality following gill-net and hook-and-line capture in the Mangrove Bay Sanctuary Zone on Ningaloo Reef, Australia (Oh et al., 2017a). In contrast, however, another study out of Western Australia reported that, when facing unspecified capture durations in gill-nets, juvenile and adult *C. melanopterus* were more susceptible to immediate mortality than *N. acutidens* (Dapp et al., 2017). Local adaptation to environmental conditions at the population level may influence these contrasting trends (Eliason et al., 2011; Di Santo, 2016); although, temperature data were not reported from the Western Australia study (Dapp et al., 2017). The model generated by Dapp et al. (2017) to estimate immediate mortality of *C. melanopterus* would have predicted 100% mortality for sharks in our study using only total length as a predictor. It is possible that differences in the duration of capture led this study to conclude that immediate mortality was 0%, whereas difficulty in identifying capture events by Dapp et al. (2017) could have allowed for sufficiently long capture durations and more realistic immediate mortality estimates. Alternatively, stress resulting from this study’s shorter capture durations may simply not have been fatal (Oh et al., 2017a). Differences in mortality estimates for *N. acutidens* may have been related to size; although, sizes of *N. acutidens* were not reported by Dapp et al. (2017). It, therefore, seems likely that these apparently contrasting findings resulted from differences in the nature and duration of the stressor (e.g. capture duration, supplementing air exposure, local environmental conditions, etc.).

This study’s exhaustive challenge was associated with a large energetic cost and long recovery for *C. melanopterus*. The mean estimated EPOC was 703.72 mg O₂ kg⁻¹ h⁻¹, and recovery was estimated to take 8.42 h. Comparatively, chasing juvenile lemon sharks (*N. brevirostris*) to exhaustion without air exposure resulted in an EPOC of 154.10 mg O₂ kg⁻¹ and 5.40 h of recovery at 30°C (Bouyoucos et al., 2017b). Gill-net capture and air exposure may result in a larger EPOC than exhaustive chasing because oxygen uptake is impeded, such that recovery cannot begin until oxygen uptake is resumed; a chased fish in water can still meet some of its energy demand aerobically and even begin to recover. The EPOC estimated for *C. melanopterus* is much larger than measured for other elasmobranchs (Brett and Blackburn, 1978; Bouyoucos et al., 2017a, b), but it is similar to values reported for a tropical coral reef fish (*Pomacentris ambioensis*) at comparable temperatures (28–29°C) (Killen et al., 2014). However, recovery times for *P. ambioensis* were under one hour, which likely relates to this species’ impressive AS that is almost ten times that of *C. melanopterus* (Killen et al., 2014). Assuming that MO₂ scales with swimming speeds similarly among carcharhinid sharks (Carlson et al., 2004), a routine swimming MO₂ of 195.87 mg O₂ kg⁻¹ h⁻¹ can be estimated for *C. melanopterus* using a power-performance slope of 0.36, a routine swimming speed of 0.80 body lengths s⁻¹ for captive *C. melanopterus*, and this study’s estimate of MO₂Min (*Webb and Keyes, 1982; Bushnell et al., 1989*). Applying an oxygen equivalent of 14.14 J mg O₂⁻¹, *C. melanopterus* would have a daily metabolic rate of 66.47 kJ kg⁻¹ d⁻¹ for swimming alone, and an EPOC of 9.95 kJ kg⁻¹ from an exhaustive challenge would increase daily energy expenditure for swimming by 14.9% (Elliot and Davison, 1975). Around Moorea, neonatal *C. melanopterus* and (*N. acutidens*) must quickly transition from relying on endogenous fuel stores to energy acquired through hunting (Matich et al., 2015). Energetically costly one-off exhaustive challenges, like incidental capture, could precede starvation in neonatal sharks, especially for populations with high natural mortality.

In conclusion, within a narrow range of temperatures, neonatal *C. melanopterus* and *N. acutidens* are resilient to brief durations of gill-net capture. However, we are unaware of these species’ physiological resilience to longer durations of capture with different gear types, to longer periods of air exposure, or at temperatures beyond 28–31°C. As such, artisanal and recreational fisheries bycatch mortality could still pose a threat to Moorea’s neonate and juvenile shark populations. Indeed, longer gill-net capture durations could be...
Moving forward, studies are needed to define environmental conditions that limit physiological performance and to fully characterize recovery following a challenge. Furthermore, defining changes in routine energy requirements and reserves of neonates exposed to stressors in relation to the quality and availability of shelter and prey will be important for estimating sharks’ likelihood of facing predation and starvation, respectively. Together, these data have the potential to improve our understanding of how anthropogenic and environmental stressors affect the survivorship of neonate and juvenile reef sharks in important habitats like shark nursery areas. Understanding the vulnerability of shark populations to manageable stressors, like fishing pressure, is an important step toward improving the efficacy of MPAs as conservation tools for sharks, globally.

**Supplementary material**

Supplementary material is available at *Conservation Physiology* online.

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