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Differential expression profiling of heat stressed tardigrades reveals major shift in the transcriptome

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ABSTRACT
Tardigrades are renowned for their extreme stress tolerance, which includes the ability to endure complete desiccation, high levels of radiation and very low sub-zero temperatures. Nevertheless, tardigrades appear to be vulnerable to high temperatures and thus the potential effects of global warming. Here, we provide the first analysis of transcriptome data obtained from heat stressed specimens of the eutardigrade Ramazzottius variornatus, with the aim of providing new insights into the molecular processes affected by high temperatures. Specifically, we compare RNA-seq datasets obtained from active, heat-exposed (35 °C) tardigrades to that of active controls kept at 5 °C. Our data reveal a surprising shift in transcription, involving 9634 differentially expressed transcripts, corresponding to ~35% of the transcriptome. The latter data are in striking contrast to the hitherto observed constitutive expression underlying tardigrade extreme stress tolerance and entrance into the latent state of life, known as cryptobiosis. Thus, when examining the molecular response, heat-stress appears to be more stressful for R. variornatus than extreme conditions, such as desiccation or freezing. A gene ontology analysis reveals that the heat stress response involves a change in transcription and presumably translation, including an adjustment of metabolism, and, putatively, preparation for encystment and subsequent diapause. Among the differentially expressed transcripts we find heat-shock proteins as well as the eutardigrade specific proteins (CAHS, SAHS, MAHS, BvLEAM, and Dsup). The latter proteins thus seem to contribute to a general stress response, and may not be directly related to cryptobiosis.

1. Introduction
Tardigrada constitutes a group of microscopic animals with a well-defined cephalic region and a trunk bearing four pairs of legs (Møbjerg et al., 2018). Distributed worldwide, tardigrades, also known as water bears, populate highly diverse microhabitats in marine, freshwater and terrestrial ecosystems (McInnes and Pugh, 2018; Nelson et al., 2018). Tardigrades divide into two major evolutionary lineages represented by eutardigrades and the more diverse heterotardigrades (Jørgensen et al., 2018; Morek et al., 2020). They are renowned for their extreme stress tolerance (Møbjerg and Neves, 2021) and their ability to enter cryptobiosis—a reversible state of suspended animation—which provides them with an extreme tolerance towards severe environmental conditions, including complete desiccation, high external solute concentrations, extremely low temperatures, high toxicant concentrations and oxygen depletion (Clegg, 2001; Guidetti et al., 2011; Møbjerg et al., 2011; Hygum et al., 2017; Sørensen-Hygum et al., 2018). Importantly, tardigrades need to be surrounded by a film of water, in order to remain in their physiologically active state. During desiccation, when tardigrades may lose more than 95% of their water (Crowe, 1972), they contract longitudinally, retract head and legs, while forming a quiescent and barrel shaped form, the so-called “tun” (Crowe and Madin, 1974; Bertolani et al., 2004; Halberg et al., 2013).

Although their ability to endure a range of extreme conditions, tardigrades were recently reported to be sensitive to prolonged periods of high temperature exposure in active as well as desiccated tun states (Neves et al., 2020a, 2020b). Specifically, thermotolerance experiments on desiccated tuns of Ramazzottius variornatus, a eutardigrade frequently found in European roof-gutters, revealed a decrease in the 50% mortality temperature from 82.7 °C, over 63.1 °C to 56.1 °C

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following 1 h, 24 h and 1 week of high temperature exposures, respectively (Neves et al., 2020a, 2020b). In fully hydrated and thus metabolically active specimens, the 50% mortality temperature was 37.1 °C. Nevertheless, tardigrades occurring in limno-terrestrial and marine tidal environments are most likely side-stepping the damaging effects of heat exposure by undergoing desiccation induced cryptobiosis and, hence, being able to survive summer periods in a warmer Earth (Sørensen-Hygum et al., 2018; Vecchi et al., 2021). Accordingly, it remains to be seen to which extent the ongoing global warming will affect tardigrade populations in general.

The limited thermostolerance of *R. varieornatus* suggests an interference of high temperatures with tardigrade cell and macromolecule function (Mohr and Neves, 2021). Hence, bioprotectants and other macromolecules, which otherwise provide tardigrades with their extraordinary stress tolerance, might indeed be denatured following prolonged heat exposure. A variety of proteins with proposed cytoprotective function are found among tardigrades. For instance, heat shock proteins (HSPs) and late embryogenesis abundant (LEA) proteins may act as chaperones and molecular shields, preventing protein aggregation during desiccation, and HSPs might further participate in the repair of damaged proteins during rehydration (Schill et al., 2004; Schokraie et al., 2010). Noteworthy, the expression of HSP70 seems to be induced by heat stress in the eutardigrade *Milnesium inceptum* (Jönsson and Schill, 2007) and in *Milnesium inceptum* the expression of a small α-crystallin heat-shock protein gene (*Mt-shsp17.2*) increased 779-fold following heat stress (Reuner et al., 2010). More recently, it has been suggested that heat-soluble “tardigrade-unique proteins” (Yamaguchi et al., 2012; Tanaka et al., 2015; Hashimoto et al., 2016), also known as tardigrade-specific intrinsically disordered proteins, may provide structural stabilization to eutardigrade cells during desiccation (Bootby et al., 2017; Yagi-Utsumi et al., 2021). These intrinsically disordered proteins, including SAHS (secretory abundant heat-soluble proteins), CAHS (cytoplasmic abundant heat-soluble proteins), MAHS (mitochondrial abundant heat-soluble proteins) and RvLEAM (a group 3 LEA proteins) and late embryogenesis (LEA) proteins may provide structural stabilization to eutardigrade cells during desiccation (Boothby et al., 2017; Yagi-Utsumi et al., 2021). These intrinsically disordered proteins, including SAHS (secretory abundant heat-soluble proteins), CAHS (cytoplasmic abundant heat-soluble proteins), MAHS (mitochondrial abundant heat-soluble proteins) and RvLEAM (a group 3 LEA protein located in the mitochondria) are missing in the heterotardigrade lineages (Kamirli et al., 2019; Murai et al., 2021). Eutardigrades furthermore express an unstructured nuclear protein, Dsp, which supposedly binds to nucleosomes and protects DNA from reactive hydroxyl radicals (Chavez et al., 2019; Minguez-Toral et al., 2020).

Importantly, transcriptome analyses have shown that extremotolerant tardigrade species seem to constitutively express their genes during cryptobiotic survival (Hashimoto et al., 2016; Murai et al., 2021). Here, we investigate the transcriptomic response to heat stress in *R. varieornatus* with the aim of providing new insights into the molecular processes affected by high temperatures. Specifically, we compare the transcriptome of physiologically active, heat-exposed (35 °C) tardigrades to that of physiologically active controls kept at 5 °C. To the best of our knowledge, the current study presents the first analysis of RNA-seq datasets obtained from heat stressed tardigrades. Our transcriptome profiling reveals a surprising shift in the transcriptome content of 9634 differentially expressed genes, corresponding to ~35.6% of the transcriptome. Among the very large number of differentially expressed genes, several code for eutardigrade specific proteins such as SAHS, CAHS, MAHS, Dsp, and RvLEAM. In addition, two stress-related heat-shock proteins were found to be upregulated, namely a HSP70 and a small HSP, while other heat-shock proteins were found to be downregulated. A gene ontology analysis suggests a change in metabolism, transcription, and possibly translation as a response to the heat-shock. Our findings support previous observations revealing that high temperatures (> 30 °C) are considerably more stressful for the eutardigrade *R. varieornatus* than extreme conditions, such as desiccation (Hashimoto et al., 2016) or freezing at very low sub-zero temperatures (~ 80 °C) (unpublished observations).

2. Methods

A graphical representation of the methods used to evaluate the effect of exposing fully hydrated, active tardigrades to 35 °C as compared to 5 °C is provided in Fig. 1. In the figure, grey text and arrows refer to a reference transcriptome available from the European Nucleotide Archive under the project PRJEB47629, which was used for mapping the reads and quantifying differential gene expression.

2.1. Sampling of specimens, experimental setup and total RNA extraction

Specimens of Ramazzottius varieornatus Bertolani and Kinchin, 1993 (Eutardigrada, Ramazzottiidae) were obtained from sediment collected from a root gutter in Nivå, Denmark (55°56′36.53″ N, 12°30′00.90″ E), in February 2018, and identified using the original taxonomic description of the species (cf. Bertolani and Kinchin, 1993). The sediment sample was frozen under wet conditions, stored at −20 °C for several months and subsequently transferred to −80 °C. Two subsamples were thawed, diluted in ultrapure water (Millipore Milli-Q® Reference, Merck, Darmstadt, Germany) and acclimated to 5 °C in May and June 2020, respectively. The subsamples were recurrently examined for tardigrades between June and July 2020 with the aid of a stereomicroscope. Highly active, adult (large) specimens of *R. varieornatus* showing spontaneous movement of legs were collected from the subsamples and transferred to embryo dishes using hand pulled Pasteur pipettes, at room temperature (RT; 21–25 °C). Subsequently, three groups of ca. 100–180 tardigrades each – i.e. three replicates – were transferred to Eppendorf tubes with 1.5 ml of moderately hard reconstituted water (MRHW; Khamma et al., 1997) and then exposed to 35 °C using a AccuBlock Digital Dry Bath heating system (Labnet International, Edison, NJ) for approximately 24 h. After the heat shock, the tardigrades were briefly checked for locomotor activity and processed to extract total RNA. In addition, three groups of ca. 100–180 tardigrades each were kept refrigerated (5 °C) for 24 h as controls, and then processed to extract total RNA. Total RNA was extracted using the RNeasy Plus Universal Mini Kit (Qiagen, Hilden, Germany) according to the instructions of the manufacturer, with the following modifications: for disruption of the tardigrade cuticle, the specimens were processed in BeadBug™ tubes (Benchmark Scientific, Sayreville, NJ) containing 2.3-mm and 1-mm diameter zirconia/silica beads during 2 × 1 min homogenization at 50 Hz. Initial quantity and quality of the extracted RNA were evaluated with a NanoDrop® ND-1000 spectrophotometer (Peqlab Biotechnologie GmbH, Erlangen, Germany).

2.2. mRNA library construction and sequencing

The total RNA samples were sent to BGI Europe A/S for library preparation and sequencing. Concentration and purity of each RNA sample were assessed by BGI using an Agilent 4200 (Agilent Technologies, CA, USA) before mRNA was selected, reverse transcribed and sequenced using the DNA nanoball sequencing (DNBseq-G400) platform. A total of 376,485,664 (100 bp) paired end reads were generated (Table 1). We subsequently assessed the quality of the raw RNA-Seq data produced by BGI with the software FastQC (Andrews, 2010) and the produced reads were then subject to trimming of adapters and low-quality stretches using AdapterRemoval v2.0 (Schubert et al., 2016). The quality of the reads was once again evaluated after adapter trimming using FastQC to ensure that the adapter sequences were properly trimmed (Ewels et al., 2016).
2.3. Differential expression analyses

We used a reference transcriptome of the Danish population of *R. varieornatus*, recently made available (ENA PRJEB47629), for mapping sequence reads and a pseudo-aligner, Salmon version 1.1.0 (Patro et al., 2017), to quantify transcript abundance in each of the six RNA-Seq datasets (i.e. the three datasets from heat-exposed (35°C) tardigrades and three datasets from non-exposed tardigrades, kept at 5°C). The NumbRead metric estimated by Salmon was subsequently used for three differential expression analyses using the Bioconductor packages DESeq2 (Love et al., 2014), EdgeR (Robinson et al., 2010) and Limma (Ritchie et al., 2015). Only genes that were expressed in more than half of the RNA-Seq datasets were considered in the differential expression analyses (19,456 quantified transcripts were included, whereas 133 did not meet this criterion). The quasi-likelihood F-test was used for EdgeR, as this test is effective at controlling the false discovery rate (FDR) and

![Fig. 1](image.png)

Graphical representation of the methods used to evaluate the effect of exposing fully hydrated, active tardigrades to 35°C. Active specimens were randomly pooled into groups (3 × ca. 100–180) at room temperature (RT), either exposed to 35°C or kept at 5°C (controls) for 24 h and subsequently used to extract total RNA for differential gene expression analyses. Sequenced reads were mapped against a reference transcriptome. Grey text and arrows concern the preparation of this reference transcriptome (ENA PRJEB47629), which was assembled and annotated for another study.

### Table 1

Summary data for RNA samples generated from active heat exposed (replicate #1-3) and control (control #1-3) specimens of *Ramazzottius varieornatus*.

<table>
<thead>
<tr>
<th>Samples</th>
<th>No. of specimens</th>
<th>Total RNA extracted (µg)</th>
<th>Concentration of total RNA (ng/µL)</th>
<th>No. of raw reads</th>
<th>No. of clean reads</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control #1</td>
<td>154</td>
<td>0.5</td>
<td>18.7</td>
<td>58,096,968</td>
<td>49,625,720</td>
</tr>
<tr>
<td>Control #2</td>
<td>180</td>
<td>0.8</td>
<td>28.1</td>
<td>67,004,602</td>
<td>52,186,994</td>
</tr>
<tr>
<td>Control #3</td>
<td>176</td>
<td>0.7</td>
<td>25.0</td>
<td>59,392,932</td>
<td>49,817,626</td>
</tr>
<tr>
<td>Replicate #1</td>
<td>180</td>
<td>0.8</td>
<td>28.4</td>
<td>59,600,522</td>
<td>49,777,180</td>
</tr>
<tr>
<td>Replicate #2</td>
<td>102</td>
<td>0.6</td>
<td>21.8</td>
<td>63,121,868</td>
<td>49,943,728</td>
</tr>
<tr>
<td>Replicate #3</td>
<td>152</td>
<td>0.8</td>
<td>29.0</td>
<td>69,268,772</td>
<td>49,803,520</td>
</tr>
</tbody>
</table>

![Fig. 2](image.png)

Differential transcript expression analyses. Venn diagram as determined by the three differential gene expression analyses: DESeq2, EdgeR and Limma. The blue circle represents Limma + voom, the red circle represents the DEGs identified by EdgeR, and the green circle represents DESeq2. Numbers in overlapping zones represent the DEGs identified in more than one analysis. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

NumbRead metric estimated by Salmon was subsequently used for three differential expression analyses using the Bioconductor packages DESeq2 (Love et al., 2014), EdgeR (Robinson et al., 2010) and Limma (Ritchie et al., 2015). Only genes that were expressed in more than half of the RNA-Seq datasets were considered in the differential expression analyses (19,456 quantified transcripts were included, whereas 133 did not meet this criterion). The quasi-likelihood F-test was used for EdgeR, as this test is effective at controlling the false discovery rate (FDR) and...
most appropriate for experiments with few replicates. An FDR adjusted p-value of $\leq 0.05$ was used to determine whether a gene was up- or down-regulated in the heat-exposed ($35 ^\circ C$) tardigrades as compared to the controls kept at $5 ^\circ C$. The Venn diagram (Fig. 2) and MA-Plot (Fig. 3) were generated in R (R Core Team, 2016) and edited with Illustrator CS5 version (Adobe Inc.).

2.4. Enrichment analysis

We performed a Gene Ontology (GO) enrichment analysis on the transcripts identified in the differential expression analyses using TopGO version 2.42.0 (Alexa and Rahnenfuhrer, 2020), with GO terms derived from the annotated reference transcriptome (ENA PRJEB47629). Fisher’s exact test was implemented to compute the significant GO enrichments at the $\alpha$-level of $p \leq 0.05$.

3. Results

Gene expression profiles for the heat-exposed ($35 ^\circ C$) and the non-exposed ($5 ^\circ C$) tardigrades revealed a substantial number of differentially expressed genes (DEGs), independently of the method used to perform the differential expression analysis. Indeed, a total of 11,015 different DEGs were identified across the three methods (DESeq2, EdgeR and Limma): 9634 of these transcripts (i.e., 35.6% of the 27,054 assembled unigenes in the reference transcriptome) were found by all three methods as shown in the Venn diagram represented in Fig. 2. Limma, EdgeR and DESeq2 exclusively identified 199, 178 and 30, respectively. 32 DEGs were shared only by EdgeR and DESeq2, 87 DEGs by Limma and DESeq2 and 855 DEGs were shared only by EdgeR and Limma. Out of the 9783 DEGs found by DESeq2, 4901 of the transcripts were upregulated and 4882 were downregulated in respect to the controls kept at $5 ^\circ C$. The Venn diagram (Fig. 2) and MA-Plot (Fig. 3) were generated in R (R Core Team, 2016) and edited with Illustrator CS5 version (Adobe Inc.).

<table>
<thead>
<tr>
<th>Method of Differential Expression Analysis</th>
<th>DESeq2</th>
<th>EdgeR</th>
<th>Limma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upregulated genes</td>
<td>4901</td>
<td>5382</td>
<td>5489</td>
</tr>
<tr>
<td>Downregulated genes</td>
<td>4882</td>
<td>5317</td>
<td>5286</td>
</tr>
</tbody>
</table>

Table 2

Differential gene expression analyses of RNA-seq data generated from control and heat exposed tardigrades with the number of upregulated and downregulated genes found by each of the three methods (DESeq2, EdgeR, and Limma).

Table 3

List of enriched subcategories with corresponding GO IDs, identified by TopGO following the differential gene expression analyses considering an FDR $\leq 0.05$.

<table>
<thead>
<tr>
<th>GO ID</th>
<th>GO subcategories</th>
<th>Fold enrichment</th>
</tr>
</thead>
<tbody>
<tr>
<td>GO:0016887</td>
<td>ATPase activity</td>
<td>1.007919366</td>
</tr>
<tr>
<td>GO:0046982</td>
<td>Protein heterodimerization activity</td>
<td>1.529411765</td>
</tr>
<tr>
<td>GO:0043565</td>
<td>Sequence-specific DNA binding</td>
<td>1.198156682</td>
</tr>
<tr>
<td>GO:0003735</td>
<td>Structural constituent of ribosome</td>
<td>1.218879668</td>
</tr>
<tr>
<td>GO:0008061</td>
<td>Chitin binding</td>
<td>1.25394272</td>
</tr>
<tr>
<td>GO:0008137</td>
<td>NADH dehydrogenase (ubiquinone) activity</td>
<td>1.420765027</td>
</tr>
<tr>
<td>GO:0003743</td>
<td>Translation initiation factor activity</td>
<td>1.256830601</td>
</tr>
</tbody>
</table>

Proteins that have been shown previously to be involved in stress responses in tardigrades (e.g., heat shock and tardigrade disordered proteins), were sought after among the annotated DEGs. In our analysis, 12 Heat shock transcripts were identified as differentially expressed by all three methods. Two of these transcripts were upregulated, whereas the remaining 10 were downregulated. The two upregulated transcripts were a HSP70 and a small HSP (annotated as HSP67Bb), while the downregulated transcripts encompassed one HSP90, three HSP70 isoforms, one HSP68, one HSP60, and four small HSPs. Among the upregulated transcripts, two SAHS genes, i.e. SAHS1 and SAHS2, were identified by all three methods. Seven CAHS transcripts were annotated and identified as DEGs by all three methods (i.e. CAHS1, CAHS2, CAHS3, CAHS68135, CAHS77611, CAHS86272 and CAHS89226). All the identified CAHS transcripts were upregulated, with the exception of CAHS68135, which was downregulated. In addition, one MAHS transcript was identified, and all three analyses found it to be upregulated in the heat-exposed tardigrades when compared to the controls kept at $5 ^\circ C$. Likewise, the group 3 LEA gene RvLEAM, a mitochondrial protectant like MAHS, was found to be differentially expressed and

![Fig. 3](https://example.com/fig3.png) Differential transcript expression analysis. MA-Plots of DESeq2 data, showing the log2 fold change against the mean of normalized counts. Differentially expressed genes are highlighted in blue: DESeq2 found 4901 upregulated and 4882 downregulated genes in tardigrades kept at 35 $^\circ C$ for 24 h as compared to controls (5 $^\circ C$). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
upregulated. Furthermore, a Dsup isoform was found to be upregulated in the differential expression analyses.

4. Discussion

4.1. Differential gene expression

Our results revealed that major transcriptional perturbation occurs in active specimens of *R. varieornatus* as a response to temperature increment. The latter is in clear contrast to the constitutive expression observed during cryptobiotic survival in both eu- and heterotardigrades (Hashimoto et al., 2016; Murai et al., 2021). Importantly, high temperatures seem to transform tardigrade physiology, with 35.6% of the transcriptome being differentially expressed following 24 h heat stress. Among the 9634 differentially expressed transcripts identified by all three methods used to perform differential expression analyses (i.e., DESeq2, EdgeR and Limma), we found 12 DEGs related to heat-shock proteins, seven CAHS, two SAHS, one MAHS, one LEA and one Dsup gene. Of these transcripts, all DEGs coding for tardigrade intrinsically disordered proteins were upregulated, except for one CAHS. In contrast, only two of the heat-shock related transcripts were upregulated (a small heat-shock protein and a HSP70), whereas the remaining 10 were downregulated. We hypothesize that the small heat-shock protein could bind to denatured proteins preventing aggregation and that the HSP70 isoform could assist in renaturing dysfunctional proteins at the expense of ATP (Willsie and Clegg, 2002; Jagla et al., 2018; Hibbsman et al., 2020), thus counteracting the effect of heating on tardigrade proteins. The upregulation of Dsup indicates that this damage suppressor may have an important role in protecting DNA during heat-stress. Specifically, Dsup has been shown to protect chromosomesal DNA by binding to nucleosomes thereby shielding the DNA from reactive hydroxyl radicals (Hashimoto et al., 2016; Chavez et al., 2019; Minguez-Toral et al., 2020), thus counteracting the effect of heating on tardigrade proteins.

ATPase activity is among the GO molecular function categories (Table S3) that were found to be enriched among the differentially expressed transcripts in heat-exposed versus non-exposed *R. varieornatus*. ATPase activity represents a common molecular function utilized by a wide array of enzymes that hydrolyze ATP, providing the necessary energy to catalyze chemical reactions and e.g. transport ions and other molecules against their electrochemical gradients. NADH dehydrogenase (ubiquinone) activity is another overrepresented category in our analysis. This enrichment could reflect a change in mitochondrial inner membrane respiratory chain complex I activity. The mitochondrial respiratory complex I accepts electrons from NADH + H+ and transfers these to the ubiquinone, while simultaneously moving protons out of the mitochondrial matrix, thereby fueling the electrochemical gradient that drives ATP production (Whitehouse et al., 2019). Thus, both the overrepresentation of ATPase activity and NADH dehydrogenase (ubiquinone) activity indicate a change in the metabolism of the heat stressed tardigrades. Given the higher temperature, it seems likely that chemical energy is required at a larger scale in the heat-exposed, as compared to the non-exposed, tardigrades to catalyze cellular processes (e.g., DNA and RNA synthesis, intracellular signaling, transmembrane transport, etc.). Future studies could provide important further insight into the metabolic changes that take place in tardigrade cells following temperature increments.

Two other overrepresented GO molecular function categories, protein heterodimerization activity and sequence-specific DNA binding proteins, have a common denominator: transcription factors. Heterodimerization, a result of two non-identical proteins associating through non-covalent interaction to form a complex, is common among transcription factors, receptors and some enzymes (Jordan and Devi, 1999; Funnell and Crossley, 2012; Singh and Jois, 2018). Moreover, sequence-specific DNA binding proteins may also represent transcription factors that regulate gene expression (Suter, 2020). As such, these two overrepresented categories are in agreement with a change in transcription as a result of heating.

The overrepresented gene ontologies show a pattern revolving around a change, not only in transcription, but also translation. Indeed, overrepresented GO categories also include structural constituents of ribosome and translation initiation factor activity. Therefore, our analysis further suggests that the translation of proteins is being regulated as a response to the heating stimulus. This further supports the observation that heat-exposed tardigrades change their metabolism, transcription and translation to match the increase in temperature and counteract its effects.

A key feature in the body plan of tardigrades is the presence of a cuticular cuticle (Baccetti and Rosati, 1971; Bussers and Jeuniaux, 1973; Greven et al., 2016). As chitin binding is also an overrepresented GO term (Table 3), we hypothesize that heat-exposed *R. varieornatus* may be responding to the unfavorable change in their environment by preparing for encystment and diapause. Specifically, the observed regulation in transcription of chitin binding proteins could indicate that the heat-exposed tardigrades are in the process of shedding their cuticle. Encystation in tardigrades involves formation of new cuticle layers that are shed and subsequently provide protection against unfavorable environmental conditions (Guidetti and Mobberg, 2018; Mobberg and Neves, 2021). Encystment is not normally encountered in strong cryptobionts, such as *R. variocornatus*. Nevertheless, cysts of a *Ramazzottius* species were reported by Murray (1907). Interestingly, cyst formation
was also reported in the strong cryptobiont Echiniscoides sigismundi, a marine tidal heterotardigrade, following experiments on osmotic stress tolerance (Clausen et al., 2014). Thus, although encystment is not the default strategy for strong cryptobionts, it may represent an alternative strategy, when they are presented with a stress factor (e.g. high temperatures or fluctuating salinities) that does not by itself induce cryptobiosis. Clearly more comprehensive studies on R. varieornatus and other strong cryptobionts as well as species well-known to form cysts are required to clarify whether high temperatures induce encystment.

Our results show that 

R. varieornatus

reacts to heat-stress with a change in transcription and translation and with an adjustment of other genes with importance for tardigrade stress responses. Under heat stress. A more detailed inspection of the large number of differentially expressed genes will be required, as well as investigations into the influence of habitat. In order to explore possible climate-correlated thermotolerance adaptations, latitudinal comparisons of thermotolerance in species and populations inhabiting different climate regions is needed.

4.3. Concluding remarks

Our findings indicate that heat-shock proteins as well as eutardigrade specific proteins are involved in the heat stress response of

R. varieornatus.

Yet, these proteins only represent a minute fraction of the thousands of genes found to be differentially expressed following heat stress. A more detailed inspection of the large number of differentially expressed transcripts found in our analyses would likely uncover other genes with importance for tardigrade stress responses. Understanding the role of tardigrade stress-related genes, and the cellular processes they are involved in, can potentially be of great biomedical and biotechnological interest. Indeed, the potential translational application of tardigrade specific proteins was recently shown in studies revealing that the RNA-associating damage suppressor (Dsp) protein renders protection against UV-C and hydroxyl radicals to Dupont transformed human HEK293 cells (Hashimoto et al., 2016; Ricci et al., 2021). Moreover, tobacco plants that express the codon-optimized Dsp gene seem to have an enhanced tolerance towards UV and X-rays (Kirke et al., 2020), revealing a potential for tardigrade specific proteins in the engineering of crops that better tolerate the more stressful environmental conditions associated with global warming.

Data availability

The RNA-Seq data generated and used for differential expression analyses in this study are available at the European Nucleotide Archive under project PRJE49649.

Code availability

The bioinformatic pipeline used to analyse the RNA-Seq datasets is available in GitHub.

Author contributions

R.C.N. and N.M. conceived the study and designed the experiments. N.M. provided the tardigrade sample and R.C.N. performed the experiments and extracted total RNA. M.K., J.R.M., A.M. conducted the bioinformatics analyses. The study was supervised by N.M. and M.T.P.G. All authors contributed to the interpretation of data. A.M. and R.C.N. prepared figures and tables. R.C.N., A.M., N.M. wrote the manuscript with inputs from the other authors.

Declaration of Competing Interest

The authors report no conflict of interest.

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R.C. Neves et al.

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