Exploring the limit of metazoan thermal tolerance via comparative proteomics: thermally induced changes in protein abundance by two hydrothermal vent polychaetes

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Temperatures around hydrothermal vents are highly variable, ranging from near freezing up to 300°C. Nevertheless, animals thrive around vents, some of which live near the known limits of animal thermotolerance. Paralvinella sulfincola, an extremely thermotolerant vent polychaete, and Paralvinella palmiformis, a cooler-adapted congener, are found along the Juan de Fuca Ridge in the northwestern Pacific. We conducted shipboard high-pressure thermotolerance experiments on both species to characterize the physiological adaptations underlying P. sulfincola’s pronounced thermotolerance. Quantitative proteomics, expressed sequence tag (EST) libraries and glutathione assays revealed that P. sulfincola (i) exhibited an upregulation in the synthesis and recycling of glutathione with increasing temperature, (ii) downregulated nicotinamide adenine dinucleotide (NADH) and succinate dehydrogenases (key enzymes in oxidative phosphorylation) with increasing temperature, and (iii) maintained elevated levels of heat shock proteins (HSPs) across all treatments. In contrast, P. palmiformis exhibited more typical responses to increasing temperatures (e.g. increasing HSPs at higher temperatures). These data reveal differences in how a mesotolerant and extremely thermotolerant eukaryote respond to thermal stress, and suggest that P. sulfincola’s capacity to mitigate oxidative stress via increased synthesis of antioxidants and decreased flux through the mitochondrial electron transport chain enable pronounced thermotolerance. Ultimately, oxidative stress may be the key factor in limiting all metazoan thermotolerance.

Keywords: proteomics; hydrothermal vents; thermotolerance; oxidative stress; Paralvinella

1. INTRODUCTION

Physiological adaptations to thermal stress are ubiquitous among all organisms. While prokaryotes have a known upper thermal limit of at least 122°C [1], metazoans have a much lower thermal tolerance, with 45–47°C as the currently accepted temperatures at which metazoans can maintain homeostasis [2] (though unicellular eukaryotic fungi are known to grow at 60–62°C [3]). Protein thermal stability and degradation [4], mitochondrial dysfunction [5], membrane and structural instability [2], and limitations in gas transport have all been implicated as possible modes of physiological failure in eukaryotes at elevated temperatures [6–8].

There have been numerous studies to date on metazoan thermotolerance (see [9,10]). The few that have focused on highly thermotolerant animals such as desert ants and hot spring ostracods have largely examined their response to acute thermal exposure [11,12]. Recently, some studies have employed proteomics to examine responses to chronic thermal stress in mesotolerant animals [7,13]. However, there remains a limited amount of biomolecular data for extremely thermotolerant metazoans [14]. It remains to be determined how highly thermotolerant organisms respond to chronic thermal exposure, and which physiological or biochemical adaptations enable them to ameliorate physiological perturbations that arise at higher temperatures.

Deep-sea hydrothermal vents are home to some of the most thermotolerant animals known. These include the polychaetes Alvinella pompejana and Paralvinella sulfincola. To date, numerous studies have investigated the thermal tolerance of A. pompejana, beginning with the observation that A. pompejana lives upon 81°C substratum [15]. Subsequent to that, and in contrast to the in situ observations, in vitro research on A. pompejana has suggested that key enzymes and structural components are not stable after chronic exposure to elevated temperatures [16]. A recent study of A. pompejana protein expression via two-dimensional gel electrophoresis compared physiological responses with different oxygen concentrations.
[14], though its response to chronic exposure remains unconstrained. Unfortunately, A. pompejana are not easily amenable to in vivo experimentation [17], making it difficult to address chronic thermal tolerance in this species. *Paralvinella sulfincola* thrives on hydrothermal sulphides in the northwest Pacific. *Paralvinella sulfincola* are found at temperatures up to 88.5 °C [18], and in vivo laboratory studies of *P. sulfincola* have experimentally demonstrated the broadest known range of thermal tolerance in metazoans (5–48 °C [19,20]). *Paralvinella palmiformis*—a closely related congener—is also found in these environs but exhibits markedly different thermal tolerances [19,20]. Both are amenable to in vivo recovery and laboratory experimentation, which affords the unique opportunity to elucidate the biochemical responses of meso- and thermotolerant metazoans to chronic thermal exposure in a comparative phylogenetic context.

To better understand the biochemical mechanisms that underlie extreme thermal tolerance, we present data from a series of in vivo high-pressure laboratory experiments in which we characterized the chronic thermal tolerance of both species, and measured quantitative changes in protein expression of live *P. sulfincola* and *P. palmiformis* over their thermal range, including temperatures near each species’ ultimate incipient lethal temperature (ULT; the temperature beyond which 50% of the population cannot survive indefinitely [21,22]). A normalized expressed sequence tag (EST) library served as the database for the proteomic analyses (EST libraries are inherently qualitative, so all data shown here are from the quantitative proteomic analyses unless otherwise noted; see electronic supplementary material, methods). We also quantified differences in antioxidant abundance between both species and across thermal treatments. These data reveal notable differences in chronic thermal tolerance, and significant differences in protein abundance and antioxidant concentrations between these two congers, primarily related to mitigating oxidative stress across their thermal ranges and at their respective ULTs. The results of this study provide the first direct empirical evidence that oxidative stress may be the primary stressor at *P. sulfincola’s* upper thermal limit and illustrate the means by which *P. sulfincola* mitigates this stress.

2. RESULTS AND DISCUSSION

(a) Constraining *P. sulfincola* and *P. palmiformis* chronic thermal tolerance

To constrain chronic thermal tolerance of these two paralvinellid species, we conducted a series of incubations in which worms were acclimated to 20 °C overnight, then incubated over a range of temperatures, for up to 12 h at each temperature (these acclimation and exposure periods are longer than previous studies [19,20], but are also limited by the logistical considerations of working aboard a ship). Dissolved oxygen concentrations were monitored during these incubations to ensure that oxygen concentrations were not limiting owing to insufficient fluid flow through the chambers (lower fluid flows increase fluid residence time and can result in decreased oxygen concentrations in the chambers). These data reveal that *P. sulfincola* is capable of surviving chronic (greater than or equal to 12 h) exposure up to 45 °C. We thus define 45 °C as *P. sulfincola’s* ULT, as we observed mortality during chronic exposure to higher temperatures (figure 1). *Paralvinella palmiformis* also had a comparably wide thermal range, surviving chronic exposure to temperatures between 0 °C and 38 °C. *Paralvinella palmiformis*, however, was not as thermotolerant as *P. sulfincola*, and died within 6 h of incubation at 40 °C (figure 1). Collectively, these reveal that species exhibit broad thermal ranges, but only *P. sulfincola* is capable of chronic exposure to 45 °C (well above that tolerated by most eukaryotes). Based on these data, we selected discrete temperatures that represent the lower, middle and upper temperature ranges of both *P. sulfincola* and *P. palmiformis*, and conducted a series of thermal exposure experiments at these temperatures on freshly collected and thermally acclimated assemblages of *P. sulfincola* and *P. palmiformis* (see §4). Tissues recovered from these worms were used for the proteomic analyses and glutathione quantification.

(b) Differences in expression of molecular chaperones

A total of 27 molecular chaperones and co-chaperones were examined in our analysis, representing members of all detected HSPs. We observed that *P. sulfincola* exhibited elevated levels of all major molecular chaperones, even those previously categorized as inducible, over all temperature treatments including the control (figure 2), while *P. palmiformis* exhibited higher chaperone production primarily near the ULT. Molecular chaperones such as HSPs mitigate thermal stress primarily by minimizing protein dysfunction through catalysing nascent protein folding in the endoplasmic reticulum (ER), and reforming
misfolded proteins [23,24]. While many chaperones are constitutively expressed, a large number of chaperones are upregulated during periods of cellular stress—the so-called inducible forms [25].

Key chaperone families are discussed in the paragraphs below. Their abundance and changes in response to treatments are presented in table 1a, and their statistical significance is presented in electronic supplementary material, tables S1 and S2.

(i) Heat shock protein 70
The 70 kDa heat shock proteins (HSP70 family) are highly conserved across domains of life [24]. Multiple isoforms in the family are constitutive, while others are induced by thermal stress [25,26]. In P. sulfincola, GRP75, a member of the HSP70 family and a homologue to the human constitutive mitochondrial protein HSPA9 [27], exhibited the highest abundance of all molecular chaperones across all treatments. In contrast, GRP75 expression in P. palmiformis exhibited a moderate increase in abundance at 38°C (relative to the cooler thermal treatments). The apparently constitutive and elevated abundance of HSP70 proteins in P. sulfincola suggest they may be kept abundant to cope with the rapid changes in temperature typically encountered by this species, which includes maintaining physiological function near the organism’s UILT.

(ii) Heat shock protein 90 chaperone proteins
Although less well characterized than the HSP70 family, HSP90s are flexible dimer ATPases that bind to a variety of proteins, including steroid hormone receptors, transcription factors and protein kinases [28,29]. The HSP90 protein GRP94 (a luminal protein associated with the ER [30]) was detected in P. sulfincola and exhibited constitutive expression across all treatments (with a probability of differential expression, or Pr(DE) of 0.11). GRP94 was also observed in P. palmiformis, and its abundance appears to have increased with temperature (12°C → 38°C—log 1.55, Pr(DE) 0.66). Co-chaperones including FKBP52 and HOP were observed in both P. sulfincola and P. palmiformis. FKBP52 exhibited a highly significant increase with temperature in both worms (Pr(DE) 1.00). HOP, which modulates HSP70/90 interactions, was also upregulated with temperature in both P. sulfincola and P. palmiformis at their highest treatments (P. sulfincola, 10°C → 45°C—log 0.98, Pr(DE) 0.82; P. palmiformis, 12°C → 38°C—log 0.98, Pr(DE) 0.51). The HSP90 activator AHA1 was substantially upregulated at 45°C in P. sulfincola (10°C → 45°C—log 3.58, Pr(DE) 1.00) but not in P. palmiformis. Conspicuously, the HSP90 inhibitor CDC37 remained constant in P. sulfincola and significantly decreased in P. palmiformis in higher thermal regimes. The patterns observed here suggest that HSP90 is constitutively expressed in P. sulfincola, though this activity is apparently being regulated in both species. These observations are also consistent with the aforementioned hypothesis that P. sulfincola maintains a pool of molecular chaperones to cope with acute temperature fluctuations.

(iii) Heat shock protein 60 and heat shock protein 27 chaperone proteins
HSP60 is a mitochondrial molecular chaperone known to facilitate thermal tolerance in eukaryotes [31]. HSP60 was very abundant in both species and across all treatments. This trend was mirrored in the HSP60 co-chaperone, HSP10, which assists HSP60 in protein folding during periods of stress [32]. These findings suggest that both species maintain pools of HSP60 and HSP10 to mitigate damage to mitochondrial proteins.

The small 27 kDa heat shock protein (sHSP), found throughout cellular compartments and the cytosol, responds to both thermal and oxidative stress by binding to damaged or misfolded proteins and forming reservoirs for other chaperones to correctly refold or initiate proteolytic degradation [33]. It is also known to upregulate key enzymes in the glutathione pathway [33,34]. HSP27 was highly abundant across all treatments in P. sulfincola. However, HSP27 increased only at the highest temperature in P. palmiformis (12°C → 38°C—log 2.09, Pr(DE) 1.00). We posit that the abundance of HSP27 relates to oxidative stress response and the glutathione pathway (discussed in detail below).

(iv) Foldases
Foldases catalyse rate-limiting steps in protein folding and play a key role in the cellular ‘unfolded protein response’ (i.e. a response to an accumulation of unfolded and misfolded proteins in the ER, which aims to restore normal function by halting protein translation and signalling the production of molecular chaperones involved in protein folding [35]). Foldases involved in the unfolded protein response were detected in both species. Of note, the foldase PDIA1, a protein-thiol oxidoreductase that acts as both a chaperone and a foldase [35,36], was constitutive and highly abundant across all treatments in P. sulfincola (Pr(DE) 0.001). In P. palmiformis, PDIA1 abundance increased as a function of temperature (Pr(DE) 0.914).
Table 1. Paralvinella sulfincola and P. palmiformis protein abundance during thermal exposure, including select P. sulfincola and P. palmiformis proteins as discussed in the text. EST is the (i) isotig or (c) contig identifier for each protein. Log change is the shift in abundance between treatments (i.e. log Δ P. palmiformis—12→21 is the protein log-fold change between P. palmiformis treatments 12°C and 21°C). Counts are combined between all three technical replicates and normalized to treatment library sizes. Red boxes indicate a significant increase in protein abundance (Pr(DE) > 0.9); blue boxes indicate a significant decrease in protein abundance (Pr(DE) > 0.9).

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Collectively, these data on chaperone and foldase abundance suggest that *P. sulfincola* maintains pools of chaperones across all thermal regimes. We posit, however, that the representation and abundance of these chaperones do not fully explain the observed differences in thermotolerance. Indeed, if the abundance of chaperones and foldases was the sole factors in conferring extreme thermotolerance, then *P. palmiformis* could be expected to have a thermal tolerance similar to that of *P. sulfincola* because the proportion of chaperones (relative to total protein) between these two species was equivalent at their respective highest thermal treatments. We further posit that constitutively elevated HSP abundances in *P. sulfincola* are more likely to be a reflection of its ecological niche *in situ*, enabling it to survive acute, rapid shifts in temperatures caused by its proximity to hot vent fluid, but do not alone explain their chronic thermal tolerance.

### (c) Response to oxidative stress

In both species, at elevated temperatures, proteins related to the mitigation of oxidative stress exhibited the largest increases in abundance. In mitochondria, superoxide (O$_2^-$) is generated in complexes I/III during respiration, while other reactive oxygen species (ROS), such as in the hydroxyl radical (HO•) and uncharged hydrogen peroxide (H$_2$O$_2$), are produced in the outer and inner membranes (see [37,38]). At normative conditions, mitochondria consume more than 90 per cent of all cellular O$_2$, while also producing the majority of ROS [39]. However, studies have shown that elevated temperatures increase oxidative stress in mesotolerant euakaryotes [6,40,41], as mitochondrial respiration rates increase at elevated temperatures [4].

In *P. sulfincola*, glutathione appears to play a prominent role in mitigating oxidative stress. Glutathione (L-$\gamma$-glutamyl-L-cysteinylglycine, or GSH) is a tripeptide thiol that is the primary non-protein antioxidant in metazoans. Found in millimolar concentrations in many metazoans, GSH mitigates oxidative stress by chemically reducing hydrogen peroxide and other toxic compounds [37,42]. The enzyme glutathione peroxidase (GPx, 1.11.1.9) catalyses this reduction, yielding glutathione disulphide (GSSG). GSSG is reverted back to GSH by glutathione reductase (GSR, enzyme commission (EC) 1.8.1.7). Regulation of GSH metabolism serves as an indicator of cellular oxidative stress levels [42]. Because cysteine is the required peptide for *de novo* GSH synthesis, and the rare amino acid selenocysteine is required for the synthesis of GPx, increases in cysteine (and in particular selenocysteine) are good indicators for increases in GSH cycling, and were observed in *P. sulfincola* treatments (table 1; electronic supplementary material, tables S1 and S2).

Figure 3 depicts glutathione metabolism, including the enzymes GPx and GSR, in *P. sulfincola* and *P. palmiformis* as a function of thermal exposure. Glutathione peroxidase 3 (GPx-3, a cytosolic form), which reduces lipid hydroperoxides to alcohols and hydrogen peroxide to water, showed significant increases in abundance at both the medium- and high-temperature treatments in *P. sulfincola*, as well as at the highest temperature treatment in *P. palmiformis*. Notably, *P. sulfincola* also significantly increases its GSR protein abundance, while *P. palmiformis* significantly decreases it at higher temperatures.

Cystathionine $\beta$-synthase (CBS, EC 4.2.1.22), central to both cysteine and selenocysteine synthesis, exhibited the largest increase in abundance as a function of temperature in *P. sulfincola* (*P. sulfincola*, 10°C→45°C—log 5.74, Pr(DE) 1.00). *Paralvinella palmiformis* CBS exhibited a comparable response (*P. palmiformis*, 12°C→38°C—log 5.55, Pr(DE) 1.00).

Selenide water dikinase (SelD, EC 2.7.9.3) and gamma-glutamylcysteine synthetase (GCS, EC 6.3.2.2) are two ATP-dependent, rate-governing steps within the glutathione pathway that were detected in both *P. sulfincola* and *P. palmiformis* (figure 3). SelD is essential for *de novo* synthesis of selenoproteins, and increased in *P. sulfincola* at both 30°C and 45°C. No differences in abundance were observed in *P. palmiformis* across all treatments. GCS is the rate-governing step in the production of GSH [42], and in *P. sulfincola* exhibited a steady increase in abundance with temperature. In *P. palmiformis*, however, GCS was only detected in the 38°C treatment (12°C→38°C—log 3.81). These data demonstrate that GSH is being synthesized at higher rates in relation to increasing thermal stress in both species, though the abundance of proteins related to GSH production are much higher in *P. sulfincola*. We further suggest that the differences may be indicative of mitochondrial dysfunction and uncoupling in *P. palmiformis*, possibly owing to lipid peroxidation from increasing ROS activity, as has previously been observed in cold-water marine molluscs exposed to heat stress and functional hypoxia [6,39].

To further investigate the effect of thermal and oxidative stress on GSH production and recycling, total GSH (tGSH) levels and GSH/GSSG ratios (the ratio of the reduced and oxidized forms) were measured from *P. sulfincola* and *P. palmiformis* worms exposed to the medium- and high-temperature treatments (see electronic supplementary material, figure S1). tGSH concentrations in *P. sulfincola* were about half those observed in *P. palmiformis*. There were no measurable differences in the GSH/GSSG ratio among *P. sulfincola* worms across all thermal treatments. However, in higher thermal treatments, *P. palmiformis* exhibited a twofold decrease in the pool of tGSH. Also, the GSH/GSSG ratio in *P. palmiformis* exhibited more than a threefold drop at those same temperatures. These data reveal that *P. palmiformis* is less effective at reducing GSSG at 38°C. These data further suggest that *P. sulfincola* is able to continue GSH resynthesis (via GSSG reduction) near its UILT, allowing it to better mitigate oxidative damage at elevated temperatures.

Superoxide dismutase (SOD, EC 1.15.1.1) is another ubiquitous enzyme that is responsible for catalysing the reduction of O$_2^-$ to H$_2$O$_2$. There are two forms of this metalloprotein; Cu/Zn SOD (iso1g03775) is primarily found in the cytosol, whereas Mn SOD (iso1g06674) is located in the mitochondria [43]. *Paralvinella sulfincola* showed no differences in the abundance of either SOD across all treatments, though *P. palmiformis* exhibited significant increases in both Mn SOD (Pr(DE) 0.997) and Cu/Zn SOD (Pr(DE) 0.999) with increasing temperature (see electronic supplementary material, tables S1 and S2).

### (d) Shifts in oxidative phosphorylation

In euakaryotes, oxidative phosphorylation within the electron transport chain is responsible for the majority of ATP
specifically, a large portion of ROS is generated by nicotinamide adenine dinucleotide (NADH) dehydrogenase (complex I). Succinate dehydrogenase (complex II) may not contribute directly to ROS formation, but it funnels electrons to complex III, which does produce ROS. *Paralvinella sulfincola* exhibited a significant reduction in abundance of these two key proteins with increasing temperature (NADH dehydrogenase from 10°C to 45°C—log 2.01, and succinate dehydrogenase from 10°C to 45°C—log 1.00; see electronic supplementary material, tables S1 and S2). In *P. palmiformis*, the decrease in NADH dehydrogenase was less pronounced (12°C to 38°C—log 0.48), and there was an increase in abundance of succinate dehydrogenase with temperature (12°C to 38°C—log 1.53). The observed patterns of NADH dehydrogenase are in contrast to a previous study of the mussels *Mytilus galloprovincialis* (a warmer-adapted species) and *Mytilus trossulus* (a cooler-adapted species) [7]. In the previous study, *M. trossulus* exhibited a downregulation in the NADH dehydrogenase with increased heat stress, while no change was observed in *M. galloprovincialis*. While the reasons underlying these differences among these organisms remain unknown, our data are consistent with the hypothesis that *P. sulfincola* is actively repressing ROS formation at high temperatures by lessening endogenous generation via the electron transport chain (ETC), and depending more heavily on anaerobic respiration at elevated temperatures.

(e) **Global changes in protein abundance and emerging hypotheses**

Global, quantitative proteomic analyses further provide an opportunity to examine how other biochemical systems respond to experimental treatments. To this end, iPath [44] was used to map changes in global protein abundance within 139 KEGG metabolic pathways (summarized in electronic supplementary material, figures S3 and S2–d). In general, we observed that *P. palmiformis* exhibited a greater number of significant changes in protein abundance during the treatments. In *P. palmiformis*, increases in temperature...
from 12°C to 21°C led to significant increases in proteins associated with nitrogen metabolism, amino acid synthesis, glycan synthesis and nucleotide metabolism, while proteins involved in pentose phosphate synthesis and pyrimidine metabolism decreased during this treatment (see supplementary material, figure S2c). Increasing temperature from 21°C to 38°C led to significant increases in proteins involved with the tricarboxylic acid cycle (TCA cycle), xenobiotic degradation, fatty acid biosynthesis, nitrogen and amino acid metabolism. Proteins associated with nucleotide metabolism, in general, decreased during this latter treatment (see electronic supplementary material, figure S2d). In P. sulfincola, increases in temperature from 10°C to 30°C led to significant increases in proteins involved in glycan biosynthesis, taurine and hypotaurine metabolism, as well as cysteine and methionine metabolism, while starch metabolism and amino acid biosynthesis were downregulated during this treatment (see electronic supplementary material, figure S2a). Increasing temperature from 30°C to 45°C led to significant increases in proteins involved in fatty acid biosynthesis, and extensive decreases in the TCA and associated cycles and nitrogen metabolism (see electronic supplementary material, figure S2b).

Most striking (and relevant to this study) were the differences in the abundance of TCA and pentose phosphate proteins observed during these treatments. At temperatures near their UILT, P. sulfincola decreased the abundance of TCA cycle proteins, while P. palmiformis increased the abundance of TCA cycle proteins. The opposite pattern was observed in the pentose phosphate pathway, with P. sulfincola and P. palmiformis exhibiting increased and decreased protein abundance, respectively. Although one cannot infer the degree to which metabolic flux changes through a pathway from these or any proteomic/transcriptomic data [45,46], these trends are consistent with a decreased emphasis on aerobic respiration in P. sulfincola (the TCA cycle feeds reducing equivalents in the form of NADH into the ETC, which exacerbates ROS production) and an increased need for reducing equivalents to maintain antioxidant activity (the pentose phosphate pathway produces NADPH, which is used to produce GSH [47]). We speculate that the observed broad-scale upregulation by P. palmiformis at elevated temperatures may reflect the organism’s effort to maintain homeostasis. Future studies should be aimed at addressing these hypotheses to further our understanding of how increasing temperature influences metabolism in these two species. Moreover, future studies should examine the degree to which duration of exposure, as well as acclimation to warmer temperatures, influences the patterns of protein abundance observed in these species. It also remains to be determined how other environmental factors (H₂S, CO₂, and pH) impact thermal tolerance. Finally, studies might also consider the influence of post-translational protein modifications to physiological functions at elevated temperatures.

4. MATERIAL AND METHODS

(a) Animal collection and experimental apparatus

Paralvinella sulfincola and P. palmiformis ‘palm worms’ were collected from hydrothermal vents in the Main Endeavour field along the Juan de Fuca Ridge (47°57’ N, 129°5’ W) at a depth of 2200 m during the R/V Atlantis expedition 15–34 in July 2008. Organisms were collected by the DSV
Alvin on dives no. 4409–4423, using a multi-chamber suction sampler or an insulated sample recovery box. Upon recovery to 1 atm, worms were transferred to a 4°C cold room and visually sorted based on segment number and gill morphology. Aggregations of mucus and minerals were removed from the animals before transfer into a flow-through high-pressure aquaria system (see electronic supplementary material, methods and figure S4).

(b) Experimental design

Though critical thermal maxima (CT\textsubscript{max}) of both species and thermal preference of \textit{P. sulfincola} were previously examined \cite{19,20}, we augmented these data to better establish \textit{P. sulfincola} and \textit{P. palmiformis} chronic thermal tolerance (figure 1). A total of 165 \textit{P. sulfincola} and 108 \textit{P. palmiformis} were used in this chronic (greater than or equal to 12 h) thermal tolerance study. Less than 5 per cent of individuals died during treatments; these deaths were typically attributable to damage during recovery and handling. Based on these data, three temperatures were chosen that span the chronic thermal tolerance range of each species (\textit{P. sulfincola} = 10\textdegree C, 30\textdegree C and 45\textdegree C; \textit{P. palmiformis} = 12\textdegree C, 21\textdegree C and 38\textdegree C; figure 1). These temperatures were chosen to represent the organisms’ protein profiles at their lower, nominal and upper (approaching UI\textsubscript{LT}) temperature regimes. At each thermal exposure treatment, six to nine worms were maintained at constant pressure and temperature for greater than 12 h for global protein expression analysis. To minimize the effects of collection and handling, worms were acclimated in each system at 20\textdegree C for 12 h prior to experimentation. At the conclusion of each trial, the chambers were quickly depressurized, and worm health was assessed by looking for embolisms, motor dysfunction or other damage that might have arisen from thermal exposure or other experimental handling. Healthy worms were selected, and their branchiae and body tissues were separated and flash frozen in liquid nitrogen for subsequent protein extraction.

(c) Protein extraction

Gills from three \textit{P. sulfincola} and three \textit{P. palmiformis} per treatment were excised, weighed on an electronic balance (Mettler Toledo, Columbus, OH), and placed into sterilized 0.5 ml glass micropestles (Wheaton, Millville, NJ) containing 24 \textmu l of 20 mM Tris pH 7.5 buffer and 6 \textmu l Protease Inhibitor Cocktail (PIC; Sigma-Aldrich, St Louis, MO). Tissue was homogenized until complete dissociation, then centrifuged at 1000g for 5 min. For protein extraction, 0.5 mg gill tissue was extracted via a modified Laemmli protein extraction \cite{51}. Extractions were then loaded in separate lanes onto 4 to 20 per cent precast Precise Protein Gels (Pierce Inc., Rockford, IL) with blank lanes between samples. The gels were bathed in a Tris–hydroxymethyl piperazineethanesulphonic acid (HEPES)–sodium dodecyl sulphate (SDS) buffer solution and electrophoresed for 45 min at 100 V. After electrophoresis, gels were rinsed and stained for 3 h using the Novex colloidal comassie blue dye (Invitrogen, Carlsbad, CA). Gels were visualized using a Gel Logic 100 gel imaging system (Kodak, Rochester, NY) and sub-sectioned into six fragments according to protein size. Three biological replicates from each treatment were pooled into one sample per fragment; total gel surface area did not exceed 1 cm\textsuperscript{2}. The pooled gel sub-sections were then washed with 1 ml of 50 per cent acetonitrile and frozen at −20\textdegree C prior to analysis.

(d) Protein analyses by tandem mass spectrometry

A total of 36 pooled samples (two species incubated at three temperatures fractionated into six equal sections) were reduced, carboxamidomethylated and digested with trypsin. Resulting peptides from each sample were analysed in triplicate using microcapillary reverse-phase high-performance liquid chromatography (HPLC) directly coupled to the nano-electrospray ionization source of a ThermoFisher LTQ-Orbitrap XL (replicate 1) or LTQ-Orbitrap Velos (replicates 2 and 3) hybrid mass spectrometer (\mu LC/MS/MS). The Orbitrap repetitively surveyed m/z range from 395 to 1600, while data-dependent MS/MS spectra on the 20 most abundant ions in each survey scan were acquired in the linear ion trap. MS/MS spectra were acquired with a relative collision energy of 30 per cent, 2.5 Da isolation width and recurring ions dynamically excluded for 60 s. Preliminary evaluation of peptide-spectrum matches (PSMs) employed the SEQUEST algorithm with a 30 ppm mass tolerance against the \textit{P. sulfincola} EST library (see electronic supplementary material, methods and table S4) and NCBI databases. Spectral counting is a method of quantification in which one compares the number of MS/MS spectra acquired for a particular protein across multiple LC-MS/MS datasets. Increases and decreases in relative protein abundance are reflected in corresponding increases and decreases in spectral counts for that protein \cite{52,53}. PSMs were accepted with mass error less than 3.0 ppm and score thresholds to attain an estimated false discovery rate of less than 1 per cent using a reverse decoy database strategy and a custom version of the Harvard Proteomics Browser Suite. A total of 172 122 peptide spectra were identified with an average of 14.6 amino acids per sequence, with MS/MS spectra populating 1296 referenced proteins (see electronic supplementary material, table S5).

(e) Glutathione measurements

Total GSH and GSSG levels were determined using the Glutathione Assay Kit (Cayman Chem, Ann Arbor, MI). Spectrophotometric readings were taken for 30 min using a Spectramax Plus\textsuperscript{384} spectrophotometer (Molecular Devices, Sunnyvale, CA). Internal standards were run with total GSH and GSSG experimental treatments, and standard curves were built from the endpoint readings.

(f) Data analyses and statistics

Bayesian analyses were used to determine statistically significant changes in protein abundance among the experimental treatments. The details of these analyses may be found in the electronic supplementary material.

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