Compensatory proteome adjustments imply tissue-specific structural and metabolic reorganization following episodic hypoxia or anoxia in the epaulette shark (Hemiscyllium ocellatum)

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Dowd WW, Renshaw GMC, Cech JJ Jr, Kültz D. Compensatory proteome adjustments imply tissue-specific structural and metabolic reorganization following episodic hypoxia or anoxia in the epaulette shark (Hemiscyllium ocellatum). Physiol Genomics 42: 93–114, 2010. First published April 6, 2010; doi:10.1152/physiolgenomics.00176.2009.—The epaulette shark (Hemiscyllium ocellatum) represents an ancestral vertebrate model of episodic hypoxia and anoxia tolerance at tropical temperatures. We used two-dimensional gel electrophoresis and mass spectrometry-based proteomics approaches, combined with a suite of physiological measures, to characterize this species’ responses to 1) one episode of anoxia plus normoxic recovery, 2) one episode of severe hypoxia plus recovery, or 3) two episodes of severe hypoxia plus recovery. We examined these responses in the cerebellum and rectal gland, two tissues with high ATP requirements. Sharks maintained plasma ionic homeostasis following all treatments, and activ-

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metabolic processes during and after oxygen limitation, the compensatory strategies of other tissues in this and other hypoxia- and anoxia-tolerant species have received far less attention.

Just as critical as the need to survive periods of oxygen limitation is the ability to cope with subsequent rapid reoxygenation. The return to normoxic conditions in mammalian systems coincides with bursts in the generation of reactive oxygen species (ROS) or reactive nitrogen species (RNS) (i.e., oxidative stress). The associated macromolecular damage and cell death (necrosis and/or apoptosis) is commonly referred to as reoxygenation/reperfusion injury (reviewed in Ref. 54). Recent work has demonstrated that elevated ROS formation may occur during hypoxia as well (54, 56, 57). Thus, the epaulette shark’s survival of episodic hypoxia on the reef platform (and episodic anoxia in the laboratory) may in some ways be more relevant than long-term anoxia tolerance to biomedical interest in ischemia/reperfusion, due to this species’ apparent ability to repeatedly suppress or minimize reoxygenation injury.

Due to the repeated, episodic nature of hypoxia in the epaulette shark’s habitat, the reef platform has been referred to as a natural “hypoxic preconditioning environment” (71). By the term preconditioning (or conditioning hormesis, 9) we refer to the common phenomenon in which exposure to an initial stress elicits physiological and biochemical responses (the preconditioned phenotype) that offer protection against a subsequent, more severe stress (the preconditioning outcome). This preconditioning outcome typically manifests in the form of reduced tissue or molecular damage and/or increased survival. Although definitive evidence for a preconditioning outcome as defined above is lacking in the epaulette shark, this may be moot given that this species likely represents a “constitutively preconditioned” (sensu 66) model of hypoxia and anoxia tolerance. Accordingly, epaulette sharks raised in captivity and thus naïve to oxygen limitation are also tolerant of episodic anoxia (13).

Clearly some of the pathways that prevent neuronal and organismal death after several hours of severe hypoxia or anoxia at tropical temperatures, as well as those conferring the ability to suppress reoxygenation injury, must be constitutively expressed in this species. However, novel or enhanced protective mechanisms that confer increased damage resistance may also be induced by preexposure to oxygen limitation. The relative costs and benefits of these two strategies should depend on the degree, frequency, and predictability of low oxygen conditions experienced in the wild relative to the costs of synthesizing, stabilizing, and degrading the necessary proteins and repairing damage. Several lines of evidence suggest that a preconditioned phenotype is activated by hypoxic pre-exposure in the epaulette shark. Laboratory exposure to multiple hypoxic episodes reduces the organismal aerobic metabolic rate and reduces the critical oxygen tension for maintenance of aerobic metabolism (71, 89). Within the epaulette shark brain, the same repeated hypoxia protocol promotes neuronal aerobic hypometabolism via a decrease in abundance of the tricarboxylic acid cycle enzyme citrate synthase (68) and results in increased localized concentrations of the inhibitory neurotransmitter GABA, particularly in nuclei of the brainstem and cerebellum (67). These adjustments may reduce both ATP turnover and excitatory stimulation and thus provide neuroprotection. Yet, we possess a minimal understanding of the molecular mechanisms underlying these and other responses to oxygen limitation in the epaulette shark brain and other tissues. Given the previous evidence for prolonged preconditioning in field-acclimatized epaulette sharks (and in mammalian models, 4, 99), the periodic and incremental nature of hypoxia on reef platforms due to its dependence on the lunar cycle, and the likely involvement of protein synthesis in late ischemic preconditioning (87), we anticipated that at least some of the epaulette shark’s compensatory response mechanisms are inducible and would be detectable after reoxygenation (e.g., 117). Therefore, in this study we exposed epaulette sharks to controlled anoxic, hypoxic, and repeated hypoxic treatments plus 2 or 24 h of normoxic recovery to assess their responses to no vs. low oxygen and to determine whether these responses vary between one and more than one hypoxic episodes. Because proteins perform the majority of cell functions, we used two-dimensional gel electrophoresis and mass spectrometry-based proteomics approaches to quantify molecular responses to oxygen limitation. Measures at the proteome level integrate changes in transcription, mRNA stability, and protein turnover in response to stressors (5, 19, 46). We focused on the 24 h recovery time point for our proteomics analyses for three reasons: 1) reoxygenation induces as or more severe damage as the period of oxygen limitation (5, 56); 2) compensatory stress response mechanisms may be more potent during the recovery period than during the actual stress period (5, 49); and 3) hypoxic episodes are naturally spaced 24 h apart on the reef platform. Thus, our proteomic data represent the phenotype at the time when the next stressful episode would likely be encountered. This 24 h lag also coincides with the onset of the delayed phase of ischemic preconditioning in mammalian models (e.g., 4).

Fig. 1. Profiles of dissolved oxygen (black line) and water temperature (dark gray line) in relation to tidal height (dotted line) on the Heron Island reef platform for both neap tides (from flood tide through nocturnal low tide on July 22, 2007; left) and spring tides (7 day plot from July 28 to August 3, 2007; right). Severe reductions in dissolved oxygen coincide with nocturnal, spring low tides. Large x-axis ticks indicate midnight; smaller ticks represent 6 h intervals. Light gray circle denotes the time at which the field blood samples were collected. Note that these measurements were taken in Austral winter; oxygen limitation is likely to be even more severe at higher summer temperatures.
We first examined responses to oxygen limitation in the cerebellum, which controls critical organismal functions (including reflex coordination) and is highly vulnerable to oxygen limitation in most species. Cerebellar activity appears to be downregulated during anoxia in the epaulette shark, as evidenced by the reversible loss of a righting reflex (85). We anticipated that adaptive molecular responses to oxygen limitation might occur in pathways related to energy metabolism, redox balance and ROS neutralization, protein stabilization and repair, and apoptosis/cell cycle. Thus, our expectation was that patterns of protein abundance in these pathways would vary depending on the hypoxic or anoxic exposure regime. We then asked whether oxygen limitation elicits similar or divergent responses in another tissue that performs a vital organismal function. The elasmobranch rectal gland uses significant amounts of aerobically derived ATP to fuel the high Na+/K+ ATPase activity that drives excretion of NaCl for ionoregulation (22), but it may also have substantial anaerobic capacity (110).

In concert with the proteomics data from both cerebellum and rectal gland, we assessed a number of indicators of metabolic and physiological status in the same individuals and tissues. At the organismal level this included measures of ventilation rate during episodes of oxygen limitation, circulating plasma glucose (a favored metabolic fuel of elasmobranch brain and rectal gland), and plasma lactate (the primary end product of anaerobic glycolysis). We anticipated that there might be some breakdown or selective downregulation of the rectal gland’s salt and extracellular volume homeostatic functions (assessed here via measurements of plasma ion concentrations) with oxygen limitation. At the biochemical level, we measured the activity of Na+/K+-ATPase, a major ATP sink in both tissues (albeit in support of different functions), to determine whether oxygen limitation and recovery elicits a decrease in ion pumping capacity. The combined activities of the apoptotic effector enzymes caspases 3 and 7 were used as an indicator of apoptosis, the programmed cell death that may result from exposure to excessive stress (46). Finally, we measured levels of glutathione to determine whether epaulette sharks have high constitutive expression of this ROS scavenger or whether the total glutathione pool could be regulated as part of the response to oxygen limitation. Based on these integrative data, we outline some of the common, treatment-specific, and tissue-specific mechanisms involved in the epaulette shark’s response to episodic anoxia and hypoxia plus reoxygenation.

MATERIALS AND METHODS

Animal Capture and Maintenance

Epaulette sharks (n = 42, mean 356 ± 21 g, range 69–646 g) were collected by hand at low tide on the reef platform surrounding Heron Island Research Station (23.4500°S, 151.9167°E) on the Great Barrier Reef. Sharks were transported in mesh bags to an outdoor 5,000-liter holding tank, where they were maintained at natural winter photoperiod, temperature (range 16.5–20°C), and salinity (35.5–37 ppt), for 37 to 158 h (mean 84 ± 7) prior to experimentation. Dissolved oxygen in the aerated holding tank averaged 6.9 ± 0.1 mg O2/l (parts per million) (>90% air saturation). The sharks were not fed in captivity. Each day 1,000 liters (~20% of the tank volume) of seawater were exchanged in the holding tank with fresh seawater. There were no differences in body mass (356 ± 21 g, P = 0.23) or total length (52.9 ± 1.2 cm, P = 0.10) among the treatments. The animals in the three hypoxic treatments were held in captivity for a shorter period before the experiments (37 ± 2 h) than those in the anoxic treatments (88 ± 9 h). This research was carried out under the auspices of the Great Barrier Reef Marine Parks Authority permits G07/24973.1 and G07/23338.1.

Anoxic and Hypoxic Exposures

Anoxic and hypoxic exposures were conducted in 65-liter glass aquaria with one animal per aquarium. Aquaria were fitted with clear plastic covers to minimize gas diffusion while allowing ambient light to enter, and the sides were wrapped in black plastic to prevent visual disturbance of the sharks. The sharks and treatments were randomly assigned to each of 10 aquaria each day to prevent systematic errors in the experimental design due to tank effects. Experiments were conducted at ambient temperature (16–21°C, mean 18.9°C), which approximated the temperatures observed on the reef platform during the study. Compressed nitrogen gas was bubbled into the chambers via two diffuser stones to create anoxic conditions (~0.1% air saturation, operationally defined as ≤0.03 mg O2/l) or hypoxic conditions (~5% air saturation, operationally defined as 0.39 ± 0.03 mg O2/l). Control normoxic tanks (operationally defined as 6.53 ± 0.10 mg O2/l) were aerated using a single diffuser stone and air pump. A small aquarium pump was used to ensure a homogenous environment within each aquarium. Oxygen concentrations were determined with a TPS WP-82Y dissolved oxygen meter fitted with a YSI 5739 probe, which was calibrated daily in air and zero-oxygen solution (saturated Na2SO3 solution).

The sharks were subjected to one of eight treatments (Fig. 2), in which A1 represents exposure to anoxia until time of loss of the righting reflex (T1RR) (85), A2 represents a second 50 min episode of anoxia after A1 and following 24 h of recovery, H1 and H2 similarly represent one or two 2 h episodes of hypoxia separated by 24 h normoxic recovery, C1 represents 2 h of normoxia (control), and the subsequent times indicate the recovery period in the holding tank before sampling: A1 + 2 h (n = 5), C1 + 2 h (n = 4), A1 + 24 h (n = 5), C1 + 24 h (n = 5), A1 + A2 (n = 5), H1 + 24 h (n = 6), H1 + H2 + 2 h (n = 6), H1 + H2 + 24 h (n = 6). The repeated low oxygen exposures (i.e., A1 and A2, H1 and H2) were spaced 24 h apart to mimic the periodicity of dissolved oxygen fluctuations on the reef platform (Fig. 1). The T1RR was chosen as the endpoint for the A1 anoxia experiments because it may indicate the onset of a deeper phase of metabolic depression (85).

Animals were identified from photographs of the unique spot patterns on the dorsal region. At the beginning of an experiment, the sharks were removed from the holding tank and placed into the.
appropriate anoxic or hypoxic aquarium (i.e., they were subjected to acute low oxygen conditions as in previous studies) (13, 68, 85). Gill ventilation rates (rhythmic gill slit openings in beats per min) were determined every 15 min. In the anoxia treatments, once the ventilation rate had stabilized at a low level (typically 3–10 beats per min after ~100 min), each animal was subsequently tested for reflex responses every 15 min until the T_{1,RR} as in Renshaw et al. (85). The animal was then returned to the holding tank for the +2 h or +24 h recovery period. Most sharks recovered swimming ability within 3–5 min of removal from anoxia, many within seconds. Control and hypoxia treatment animals were handled in a similar fashion. At the appropriate sampling time, two 0.9-ml blood samples were drawn from each animal via the caudal vein into a heparinized syringe. Animals were then euthanized with an overdose of benzocaine (~80–100 mg/l), and sex, weight, and length were determined. The blood was centrifuged at 3,800 rpm for 5 min at 4°C, after which the plasma and red blood cell pellet were separated and frozen in duplicate in liquid nitrogen. The entire rectal gland and one half of the cerebellum were quickly dissected out and snap-frozen in liquid nitrogen; the brain samples were frozen within 4 min. All plasma and tissue samples were shipped on dry ice to the University of California, Davis and then stored at −80°C until analysis.

For comparison with the experimental animals, we also collected field blood samples from epaulette sharks (n = 6; 4 females, 2 males) on the Heron Island reef platform during a hypoxic nocturnal, spring low tide (2–4 AM, Fig. 1).

**Plasma Lactate, Glucose, and Electrolytes**

The concentrations of L-lactate and t-glucose (mmol/l) were determined in triplicate on a YSI 2700 Biochemistry Analyzer. Plasma samples were also analyzed in triplicate for Na⁺ and K⁺ concentrations (mmol/l) by flame photometry (Instrumentation Laboratory 343), Cl⁻ concentration (mmol/l) using a chloride titrator (Labconco Digital Chloridometer), and osmolality (mOsmol/kg) using a vapor pressure osmometer (Wescor Vaporo 5520).

**Enzyme Activity and Glutathione Assays**

Na⁺/K⁺ ATPase activity. The activity of Na⁺/K⁺ ATPase per mg of total protein was determined in triplicate in rectal gland and brain (cerebellum) homogenates following the methods of McCormick (64), as applied to elasmobranchs by Piermarini and Evans (79a).

**Caspase 3/7 activity.** We determined the combined activity of caspases 3 and 7, two proteases that are early effectors of apoptosis, in duplicate using a microplate-based luminescent assay (Caspase Glo 3/7 assay kit, Promega) modified for tissue homogenates (18, 41). Brain and rectal gland tissue samples were homogenized in non denaturating lysis buffer (10 mM Tris-HCl pH 7.5, 100 mM NaCl, 0.1 mM EDTA, 0.2% Triton-X 100). Caspase 3/7 activity was normalized to total protein, as determined by a bicinchoninic acid (BCA) assay (Pierce). Protein concentrations of the reducing agent (≥80–100 mg/l), and sex, weight, and length were determined. The blood was centrifuged at 3,800 rpm for 5 min at 4°C, after which the plasma and red blood cell pellet were separated and frozen in duplicate in liquid nitrogen. The entire rectal gland and one half of the cerebellum were quickly dissected out and snap-frozen in liquid nitrogen; the brain samples were frozen within 4 min. All plasma and tissue samples were shipped on dry ice to the University of California, Davis and then stored at −80°C until analysis.

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**Glutathione.** The level of reduced glutathione (GSH) may decrease during oxidative stress (e.g., during ROS formation following reoxygenation) as GSH donates a reducing equivalent to reactive oxygen species or oxidized proteins and itself is oxidized to GSSG. We measured the level of GSH in brain homogenates (in the same buffer as used for caspase 3/7 activity) in duplicate using a microplate-based luminescent assay (GSH-Glo assay, Promega). We treated two additional wells of each homogenate with 1 μl of the reducing agent Tris(2-carboxyethyl)phosphine (TCEP; 900 μM) to convert all GSSG back to GSH, allowing us to determine the size of the total glutathione pool (GSH + GSSG) as well as the fraction of the glutathione pool in the oxidized and reduced states after each treatment. Luminescence was converted to GSH concentration based on a standard curve created by serial dilution of a 5 mM GSH standard in the homogenization buffer per the manufacturer’s instructions. GSH was normalized to total protein, as determined by a BCA assay (Pierce).

**Statistical Analyses of Physiological and Biochemical Data**

In most cases treatment group means were compared using one-way ANOVA followed by Tukey’s honestly significant difference post hoc test. Lactate and glucose data were first power transformed to satisfy the homogeneity of variances and normality assumptions. Data that could not be transformed to meet the ANOVA assumptions were analyzed with a Kruskal-Wallis (K-W) nonparametric test. Data from reduced GSH levels were arcsine-square-root transformed for proportions. Significance level in all cases was set at α = 0.05. The ventilation rate data were compared among treatments by fitting the data to separate log-linear regression curves for each treatment using Gauss-Newton iteration. Analyses were performed in Statistica 6.1 (StatSoft).

**Proteomics**

**Two-dimensional gel electrophoresis and image analysis.** Proteins regulated by exposure to anoxia or hypoxia in the epaulette shark cerebellum and rectal gland were identified using two-dimensional gel electrophoresis and tandem mass spectrometry following previously established methods (18, 19, 51). We selected the four 24-h recovery time points (Fig. 2) for proteomics analysis. Briefly, proteins were extracted from tissue homogenates in ice-cold RIPA buffer, precipitated in acetone with 10% trichloroacetic acid, loaded (1 mg per sample for cerebellum; 0.75 mg for rectal gland) onto 18-cm immobilized pH 3–10 nonlinear gradient strips (Bio-Rad) by passive overnight rehydration at 4°C, separated in the first dimension by isoelectric focusing, reduced with dithiothreitol and iodoacetamide, and then run on 16-cm second dimension SDS-PAGE gels, all as previously described (18, 19, 107). Two-dimensional gels were stained with Comassie brilliant blue G250, destained in purified water, and then scanned for densitometry (5, 18, 19, 107). The individual gel images (one gel per individual for both cerebellum and rectal gland) were manually aligned, fused into a master gel image for each tissue, and quantified using Delta 2D software (v3.6, Decodon, Greifswald, Germany) as in Ref. 19. Because no samples were pooled, our data approximate the true biological variability.

**Proteomics data analysis.** Protein expression levels on individual two-dimensional gels from control, anoxia, and hypoxia treatments were normalized by dividing each spot’s intensity on a given gel by the total intensity of all outlined spots on that gel. Treatment spot intensities were then compared with control values using a t-test in Delta 2D software (18, 19). Protein spots whose abundance changed significantly (P < 0.05) and with an expression ratio > 2 (increased by >100%) or <0.5 (decreased by >50%) were processed for mass spectrometry identification, where the expression ratio equals the mean spot volume in a given treatment’s gels divided by the mean spot volume in the control gels. These ratios equate to a log_{10} ratio change of ±0.30. Because very few spots met these strict criteria, we expanded the analysis to include spots with P < 0.05 and an expression ratio <0.67 or >1.5 (i.e., log_{10} ratio of ±0.18).

In addition, we performed a partial least squares discriminant analysis (PLS-DA) using SIMCA-P+ software (Umetrics, Umeå, Sweden). This multivariate statistical technique is a regression extension of principal components analysis (PCA) that maximizes the amount of among-groups variation explained by the first components. This technique outperforms PCA, which maximizes the total amount of variation in the dataset explained by the first components regardless of treatment group membership (42, 114). The PLS-DA modeling approach reveals important expression changes in noisy, highly collinear datasets that may be missed by univariate tests such as a t-test (42, 78). Based on the full PLS-DA model with all protein spots included, we performed a variable importance in the projection (VIP) analysis. The VIP is a relative measure of the global importance of a
particular feature (i.e., protein spot); spots with high VIP scores are most relevant to distinguishing among the treatment groups. The top 5% of VIP scores (49 spots for brain; 38 spots for rectal gland) were used to fit reduced PLS-DA models for each tissue, following a feature-selection approach applied to microarray data (78) and recently supported by simulation studies (75). These spots in the top 5% of VIP scores were also picked for mass spectrometry analysis.

Mass spectrometry and bioinformatics database searches. Protein spots of interest were excised in duplicate using a 0.5 mm spot puncher and digested overnight at 37°C with mass spectrometry-grade trypsin (Promega). The resulting peptide fragments were eluted from gel pieces by washing twice in 60% acetonitrile:1% trifluoroacetic acid (TFA). The peptide solution was then dried in a SpeedVac and resuspended in 1% TFA. The peptides were desalted and concentrated using μC18 ZipTips (Millipore). The final eluted peptides were spotted in 0.5 μl increments on a stainless steel MALDI target, and overlaid with a-cyano-4-hydroxycinnamic acid matrix (18, 19). Peptides were analyzed on an Applied Biosystems 4700 Proteomics Analyzer MALDI-TOF-TOF tandem mass spectrometer (Foster City, CA) as previously described (18, 19, 51). Peak lists from the resulting MS and MS/MS spectra were searched jointly against the NCBI non-redundant database using the Mascot search algorithm (http://www.matrixscience.com, 79) with mass error tolerances of 50 and 100 ppm for MS, 0.3 and 0.5 Da for MS/MS, and fixed modifications of cysteine residues (carbamidomethylation) and variable oxidation of methionine residues. The MS/MS spectra were also submitted for automated de novo peptide sequencing analysis to PEAKS Studio v4.5 software (Bioinformatics Solutions, Toronto, Canada) (60), with the same criteria as for Mascot. The resulting de novo amino acid sequences for the tryptic peptides were then searched against the PEAKS proprietary search engine. Consensus protein identifications were determined (in duplicate) based on both Mascot and PEAKS scores and from both postsource decay (PSD) and collision-induced dissociation (CID) MS/MS spectra. Matches solely from de novo sequencing scores were only accepted when >1 peptide matched to the protein. When possible, matches were confirmed by general agreement of molecular mass (M), and isoelectric point (pI) between the database match and the protein spot of interest on 2D gels. However, because many proteins have not been sequenced in elasmobranchs, this is not always possible.

Pathway and molecular network analyses. To aid in the development of functional hypotheses consistent with the proteomics data, the lists of epaulette shark proteins were analyzed separately for cerebellum and rectal gland using Ingenuity Pathways Analysis (IPA) software (Ingenuity Systems, http://www.ingenuity.com). Identifiers for the mouse or human homologs of the identified epaulette shark proteins (from both the t-test and VIP analyses) and the corresponding expression ratios and P values for each treatment were uploaded into IPA. Lists of significantly represented (based on right-tailed Fisher’s exact test) molecular functions (Gene Ontology (GO; http://www.geneontology.org) biological process annotations plus manually added annotations) and canonical cellular pathways were generated for each treatment. The lists of protein identifiers were then overlaid onto the IPA knowledge base global molecular network. Molecular interaction networks containing the anoxia or hypoxia-regulated proteins were algorithmically generated based on their specific connectivity, with each network assigned a statistical likelihood score (10). We first created a master network for each tissue containing all of the proteins of interest (i.e., those that met the t-test or PLS-DA VIP criteria in any treatment). The expression ratios for each treatment were overlaid onto this master network to enable direct, visual comparison of the expression changes across treatments.

RESULTS

Time to Loss of Righting Reflex

The time to loss of righting reflex varied considerably (69–258 min) among the experimental animals in anoxia, but there were no systematic differences among the three A1 treatments (A1+2 h, A1+24 h, A1+A2) in TLR (P = 0.27, grand mean 142 ± 16 min). The variation in TLR was not correlated with body mass (P = 0.80), time in captivity (i.e., by potential loss of energy stores) (P = 0.25), or the time of day that the experiment started (P = 0.51); there was also no difference between the sexes (P = 0.39). Thus, these data can best be explained as individual differences. Compared with an earlier study (85), the TLR in anoxia was significantly longer in our A1 treatments (142 min at 18.9°C vs. 46 min at 28°C).

Ventilation Rate Decreases Rapidly in Anoxia but not Hypoxia

Ventilation rate was elevated in all treatments initially, most likely reflecting handling stress incurred while placing the animals in the chambers. Ventilation frequency in the control animals after ~60 min was similar to that observed in the holding tank and in the wild (~16–19 beats/min). Ventilation rate decreased faster and to lower levels in the three A1 anoxia treatments than in the controls (Fig. 3). The 95% confidence intervals for the slope of the log-linear Gauss-Newton fit overlapped among the three A1 treatments (A1+2: -22.2 to
Plasma lactate concentrations increased significantly after low-oxygen exposure (P < 0.001) (Fig. 4). The highest measured plasma lactate concentrations occurred 2 h after return from anoxia to normoxic conditions (A1+2 h), but we do not know how lactate varied during the episodes of oxygen limitation. In all anoxic and hypoxic treatments plasma lactate returned to baseline values by 24 h recovery. There was a significant, but small, effect of treatment on plasma glucose concentrations (P = 0.013) (Fig. 4), with the H1+24, H1+H2+24 and Field treatments tending to have less circulating glucose than other treatments. However, the Tukey’s HSD post hoc test failed to detect any pair-wise treatment differences. The less conservative Fisher Least Significant Difference post hoc test revealed three statistically homogeneous groups. There was no significant effect of oxygen limitation on plasma osmolality (K-W; P = 0.410), sodium concentration (K-W; P = 0.098), or chloride concentration (K-W; P = 0.219) (Fig. 5, A and B). Plasma potassium concentration varied among treatments (P < 0.001) (Fig. 5C), but the changes were minor and not systematically associated with specific low-oxygen conditions.

Ion-Pumping and Apoptotic Effector Enzyme Activities Do Not Respond to Oxygen Limitation and Recovery

There were no differences in cerebellum or rectal gland Na+/K+ ATPase activities 24 h after anoxic or hypoxic exposure (Table 1). Rectal gland Na+/K+ ATPase activity was significantly greater than that in brain.

There was a significant effect of treatment on caspase 3/7 in brain tissue (P = 0.032), but the only pair-wise significant difference detected with the post hoc test was between A1+2 h and H1+24 (Fig. 6A). Notably, there were no increases in caspase 3/7 activity in brain following any of the hypoxic or anoxic treatments relative to controls, suggesting no early (2 h) or delayed (24 h) caspase-dependent apoptosis. Similarly, the rectal gland exhibited no increases in caspase 3/7 activity after oxygen limitation (Fig. 6B).

No Long-term Effect of Episodic Oxygen Limitation on the Total Glutathione Pool in Cerebellum

There was a significant effect of treatment on both the reduced (P < 0.001) and total (P < 0.001) glutathione pools in epaulette shark brain (Fig. 7B). Specifically, both the control (C1) and anoxia (A1) treatments +2 h exhibited lower levels of both reduced and total glutathione than the corresponding +24 h treatments. A similar, though statistically insignificant, trend was observed for the repeated hypoxia treatments (H1+H2+2 h vs. H1+H2+24 h). The fraction of the total glutathione pool in the reduced state (% GSH) similarly varied among treat-
ments (P = 0.003); more glutathione was oxidized in +2 h brains than in the respective +24 h group for the control (C1), anoxia (A1), and repeated hypoxia (H1+H2) treatments (Fig. 7A). These data suggest a common oxidative stressor among the control and experimental treatments, most likely as a result of handling stress when placing the animals in the experimental chambers and removing them after treatment. However, almost all glutathione was in the reduced state (GSH) for the four +24 h treatments (Fig. 7A), implying that the GSH system recovered by 24 h regardless of the treatment. Furthermore, there were no significant differences among the four +24 h treatments in the reduced or total glutathione pools.

**Small Magnitude, Treatment- and Tissue-Specific Proteome Responses to Episodic Oxygen Limitation and Reoxygenation**

**Cerebellum.** The master gel image for cerebellum contained 986 protein spots that were quantified using Delta 2D software (Supplemental Fig. S1A). Of these, 14 unique spots met the 986 protein spots that were quantified using Delta 2D software treatments (Fig. 7A). These data suggest common oxidative stressors among the control and experimental treatments, most likely as a result of handling stress when placing the animals in the experimental chambers and removing them after treatment. However, almost all glutathione was in the reduced state (GSH) for the four +24 h treatments (Fig. 7A), implying that the GSH system recovered by 24 h regardless of the treatment. Furthermore, there were no significant differences among the four +24 h treatments in the reduced or total glutathione pools.

**Table 1. Activities of Na\(^{+}/K^{+}\) ATPase (μmol ADP·h\(^{-1}\)/mg protein\(^{-1}\)) in epaulette shark rectal gland and brain (cerebellum)**

<table>
<thead>
<tr>
<th>Tissue</th>
<th>C1 + 24</th>
<th>n</th>
<th>A1 + 24</th>
<th>n</th>
<th>H1 + 24</th>
<th>n</th>
<th>H1+H2+24</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rectal gland</td>
<td>12.6 ± 2.5</td>
<td>4</td>
<td>12.8 ± 2.2</td>
<td>5</td>
<td>17.8 ± 3.9</td>
<td>6</td>
<td>13.4 ± 2.5</td>
<td>5</td>
</tr>
<tr>
<td>Brain</td>
<td>3.9 ± 0.9</td>
<td>4</td>
<td>4.0 ± 0.9</td>
<td>5</td>
<td>3.2 ± 0.1</td>
<td>5</td>
<td>3.6 ± 0.4</td>
<td>6</td>
</tr>
</tbody>
</table>

Values are means ± SE. There were no significant differences among the treatments in either tissue.

\[1\] The online version of this article contains supplemental material.

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Fig. 6. Combined activity of caspases 3 and 7, two closely related effectors of apoptosis, in epaulette shark brain (cerebellum) (A) and rectal gland (B) following control (normoxic), anoxic, and hypoxic exposure plus recovery (treatment codes as in text). Activity was determined using a luminescent assay (RLU, relative light units). Different lowercase letters indicate statistically distinguishable groups for brain as determined by Tukey’s HSD post hoc test. Values for cerebellum are means ± SE from 4–6 individuals per treatment. Individual data points are presented for rectal gland due to small samples sizes in some treatments.

Fig. 7. A: percentage of the epaulette shark brain (cerebellum) total glutathione pool in the oxidized form (GSH) following control (normoxic), anoxic, or hypoxic exposure plus recovery (treatment codes as in text). B: reduced glutathione (GSH) and total glutathione (GSH+GSSG) pools in epaulette shark cerebellum in each treatment. GSH is oxidized to GSSG during oxidative stress. Different lowercase letters or numbers represent statistically different groups for GSH and total glutathione, respectively. Values are means ± SE from 4–6 individuals per treatment.
Table 2. Number of significant changes ($P < 0.05$) in relative protein abundance in brain (cerebellum) and rectal gland of epaulette sharks 24 h after exposure to anoxia (A1 + 24) and one (H1 + 24) or two episodes (H1 + H2 + 24) of hypoxia.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Brain</th>
<th>Rectal Gland</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$r &lt; 0.5$ or $r &gt; 2$</td>
<td>$r &lt; 0.67$ or $r &gt; 1.5$</td>
</tr>
<tr>
<td>A1 + 24</td>
<td>3 (−), 0 (+)</td>
<td>10 (−), 3 (+)</td>
</tr>
<tr>
<td>H1 + 24</td>
<td>6 (−), 1 (+)</td>
<td>25 (−), 14 (+)</td>
</tr>
<tr>
<td>H1 + H2 + 24</td>
<td>4 (−), 2 (+)</td>
<td>20 (−), 15 (+)</td>
</tr>
<tr>
<td>A1 + 24 &amp; H1 + H2 + 24</td>
<td>1 (−), 0 (+)</td>
<td>10 (−), 3 (+)</td>
</tr>
<tr>
<td>A1 + 24 &amp; H1 + H2 + 24</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>H1 + 24 &amp; H1 + H2 + 24</td>
<td>1: novel protein</td>
<td>8: novel protein</td>
</tr>
<tr>
<td></td>
<td>(zgc:92871)</td>
<td>(Danio rerio) (+/+);</td>
</tr>
<tr>
<td></td>
<td>(+/+)</td>
<td>splicing factor, RS rich 5 (-/-);</td>
</tr>
<tr>
<td></td>
<td></td>
<td>calmodulin (+/+/-);</td>
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<tr>
<td></td>
<td></td>
<td>heat shock 70 kDa protein 4-like</td>
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<tr>
<td></td>
<td></td>
<td>(-/-); ubiquitin-conjugating</td>
</tr>
<tr>
<td></td>
<td></td>
<td>enzyme E2D 2 isoform 1 (+/+/-);</td>
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<tr>
<td></td>
<td></td>
<td>methylcrotonoylcoenzyme A</td>
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<td></td>
<td></td>
<td>carboxylase 2 (beta) (-/-);</td>
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</tr>
<tr>
<td></td>
<td>3: EF hand calcium binding</td>
<td>1: alpha tubulin (-/-)</td>
</tr>
<tr>
<td></td>
<td>protein 2 (-/-)</td>
<td></td>
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</tbody>
</table>

Values are given for both the strict and lenient expression ratio criteria [$r$; relative to controls (C1 + 24)]. (+) increased; (−) decreased. Individual protein spots were counted in multiple categories where appropriate.

Rectal gland. The master gel image for rectal gland contained 770 protein spots that were quantified using Delta 2D software (Supplemental Fig. S1B). Of these, 29 unique spots met the twofold change and 95% confidence criteria in one or more low oxygen treatments, and the abundances of 101 unique protein spots changed in response to episodic anoxia and/or hypoxia using the less stringent criteria (Table 2). In contrast to cerebellum, more rectal gland spots were upregulated relative to controls than downregulated. As in cerebellum, the highest number of significant changes in protein abundance occurred in H1 + 24 h, followed by H1 + H2 + 24 h and then A1 + 24 h (Table 2). A greater number of consistent changes in individual protein expression occurred across treatments in rectal gland relative to cerebellum (Table 3). The reduced rectal gland PLS-DA model based on the top 38 (5%) VIP scores performed better than expected by chance (Supplemental Fig. S2B) and clearly discriminated among the four treatment groups (Fig. 8B). Similar to cerebellum, 14 of the top 38 VIP scores (37%) were also selected by the $r$-test/ratio criteria (Table 5).

Of the 101 proteins of interest in rectal gland, 53 (52%) were identified using tandem mass spectrometry combined with bioinformatics searches (Supplemental Table S3). The significant IPA canonical pathways are presented in Supplemental Table S2. The master molecular interaction network for rectal gland (Supplemental Fig. S4) was created by merging four significantly matching networks in IPA: 1) energy production, nucleic acid metabolism, small molecule biochemistry ($P = 10^{-30}$); 2) drug metabolism, lipid metabolism, small molecule biochemistry ($P = 10^{-30}$); 3) cancer, amino acid metabolism ($P = 10^{-30}$); 4) unnamed ($P = 0.01$).

The rectal gland and cerebellum interaction networks shared four highly connected protein nodes that may warrant further examination in the context of the molecular response to episodic hypoxia or anoxia in this species (Supplemental Figs. S3 and S4): huntingtin; solute carrier family 2 (facilitated glucose transporter), member 4; hepatocyte nuclear factor 4, alpha; and peroxisome proliferator-activated receptor gamma, coactivator 1 alpha. However, it should be noted that these molecular interaction networks were constructed using data from all three treatment groups (A1+24, H1+24, H1+H2+24), but the response to oxygen limitation varied considerably among treatments.

**DISCUSSION**

Epaulette Shark as Comparative Model of Hypoxia and Anoxia Tolerance

The epaulette shark offers an appealing comparative system for studying evolutionary and ecological processes leading to “constitutively preconditioned” (66) models of hypoxia and anoxia tolerance as well as hypoxic preconditioning in vertebrates.
The duration (2 h) and severity (5% air saturation or complete anoxia) of the experimental regimen used here exceed those that would be tolerable by most naïve mammalian systems. Much previous work on tolerance of oxygen limitation in fishes has involved species from the Teleostei (bony fishes), but few studies have addressed mechanisms of hypoxia or anoxia tolerance in the Chondrichthyes (sharks, skates, rays, and chimaeras). Recent works suggest that a few sharks other than the epaulette periodically enter (39) and/or tolerate (13) severely hypoxic or anoxic waters. Unearthing the mechanisms of hypoxia and anoxia tolerance and preconditioning in this ancestral vertebrate lineage may provide important insights into the evolution of these traits and the relative importance of conserved (e.g., reduced glutamate signaling, 104) vs. unique mechanisms.

Our results indicate a suite of small-magnitude changes for proteins involved in several processes relevant to the survival of oxygen limitation and reoxygenation (31). Notably, the relative importance of these responses in the epaulette shark appears to vary depending on the tissue and treatment examined. While proteome responses to H1 and H2 were more similar to each other than to A1 (Table 3), responses to the four treatments were clearly distinguished in the PLS-DA analysis for each tissue (Fig. 8). Surprisingly, one episode of anoxia elicited fewer changes in protein expression than either one or two episodes of hypoxia. Several of the identified proteins and processes in both cerebellum and rectal gland have highly conserved functions among hypoxia or anoxia-tolerant vertebrates, although these mechanisms remain to be further validated in this species. We discuss several of the more prominent patterns in these two tissues and propose novel hypotheses for both enhanced hypoxia and anoxia tolerance and preconditioning in the epaulette shark.

Prevention of Excitotoxicity in Cerebellum: Decreased Excitatory Neurotransmission, Modified Vesicle Dynamics, and Increased Calcium Binding

A number of protein expression changes in the cerebellum were consistent with the prevention of excitotoxicity via their roles in neurotransmitter dynamics. One of the major causes of neuronal cell death in cerebral ischemia/reperfusion, excitotoxicity involves a cascade of events including elevation of cytosolic calcium concentration ([Ca^{2+}]), resulting in release of the excitatory neurotransmitter glutamate and initiation of a lethal cycle of membrane depolarization. The classical adaptive decrease in action potential firing, via decreased glutamate signaling, during oxygen limitation in tolerant animals is known as "spike arrest" (97). Several proteins potentially related to glutamate production and release were downregulated in epaulette shark cerebellum following two episodes of hypoxia and, to a lesser extent, anoxia. Foremost among these, the decrease in GLS (all abbreviations as in Tables 4 and 5) after H2 is consistent with reduced glutamate production (45) after multiple hypoxic episodes.

Several of the identified proteins play important roles in vesicle dynamics, implying adjustments in cycling of neurotransmitters and/or receptors in epaulette shark neurons after episodic oxygen limitation. Again, these changes were most pronounced following two hypoxic episodes. The SNAP protein NAPB (\(\downarrow A1\) and \(\downarrow H2\)) recruits NSF to membranes for vesicle fusion and exocytosis (i.e., for neurotransmitter release) (100). NSF itself also decreased here after H2, which could have antiexcitotoxic pre- and postsynaptic consequences via decreased glutamate release (77) and decreased stabilization of AMPA glutamate receptors in the postsynaptic membrane (6), respectively. SH3GL1 decreased after A1; this protein may be needed for vesicle fusion in the presynaptic terminal (86). Two additional proteins that may participate in postsynaptic, receptor-mediated endocytosis (CLTA \(\downarrow H1\)) and trafficking of...
Table 4. Expression ratios (relative to control CI × 24) for epaulette shark cerebellum proteins regulated by episodic anoxia (AI × 24) or hypoxia (HI × 24; H1 × H2 + 24)

<table>
<thead>
<tr>
<th>Spot</th>
<th>VIP</th>
<th>VIP Score</th>
<th>Protein Identification</th>
<th>Abbrev.</th>
<th>Expression Ratio</th>
</tr>
</thead>
<tbody>
<tr>
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<td></td>
<td></td>
<td>AI × 24</td>
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<td></td>
<td></td>
<td>P</td>
</tr>
</tbody>
</table>

**Energy-producing processes**

Glycolysis
1.5–66
glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (Paralichthys olivaceus)
GAPDH 0.82 0.242 0.65 0.005 0.69 0.002

Oxidative phosphorylation/electron transport chain
1.5–33
VIP06 1.69 mitochondrial ATP synthase alpha-subunit (Cyprinus carpio)
TP5A1 0.67 0.002 0.89 0.453 1.24 0.091
1.5–36
VIP08 1.68 ATP synthase subunit alpha (Larus ridibundus)
TP5A1 0.59 0.016 0.62 0.104 1.16 0.407
1.5–38
VIP16 1.59 NADH dehydrogenase flavoprotein 2, 24 kDa subunit (Mus musculus)
NDUFV2 0.89 0.124 0.96 0.572 0.79 0.025

Oxidative substrate switching
1.5–64
3-oxoacid CoA transferase 1 precursor (Homo sapiens)
OXCT1 1.25 0.349 1.65 0.012 1.96 0.061
1.5–26
VIP08 1.68 methylcrotonoyl-Coenzyme A carboxylase 2 (beta) (Homo sapiens)
MCCC2 1.04 0.772 0.66 0.015 0.63 0.016
1.5–31
VIP16 1.59 propionyl-CoA carboxylase (Xenopus laevis)
PCCA 1.27 0.346 1.76 0.089 1.60 0.007

Intracellular movement of energetic equivalents
VIP04 1.73 voltage-dependent anion-selective channel protein 2 (Xenopus laevis)
VDAC2 1.06 0.672 1.24 0.085 0.76 0.046
VIP23 1.56 voltage-dependent anion-selective channel protein 2 (Squalus acanthias)
VDAC2 0.76 0.021 0.71 0.013 0.86 0.202

**Energy-consuming processes**

Proteolysis in proteasome
1.5–39
proteasome 26S ATPase subunit 2 (Homo sapiens)
PSMC2 0.91 0.367 0.62 0.030 0.84 0.027
1.5–40
VIP19 1.56 proteasome 26S non-ATPase subunit 11
PSMD11 0.86 0.113 0.83 0.213 0.55 0.002

Lysosome
1.5–29
H+-ATPase beta 1 subunit, lysosomal
ATP6V1B1 1.03 0.801 1.46 0.013 1.90 0.015

Transcription and mRNA processing
2–7
splicing factor, arginine/serine-rich 5a (Danio rerio)
SFRS5 0.82 0.366 0.45 0.001 0.63 0.008
1.5–62
RNA binding motif protein 8A (Danio rerio)
RBMA8 0.78 0.264 0.76 0.289 0.60 0.047

Translation
1.5–30
Seryl tRNA synthetase (Danio rerio)
SARS 0.73 0.156 0.70 0.085 0.65 0.040
1.5–48
acidic ribosomal phosphoprotein P0 (Bos taurus)
RPLP0 0.95 0.713 0.63 0.038 1.03 0.887

VIP43 1.48 elongation factor 1-delta (Tetraodon nigroviridis)
EEF1D 0.71 0.210 0.95 0.811 0.55 0.080

**Apoptosis and cell cycle**

1.5–14
far upstream element-binding protein (Homo sapiens)
FUBP1 1.26 0.367 1.48 0.195 1.95 0.024
1.5–23
VIP28 1.53 ubiquitin-conjugating enzyme E2D 2 isoform 1 (Homo sapiens)
UBE2D2 0.64 0.075 1.59 0.027 1.55 0.031

1.5–45
protein phosphatase 1 alpha (Danio rerio)
PPP1CA 1.10 0.649 0.64 0.017 0.66 0.060
1.5–58
Ras-related nuclear protein (Ictalurus punctatus)
RAN 1.14 0.355 1.57 0.041 1.37 0.091

VIP26 1.54 high mobility group box 2 (Mus musculus)
HMGB2 0.58 0.071 0.76 0.271 0.69 0.184
<table>
<thead>
<tr>
<th>Spot</th>
<th>VIP</th>
<th>VIP Score</th>
<th>Protein Identification</th>
<th>Abbrev.</th>
<th>Expression Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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<td></td>
<td>A1 + 24</td>
</tr>
<tr>
<td>Glutamate synthesis/degradation</td>
<td></td>
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<tr>
<td>1.5–27</td>
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<td>glutaminase (Rattus norvegicus)</td>
<td>GLS</td>
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<td>1.5–15</td>
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<td>N-ethylmaleimide sensitive fusion protein (Cricetulus longicaudatus)</td>
<td>NSF</td>
<td>0.80</td>
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<td>1.5–44</td>
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<td>SH3 domain-containing Grb-2-like protein (Xenopus tropicalis)</td>
<td>SH3GL1</td>
<td>0.60</td>
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<tr>
<td>1.5–51</td>
<td></td>
<td></td>
<td>clathrin light-chain B isoform 1 (Monodelphis domestica)</td>
<td>CLTA</td>
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<td>1.5–52</td>
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<td></td>
<td>dihydropyrimidinase-like 2 (Danio rerio)/collapsin response mediator protein-1A</td>
<td>TPYSL2</td>
<td>1.11</td>
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<td>Neurotransmitters</td>
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<td>1.5–18</td>
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<td>1.52</td>
<td>calmodulin (Clemmys japonica)</td>
<td>CALM1</td>
<td>0.80</td>
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<td>Molecular chaperones</td>
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<td>heat shock 70 kDa protein 4-like (unnamed protein product, Tetraodon nigroviridis)</td>
<td>HSPA4L</td>
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<td>chain A, dimerization and U-box domains of zebrafish c-terminal of HSP70 interacting protein prohibitin (Equus caballus)</td>
<td>PHB</td>
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<td>Cytoskeleton</td>
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<td>2–4</td>
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<td>1.47</td>
<td>alpha tubulin (Gillichthys mirabilis)</td>
<td>TUBA1A</td>
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<tr>
<td>2–5</td>
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<td>alpha tubulin (Gillichthys mirabilis)</td>
<td>TUBA1A</td>
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<td>1.54</td>
<td>alpha tubulin subunit (Oncorhynchus nerka)</td>
<td>TUBA1A</td>
<td>1.11</td>
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<tr>
<td>1.5–28</td>
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<td>alpha tubulin</td>
<td>TUBA1A</td>
<td>1.51</td>
</tr>
<tr>
<td>1.5–32</td>
<td></td>
<td></td>
<td>alpha tubulin</td>
<td>TUBA1A</td>
<td>1.13</td>
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<tr>
<td>1.5–34</td>
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<td>alpha tubulin</td>
<td>TUBA1A</td>
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<td>tubulin alpha</td>
<td>TUBA1A</td>
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<td>alpha 4 tubulin</td>
<td>TUBA4A</td>
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<td>2–9</td>
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<td>tubulin folding cofactor B (cytoskeleton-associated protein 1) (LOC735205, Xenopus laevis)</td>
<td>TBCB</td>
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<td>Actin cytoskeleton and dendrite morphology</td>
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<td>1.5–42</td>
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<td>beta actin</td>
<td>ACTB</td>
<td>1.03</td>
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<td>1.5–11</td>
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<td>mark (myristoylated alanine-rich C-kinase substrate) calmodulin binding domain protein</td>
<td>MARCKS</td>
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<td>1.5–10</td>
<td></td>
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<td>dynamin 3</td>
<td>DNM3</td>
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<td>1.5–41</td>
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<td></td>
<td>glial fibrillary acidic protein (Mus musculus)</td>
<td>GFAP</td>
<td>1.06</td>
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</tbody>
</table>

Continued
ubiquinated endosome-contained proteins such as receptors (TOLLIP → H1) decreased significantly after only one episode of hypoxia, but the significance of these expression changes is unknown. Meanwhile, the lysosomal \( \text{H}^+/-\text{ATPase} \) \( \text{ATP6V1B1} \) increased after both H1 and H2, possibly indicating activation of lysosomal degradation pathways following hypoxia.

Two calcium-binding proteins were also regulated by hypoxic or anoxic exposure in brain. Expression of the EF-hand domain-containing protein NECAB2 (\( \rightarrow \) A1 and H1) is inversely correlated with cell surface expression of adenosine A2A receptors (11). Our result implies increased availability of A2A receptors following anoxia or hypoxia, consistent with the role for adenosine in the epaulette shark’s response to oxygen limitation (71, 85). Binding of adenosine to the A2A receptor in mammalian models increases \([\text{Ca}^{2+}]_i\); accordingly, we observed that CALM increased after both H1 and H2. This suggests enhanced \( \text{Ca}^{2+} \) binding capacity to prevent excitotoxicity and/or elevated calmodulin-dependent signaling, the latter of which plays a known role in the cellular response to hypoxia (95). Overall, our results imply that modulation of the controls of \([\text{Ca}^{2+}]_i\) and excitotoxicity plays an important role in the epaulette shark cerebellum’s response to episodic oxygen limitation.

### Cytoskeletal and Membrane Rearrangements in Response to Hypoxia

Another prominent result from the cerebellum was the proteomic evidence for cytoskeletal and membrane reorganization following oxygen limitation, particularly after two episodes of hypoxia (H2). Nine protein spots identified as alpha tubulins exhibited significant expression changes in cerebellum following hypoxia (4 after H1, 3 after H2), and the tubulin folding cofactor TBCB decreased after H1, suggesting remodeling of microtubule networks. We also observed changes in the abundance of an actin cytoskeleton component (ACTB → H2), anchoring proteins that link the actin cytoskeleton to the cell membrane.

**Table 4.—Continued**

<table>
<thead>
<tr>
<th>Spot</th>
<th>VIP</th>
<th>VIP Score</th>
<th>Protein Identification</th>
<th>Abbrev.</th>
<th>A1 + 24</th>
<th>P</th>
<th>H1 + 24</th>
<th>P</th>
<th>H2 + 24</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cytoskeleton anchoring</td>
<td>1.5–25</td>
<td>VIP35</td>
<td>1.51</td>
<td>ezrin (( \text{Homo sapiens} ))</td>
<td>EZR</td>
<td>0.91</td>
<td>0.603</td>
<td>1.56</td>
<td>0.028</td>
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<tr>
<td>1.5–54</td>
<td></td>
<td></td>
<td>nebulla (( \text{Homo sapiens} ))</td>
<td>NEBL</td>
<td>1.04</td>
<td>0.779</td>
<td>0.77</td>
<td>0.232</td>
<td>0.63</td>
<td>0.033</td>
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<td>1.5–43</td>
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<td></td>
<td>ribosomal protein SA (laminin-binding protein) (( \text{Homo sapiens} ))</td>
<td>RPSA</td>
<td>0.99</td>
<td>0.959</td>
<td>0.51</td>
<td>0.021</td>
<td>0.69</td>
<td>0.086</td>
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<td>cAMP-dependent protein kinase (PKA)</td>
<td>2–6</td>
<td></td>
<td>cAMP-dependent protein kinase type II-beta regulatory subunit isoform 1 (( \text{Canis familiaris} ))</td>
<td>PRKAR2B</td>
<td>1.10</td>
<td>0.774</td>
<td>0.42</td>
<td>0.047</td>
<td>0.95</td>
<td>0.907</td>
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<td>1.5–37</td>
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<td></td>
<td>cAMP-dependent protein kinase type I-alpha regulatory subunit (( \text{Mus musculus} ))</td>
<td>PRKAR1A</td>
<td>0.92</td>
<td>0.428</td>
<td>1.56</td>
<td>0.151</td>
<td>1.68</td>
<td>0.007</td>
</tr>
<tr>
<td>14-3-3 mediated phosphorylproteins adapter signaling</td>
<td>1.5–55</td>
<td>VIP15</td>
<td>1.60</td>
<td>14-3-3 theta (( \text{Danio rerio} ))</td>
<td>YWHAQ</td>
<td>0.62</td>
<td>0.097</td>
<td>0.54</td>
<td>0.049</td>
<td>0.82</td>
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<td>1.5–46</td>
<td>VIP21</td>
<td>1.56</td>
<td>3-monoxygenase/tryptophan 5-monoxygenase activation protein, gamma polypeptide (14-3-3 gamma) (( \text{Danio rerio} ))</td>
<td>YWHAG</td>
<td>0.89</td>
<td>0.033</td>
<td>1.02</td>
<td>0.696</td>
<td>0.94</td>
<td>0.273</td>
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<tr>
<td></td>
<td>VIP46</td>
<td>1.47</td>
<td>14-3-3 beta (( \text{Tetraodon nigroviridis} ))</td>
<td>YWHAB</td>
<td>0.75</td>
<td>0.030</td>
<td>0.77</td>
<td>0.068</td>
<td>0.83</td>
<td>0.136</td>
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<tr>
<td>G protein-coupled receptors</td>
<td>1.5–47</td>
<td></td>
<td>guanine nucleotide binding protein Ga(0), alpha subunit (( \text{Salmo salar} ))</td>
<td>GNAO1</td>
<td>0.91</td>
<td>0.709</td>
<td>0.59</td>
<td>0.007</td>
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<td>0.416</td>
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<tr>
<td>1.5–35</td>
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<td></td>
<td>aldehyde dehydrogenase 7 family, member A1 (( \text{Monodelphis domestica} ))</td>
<td>ALDH7A1</td>
<td>0.82</td>
<td>0.252</td>
<td>0.82</td>
<td>0.399</td>
<td>0.63</td>
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<tr>
<td>Unannotated</td>
<td>2–2</td>
<td>VIP29</td>
<td>1.53</td>
<td>novel protein (zgc:92871) (( \text{Danio rerio} ))</td>
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<td>2.93</td>
<td>0.005</td>
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Proteins were grouped by putative function based on Gene Ontology annotations and manual review of the literature. Values in boldface met the statistical (\( P < 0.05 \)) and expression ratio (\( >1.5 \)) criteria for mass spectrometry analysis in that treatment. Variable importance in the projection ranks (VIP) and scores (VIP score) based on a partial least squares discriminant analysis are also presented. Expression data for unidentified protein spots can be found in Supplemental Table S4.
Table 5. Expression ratios (relative to control C1 + 24) for epaulette shark rectal gland proteins regulated by episodic anoxia (A1 + 24) or hypoxia (H1 + 24; H1 + H2 + 24)

<table>
<thead>
<tr>
<th>Spot</th>
<th>VIP</th>
<th>VIP Score</th>
<th>Protein Identification</th>
<th>Abbrev.</th>
<th>Energy-producing processes</th>
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<td></td>
<td>A1 + 24</td>
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<td>ALDOB</td>
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<td>TP1</td>
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<td>1.5–47</td>
<td>1.56</td>
<td>glyceraldehyde-3-phosphate dehydrogenase (Paralichthys olivaceus)</td>
<td>GAPDH</td>
<td>1.72</td>
<td>0.024</td>
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<td>1.5–60</td>
<td>1.50</td>
<td>phosphoglycerate mutase 1 (Gallus gallus)</td>
<td>PGAM1</td>
<td>0.62</td>
<td>0.014</td>
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<td>2–12</td>
<td>1.54</td>
<td>pyruvate dehydrogenase (lipoamide) beta (Danio rerio)</td>
<td>PDHB</td>
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<td>1.5–48</td>
<td>1.51</td>
<td>pyruvate dehydrogenase (lipoamide) beta (Danio rerio)</td>
<td>PDHB</td>
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<td>0.031</td>
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<td>dihydrolipoamide branched chain transacylase E2 (Taeniopygia guttata)</td>
<td>DBT</td>
<td>1.34</td>
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<td>isocitrate dehydrogenase 2 (NADP+), mitochondrial</td>
<td>IDH2</td>
<td>0.80</td>
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<td>2–14</td>
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<td>ATP synthase subunit alpha, mitochondrial precursor (Salmo salar)</td>
<td>ATP5A1</td>
<td>0.64</td>
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<td></td>
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<td>mitochondrial ATP synthase alpha subunit (Scyliorhinus canicula)</td>
<td>ATP5A1</td>
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<td>0.024</td>
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<td></td>
<td>1.44</td>
<td>ATP synthase, H+ transporting, mitochondrial F1 complex, alpha subunit (Danio rerio)</td>
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<td>NADH dehydrogenase (ubiquinone) Fe-S protein 1 (Danio rerio)</td>
<td>NDUS1</td>
<td>0.88</td>
<td>0.478</td>
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<td>NADH dehydrogenase (ubiquinone) Fe-S protein 1, 75 kDa (NADH-coenzyme Q reductase) (Gallus gallus)</td>
<td>NDUS1</td>
<td>0.94</td>
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<td>NADH dehydrogenase (ubiquinone) Fe-S protein 1, 75 kDa (NADH-coenzyme Q reductase) (Gallus gallus)</td>
<td>NDUS1</td>
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<td>1.5–54</td>
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<td>similar to NADH dehydrogenase (ubiquinone) Fe-S protein 3, 30 kDa (NADH-coenzyme Q reductase)</td>
<td>NDUS3</td>
<td>0.61</td>
<td>0.024</td>
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<td>1.5–55</td>
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<td>succinate dehydrogenase complex, subunit B, iron sulfur (Ip) (Danio rerio)</td>
<td>SDHB</td>
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<td>mitochondrial aspartate aminotransferase</td>
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<td>voltage-dependent anion channel 2 (Siphus acanthias)</td>
<td>VDAC2</td>
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<td>creatine kinase (Scyliorhinus canicula)</td>
<td>CKB</td>
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<td>0.496</td>
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<td>creatine kinase (Scyliorhinus canicula)</td>
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Continued
### Table 5.—Continued

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<th>Abbrev.</th>
<th>Expression Ratio</th>
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<tr>
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<td>A1 + 24</td>
</tr>
<tr>
<td>1.5–24</td>
<td>VIP38</td>
<td>1.39</td>
<td>3-oxoacid CoA transferase (oxct1) (Xenopus tropicalis)</td>
<td>OXCT1</td>
<td>0.98</td>
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<td>2–11</td>
<td>VIP37</td>
<td>1.40</td>
<td>3-hydroxyacyl-CoA dehydrogenase</td>
<td>HADH</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>acyl-CoA dehydrogenase (EC 1.3.99.3) precursor, short-chain-specific</td>
<td>ACADS</td>
<td>0.90</td>
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</tbody>
</table>

#### Oxidative substrate switching

- 3-oxoacid CoA transferase
- 3-hydroxyacyl-CoA dehydrogenase

#### Proteolysis

- Ubiquitin carboxyl-terminal esterase L3 (thioesterase)
- Similar to Cytidylate kinase (UMP-CMP kinase), partial (Ornithorhynchus anatinus)
- Annexin VII

#### Apoptosis and cell cycle

### Energy-consuming processes

#### Vesicles, endocytosis, intracellular trafficking

- RAB37, member RAS oncogene family (Homo sapiens)
- Globlastoma amplified sequence (NipSnap2)

#### Molecular chaperones

- Peptidylprolyl isomerase B (Danio rerio)
- Heat shock 70 kDa protein 9B (Gallus gallus)
- Heat shock cognate 70 kDa (Carassius auratus)

#### Microtubular skeleton

- Tubulin beta-1 chain (Gadus morhua)
- Tropomyosin (Takifugu rubripes)
- Fast tropomyosin (Scyliorhinus retifer)
- Cytoskeletal tropomyosin 4 (Coturnix coturnix)
- Tropomodulin 2 (Danio rerio)

#### Mitochondrial cristae structure

- Mitofilin (mitochondrial inner membrane protein) (Homo sapiens)

*Continued*
membrane (EZR ↑ H1) or to the extracellular matrix at focal adhesions (NEBL ↓ H2, 27), and an actin stabilizer (MARCKS ↓ H2). The latter is of particular interest here, because proteasomal degradation of actin-stabilizing MARCKS is involved in transient, translation-independent preconditioning in cultured neurons. This short-term effect was linked to a decrease in the number of postsynaptic dendritic spines, decreased NMDA-type glutamate receptor availability, and decreased glutamate/NMDA excitotoxicity (65). In accord with this mechanism, we also observed decreases in the membrane-associated GTPase DN3M after both H1 and H2. Recent work demonstrated that DN3M associates with metatropic glutamate receptors in postsynaptic dendrite spines, and, more importantly, that changes in DN3M expression lead to remodeling of dendritic spine morphology, possibly via interactions with the actin cytoskeleton (25). Morphological changes in dendrites (a.k.a., synaptic structural plasticity) are known to be involved in memory and learning (29), and they contribute to dynamic and reversible responses of neurons to stimuli (69, 72).

On the basis of our data, we hypothesize that an analogous mechanism of dendrite spine retraction contributes to prolonged excitotoxicity prevention following hypoxic exposure (especially after H2, but not anoxia) in the epaulette shark cerebellum. Because the density of dendritic spines is positively correlated with the availability of AMPA glutamate receptors (73), such structural plasticity would represent a potentially powerful preconditioning mechanism for enhanced damage prevention following repeated hypoxia in the epaulette shark brain. Further work is needed to clarify its significance in this context.

The evidence for cytoskeletal and membrane changes in response to hypoxia was not limited to the cerebellum. The rectal gland exhibited upregulation after H1 of three spots identified as tropomyosins, as well as TMOD2. Together, these proteins regulate the interaction of actin and myosin as well as the length of actin filaments. Three proteins from the ezrin/radixin/moesin family of membrane-actin cytoskeletal cross-linkers exhibited the opposite expression pattern compared with the tropomyosins, decreasing after H1 but returning to control levels after H2. Together, these results may indicate transient reorganization of rectal gland cytoskeletal components after one episode of hypoxia. A recent study demonstrated rapid morphological changes at both the tissue and cellular levels following feeding in the dogfish rectal gland (63), and we hypothesize that analogous mechanisms contribute to the epaulette shark rectal gland’s response to oxygen limitation here.

Other proteins whose expression changed in the rectal gland were placed in the “vesicles, endocytosis, and intracellular trafficking” category and play putative roles in membrane dynamics as well. However, the implications for membrane plasticity are less clear in this tissue. GBAS (a.k.a. nipsnap2; ↑ A1) belongs to a protein family implicated in vesicular transport (8), although its closest relative nipsnap1 also associates with the inner mitochondrial membrane and perhaps the electron transport chain.

Proteins were grouped by putative function based on Gene Ontology annotations and manual review of the literature. All columns as in Table 4. Expression data for unidentified protein spots can be found in Supplemental Table S5.
(14). Similarly, transient downregulation of IMMT after H1 may indicate mitochondrial cristae structural reorganization and could have implications for metabolite (ADP) flux (38, 121). The function of RAB37 (↓ H1), a GTPase associated with secretory granules in pancreatic beta cells (7), is also unknown in this context.

**Greater Efficiency in Metabolic Coupling Rather Than Decreased Capacity**

Evidence for a decrease in aerobic or anaerobic energy production capacity, one aspect of canonical metabolic depression, was limited in the present study. The data for plasma lactate levels suggest an organismal shift to anaerobic ATP production, particularly during anoxia, consistent with a continued need for cellular energy supply in the absence of oxygen. Circulating lactate at A1+2 h increased >10-fold over control (C1+2 h) levels; epaulette shark plasma lactate also peaked at anoxia +2 h in a previous study (13). If sustained throughout anaerobic episodes, this increase could compensate for much of the 10- to 18-fold loss of efficiency when relying solely on anaerobic compared with aerobic glucose metabolism (33). Because there was no A1+A2+2 h treatment (A1+A2 lactate increased ~3-fold over C1+2 h), we could not determine whether the response to a second anoxic episode involved a compensatory decrease in organismal lactate production. While both brain and rectal gland exhibited some proteome adjustments consistent with changes in glycolytic flux after oxygen limitation, these responses were not consistent. Importantly, the proteomic and metabolic responses to anoxia in other tissues that constitute a greater fraction of the organismal metabolic rate remain to be investigated.

Interestingly, the proteomic data for rectal gland indicate adjustments that could increase the efficiency of shuttling of pyruvate, the end product of cytosolic glycolysis, to the mitochondria, facilitating its complete oxidation via the tricarboxylic acid (TCA) cycle and electron transport chain (ETC). Two spots identified as PDHB (E1 of the pyruvate dehydrogenase complex) increased significantly after both A1 and H2 in rectal gland, and a third component of the complex (DBT, the E2 catalytic center) also increased after H2. We have no concrete explanation for these results, but it is tempting to speculate that enhanced PDH complex activity would tightly couple cytosolic pyruvate production to mitochondrial oxidative phosphorylation when oxygen is limiting but still available (i.e., during hypoxia). Such tight coupling could also help to prevent buildup of H+ ions from net ATP hydrolysis during conditions when oxygen is limiting (32). It should also enhance aerobic metabolic efficiency in the gland during the reoxygenation period following anoxia. No analogous changes in the pyruvate dehydrogenase complex were noted in brain.

While our data offer little evidence for systematic changes in anaerobic metabolism following oxygen limitation, previous work in this species clearly demonstrated a decrease in the normoxic aerobic metabolic rate after exposure to multiple hypoxic episodes (89). This result implies decreased flux through the TCA cycle and/or the ETC, where oxygen serves as the terminal electron acceptor. Our data indicate only minimal adjustments in the TCA cycle for the tissues examined, while the patterns for the ETC are again tissue- and treatment-specific. No TCA enzymes changed abundance following any treatment in cerebellum. In the ETC, two spots identified as ATP5A1 (ETC complex V) decreased after anoxia, possibly indicating reduced oxidative phosphorylation capacity. ATP5B increased after H1 in cerebellum, while NDUFV2 (ETC complex I) decreased after H2. ATP5B contains the catalytic site of ATP synthase (complex V), possibly indicating a more efficient coupling of ATP synthesis to the H+ ion gradient across the inner mitochondrial membrane after one episode of hypoxia (analogous to the PDHB argument in rectal gland above). Notably, expression of ATP5B also increased in two chemical preconditioning models (1, 90) and after hepatic ischemic preconditioning in mouse, where it was proposed to offer protection from reperfusion injury (118). In rectal gland, the only observed changes in TCA cycle enzymes were an increase in IDH2 after H1 (see below) and a decrease in SDHB (also a component of ETC complex II) after H1. Other ETC proteins in complex I (NDUFS1, NDUFS3) and complex V (ATP5A1) were also downregulated to varying degrees after A1 or H1, but generally not after H2. It is possible that the organismal decreases in aerobic metabolic rate observed in Ref. 89 reflect the contributions of other tissues and/or require more than two episodes of hypoxia to take effect.

Although the proteomic results for aerobic metabolic pathways in the tissues examined are somewhat ambiguous, the organismal response to repeated episodes of hypoxia reflected continued reliance on oxidative metabolism; there was no evidence for decreased ventilatory rate during H2 compared with H1. In addition, epaulette shark hemoglobin appears to have a high affinity for oxygen, and the critical oxygen tension further decreases after preexposure to hypoxia (89). These results suggest that physiological responses to hypoxia in this species work to maintain oxygen delivery to the tissues during successive hypoxic episodes, rather than depressing aerobic metabolism. Notably, lactate levels still increased more than threefold at H1+H2+2 h relative to C1+2 h. This indicates that anaerobic pathways must be invoked to supplement aerobic ATP production during hypoxia, allowing epaulette sharks to sustain metabolic activity under these conditions.

**Limited Evidence for Decreased Energy Demand**

Reduced energy demand is the other hallmark of species that survive prolonged oxygen limitation by invoking metabolic depression, but, as for energy production, our data provide inconclusive evidence for a decrease in energy-consuming processes following episodic oxygen limitation. Transcription and translation (i.e., protein synthesis) account for a large fraction of the cellular energy budget (88). Several proteins involved in these functions were significantly downregulated in brain following hypoxia, but not anoxia. SFRS5 (↓ H1 and H2) participates in mRNA splicing, as does RBM8A (↓ H2) (94). The opposite pattern of SFRS5 expression appeared in ischemic mouse brain (93). The tRNA transporter SARS (↓ H2) and the 60S ribosome subunit RPLP0 (↓ H1) are necessary for (ATP-dependent) translation at the ribosome. Although these results may be consistent with decreased protein synthesis in brain following hypoxia and recovery, these proteins represent only a small fraction of the large complexes that carry out transcription and translation. We found no evidence
for suppression of these two processes in the rectal gland following any of the treatments.

Other proteins involved in protein degradation via the proteasome decreased in cerebellum following one or more episodes of hypoxia and reoxygenation, possibly indicating reduced protein turnover in this tissue (49). PSMC2 (\( \downarrow \) H1 and H2) forms part of the 19S regulatory proteasome cap, where it unfolds and deubiquitinates proteins prior to their degradation. The 19S proteasome lid component PSMD11 (\( \downarrow \) H2) plays a critical role in assembly and stabilization of the entire 26S proteasome as well as in its substrate specificity (35, 92). Whether these expression changes are sufficient to globally or selectively suppress proteasome-dependent proteolysis remains to be determined. In rectal gland, no proteasome constituents were regulated by oxygen limitation.

Ion pumping represents another energetically costly process for cells, particularly in the brain and rectal gland. Our data provide no evidence for a preconditioning effect on the major monovalent cation-pumping protein Na\(^+\)/K\(^+\)-ATPase in either tissue. We interpret this finding as evidence for the critical cellular and organismal need to maintain ionic homeostasis and membrane polarity during episodic oxygen limitation (112) and, equally as important, during reoxygenation. Accordingly, our plasma electrolyte data reveal no pervasive loss of ionic homeostasis following any treatment (at +2 h or +24 h). However, we cannot rule out posttranslational control of Na\(^+\)/K\(^+\)-ATPase during the episodes of oxygen limitation (3, 61).

**Substrate Switching and More Efficient Energetic Shuttling to Support Continued Metabolic Activity**

The proteome responses following episodic hypoxia, but not anoxia, revealed divergent patterns of substrate utilization for sustained ATP production in cerebellum and rectal gland. The glycolytic enzyme GAPDH, which either increases (30) or is used as a housekeeping gene (91) in mammalian brain ischemia models, decreased in epaulette shark cerebellum after both H1 and H2, possibly indicating a decreased reliance on glucose metabolism. Our data for cerebellum suggest a compensatory increase in ketone body catabolism after hypoxia (OXCT1 \( \uparrow \) H1), while the decrease in MCCC2 (\( \downarrow \) H1 and H2) implies decreased reliance on branched-chain amino acid catabolism (especially leucine). Ketone bodies, including \( \beta \)-hydroxybutyrate, are a favored substrate in the brain and in elasmobranch tissues in general (2, 16). In contrast, the rectal gland exhibited decreased abundance of OXCT1 after H1 and H2, implying decreased reliance on ketone bodies in this tissue. The rectal gland was recently demonstrated to rely heavily on glucose as a substrate, while \( \beta \)-hydroxybutyrate could only supplement glucose-fueled salt secretion (110). Interestingly, both tissues exhibited an increase in enzymes related to fatty acid metabolism following hypoxia, but not anoxia (odd chain fatty acids: PCCA \( \uparrow \) H2 in brain; short-chain fatty acids: ACADS \( \uparrow \) H1 and \( \uparrow \) H2 in rectal gland). Although fatty acids are not considered an important fuel in elasmobranchs (2), their use may supplement ATP production at critical times. It will be informative to measure circulating and cellular levels of these alternative fuels in the epaulette shark in this context.

Another prominent feature in rectal gland, but not brain, was the increased abundance of several proteins involved in intracellular shuttling of energetic equivalents. The increase in mitochondrial GOT2 after anoxia suggests enhanced unidirectional flux of cytosolic NADH produced in glycolysis to the ETC via the malate aspartate shuttle (15). Such an increase would likely be more important during reoxygenation than anoxia per se, because oxygen limitation would prevent oxidation of NADH and result in an increased mitochondrial NADH:NAD\(^+\) ratio (120). It is also known that GOT2 aids the recovery from ischemia in rat heart (80). Two spots identified as creatine kinase B (CKB) increased after episodic hypoxia, one after H1 and the other after both H1 and H2. Creatine kinase plays a role in the phosphocreatine circuit, which buffers intracellular ATP levels and transfers mitochondrial ATP equivalents throughout the cell (109). Given that this enzyme physically associates with Na\(^+\)/K\(^+\)-ATPase (28) and plays a role in teleost gill ionoregulation (47), increased CKB following episodic hypoxia may represent increased efficiency of localized ATP buffering to support continued ion pumping function in rectal gland.

**Detoxification of Metabolic Byproducts and Amelioration of Oxidative Stress, Particularly in Rectal Gland**

Episodic hypoxia induced a number of proteins in the epaulette shark rectal gland for coping with the toxic byproducts of continued metabolic activity during subsequent episodes of oxygen limitation and reoxygenation. These results generally conform with the lack of change in rectal gland Na\(^+\)/K\(^+\)-ATPase activity following all treatments and the recent “transport at any cost” theory of rectal gland function (110). A variety of proteins involved in the amelioration of oxidative stress increased in rectal gland. Hypoxia elicited more compensatory proteome adjustments in this category than anoxia, consistent with recent evidence for ROS generation [particularly superoxide radical (O\(_2\)\(^\cdot\)) and hydrogen peroxide (H\(_2\)O\(_2\))] at the electron transport chain during hypoxia (54). The increase in FTH1 after H1 suggests enhanced cellular sequestration of Fe\(^{2+}\) ions, most likely to prevent oxygen radical formation via Fenton chemistry (62). The observed increases in COQ5 (\( \uparrow \) H1 and H2) indicate elevated ubiquinone biosynthesis. Ubiquinone has antioxidant properties (55), and it acts as a cofactor for mitochondrial uncoupling proteins that reduce oxygen radical generation at the ETC (20, 34).

Several other rectal gland proteins have more indirect roles in combating oxidative stress. The top-ranking VIP score and most highly upregulated protein for rectal gland, and the only identified protein to change expression in all three treatments in either tissue, was glyoxalase domain containing 5 (GLOD5; \( \uparrow \) 4.3 \( \times \) A1, \( \uparrow \) 3.4 \( \times \) H1, \( \uparrow \) 2.9 \( \times \) H2). Although this particular gene product has not been well characterized, the glyoxalase system detoxifies methylglyoxal, a cytotoxic byproduct of glycolysis (111). Overexpression of glyoxalase I in rat kidneys rescues them from oxidative stress and apoptosis (48). The deubiquitinating enzyme UCHL3 also increased significantly in rectal gland after one episode of hypoxia. This enzyme may function in recycling ubiquitin that was incorrectly bound to GSH or other nucleophiles (68). Finally, although we classified mitochondrial IDH2 (\( \uparrow \) H1) as a TCA cycle enzyme, we propose that it plays more of a role in alleviating mitochondrial oxidative stress than in TCA flux in rectal gland in response to
hypoxia. Recently, IDH2’s ability to supply NADPH reducing equivalents (possibly for recycling of GSSG to GSH) was linked to prevention of ROS-mediated apoptosis in response to several stressors (43, 52, 96).

In parallel to the IDH2 hypothesis following hypoxia, expression changes of several rectal gland glycolytic enzymes following anoxia may indicate compensatory adjustments in the oxidative stress response more than changes in energy production. GAPDH abundance increased significantly in rectal gland after anoxia, but glycolytic capacity downstream of production. GAPDH abundance increased significantly in response to several stressors (43, 52, 96).

The oxidative stress response more than changes in energy following anoxia may indicate compensatory adjustments in the oxidative stress response to subsequent episodes of anoxia and reoxygenation.

In the rectal gland, episodic oxygen limitation also induced several proteins with chaperone functions related to oxidative stress. Overexpression of the endoplasmic reticulum (ER) chaperone PPIB (↑ A1 and H1), implicated in adaptation to metabolic and oxidative stress in the kidney (40), inhibits cytosolic Ca^{2+} loading, ROS production, and apoptosis resulting from an excess of unfolded proteins (44). The Parkinson’s disease-associated chaperone PARK7 (↑ H1) is induced by oxidative stress and protects rat neurons from such insult (53).

Finally, induction by oxidative stress (H_{2}O_{2}) of the chaperone DNAJB11/Hsp40 (↑ H1 in rectal gland), a regulator of HSP70 activity, is highly conserved (108). Surprisingly, two members of the HSP70 superfamily were downregulated in rectal gland following anoxia and/or hypoxia (HSPA9 ↓ all treatments; HSPA8 ↓ H1 and H2), suggesting that these chaperones play a less of a role in this tissue relative to those that were upregulated.

Proteomic adjustments related to oxidative stress were far less common in cerebellum, including the only highly conserved protein antiquitin (ALDH7A1; ↓ H2) and the inner mitochondrial membrane chaperone PHB (↑ H2) (70). We also observed a decrease in HSPA4L (also known as osmotic stress protein 94) after both H1 and H2 in the cerebellum. In contrast, heat shock proteins are constitutively expressed at high levels in anoxia-tolerant turtle brains and further upregulated to varying degrees during anoxia (83), and they play a role in ischemic preconditioning in mammals (50). Whether ROS production is inherently lower in brain than in rectal gland, antioxidant expression is higher, or tissue-specific changes in substrate utilization lead to varying levels of ROS formation remains to be determined. The total glutathione levels in epaulette shark brain were unexceptional compared with tissues of other fishes (76, 102). In addition, our glutathione data do not indicate dramatic enhancement of the pool of this antioxidant at ±24 h, although the consistent changes across control and experimental groups imply that our handling controls worked as designed. Further data, such as activities of glutathione peroxidase and reductase, are needed to determine whether the glutathione system plays a role in the epaulette shark’s response to episodic oxygen limitation.

**Modulation of Cellular Prosurvival Signaling**

It is important to note that transient metabolic depression during episodes of anoxia or hypoxia in epaulette sharks could be mediated by translation-independent mechanisms, particularly posttranslational modifications of existing proteins via signaling cascades, which would not be detected with our experimental design. In our dataset, the abundance of several key signaling proteins changed after episodic hypoxia, but not anoxia, with potential ramifications for metabolic regulation during future periods of oxygen limitation. Adjustments in brain included a transient decrease in the cAMP-dependent protein kinase (PKA) regulatory subunit PRKAR2B after H1, followed by an increase after H2 in another regulatory subunit PRKAR1A. These regulatory subunits bind to the catalytic PKA subunit (PKAc) with different affinities (PRKAR1A binds more tightly, 105) and different sensitivities to cAMP (PRKAR2B more responsive to cAMP, 17). The changing ratio of PRKAR1A to PRKAR2B following episodic hypoxia suggests modulation of PKAc pathways downstream of G protein-coupled receptors (GPCRs), which include glutamate receptors. Furthermore, the GPCR complex member GNAO1, involved in cAMP stimulation of PKA and downstream effects on mitochondrial activity (12), also decreased after H1. PKAc has numerous other downstream targets, including Na^{+}/K^{+}-ATPase (3). In contrast, PRKAR1A decreased after H1 in rectal gland, suggesting tissue-specific regulation of the PKA signaling pathway in response to hypoxia. Both tissues also exhibited modulation of several 14-3-3 proteins, which bind phosphorylated signaling proteins and regulate their activity. Both PKA and 14-3-3 signaling cascades represent promising avenues for future research in this species.

An intriguing result in the rectal gland was the upregulation of cystathionase (CTH) after H2. CTH’s production of the signaling gas hydrogen sulfide (H_{2}S) is inversely related to oxygen levels in fish (74), and H_{2}S formation has very recently been shown to rescue cells from ischemia/reperfusion injury and apoptosis in several tissues (21, 23, 106). The role of CTH and H_{2}S in hypoxia response in the epaulette shark rectal gland may represent a promising area for further study.

Ultimately, a cell’s fate in the face of stressful conditions depends in part on the net balance of pro- and antiapoptotic signaling. In this study, we found no biochemical evidence for increased caspase 3/7-dependent apoptosis in cerebellum or rectal gland at any point following anoxia or hypoxia, nor have previous studies found morphological evidence for apoptosis following repeated hypoxia in the epaulette shark brain (67, 84). We have also found no late apoptotic DNA fragmentation in epaulette shark gill in the same treatments using a TUNEL assay (W. W. Dowd, unpublished observations). These results suggest net prosurvival signaling in epaulette shark tissues during and after periods of oxygen limitation. Accordingly, several proteins potentially involved in antiapoptotic signaling pathways were induced after episodic hypoxia, but not anoxia, in cerebellum. The transcription factor FUBP1 (↑ H2) is required at the promoter of c-myc, an oncogene that typically promotes cell cycle progression. We hypothesize that increased...
FUBP1 after multiple hypoxic episodes may stimulate a form of c-myc “insurance” against programmed cell death (36); c-myc may play an analogous role in anoxia-tolerant turtles (26). Two other major players in apoptosis are p53 (generally proapoptotic) and nuclear factor kappa B (NF-κB, generally antiapoptotic). The ubiquitin-conjugating enzyme UBE2D2 († H1 H2) ubiquitinates both p53 and inhibitor of NF-κB (IkB) and targets them for proteasomal degradation, thus releasing antiapoptotic NF-κB for translocation to the nucleus (24). RAN († H1) also plays a role in shutting of NF-κB to the nucleus (115), and its overexpression protects glioblastoma cells from apoptosis (116). In rectal gland, only the mitochondrial ALDH4A1 increased after H1 here, where it may act as a negative regulator of p53-dependent ROS formation to inhibit apoptosis (119).

Summary

Despite the fact that captive epaulette sharks naïve to oxygen limitation exhibit exceptional intrinsic tolerance at tropical temperatures (13), our data suggest that these sharks enhance a number of compensatory molecular mechanisms in response to episodic hypoxic or anoxic exposure. These tissue and treatment-specific responses should increase the resistance to subsequent episodes of oxygen limitation, and we interpret these compensatory changes as a preconditioned phenotype. In contrast to studies of prolonged hypoxia and anoxia tolerance in other species, episodic oxygen limitation and recovery did not induce sustained metabolic depression in the epaulette shark tissues examined. Rather, these sharks apparently adjust their physiology and proteome to sustain metabolic activity and cope with the consequences, especially following exposure to hypoxia. Because the majority of the observed proteome responses were relatively minor (very few expression ratios >2 or <0.5), the epaulette shark’s preconditioning phenotypic response appears quite subtle. We interpret these patterns as fine-tuning of a constitutively expressed suite of compensatory mechanisms that likely evolved under strong selective pressure in the epaulette shark’s coral reef habitat.

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DISCLOSURES

No conflicts of interest, financial or otherwise, are declared by the author(s).

REFERENCES

PAEPAULTE SHARK HYPOXIA AND ANOXIA PROTEOMICS


