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The Piezo-Hyperthermophilic Archaeon Thermococcus piezophilus Regulates Its Energy Efficiency System to Cope With Large Hydrostatic Pressure Variations

Yann Moalic1,2, Jordan Hartunians1,2, Cécile Dalmasso1,2, Damien Courtine1,2, Myriam Georges1,2, Philippe Oger3, Zongze Shao2,4, Mohamed Jebbar1,2 and Karine Alain1,2*

1 Univ Brest, CNRS, Ifremer, Laboratoire de Microbiologie des Environnements Extrêmes LM2E, UMR 6197, IUEM, Plouzané, France, 2 IRP 1211 MicrobSea, Sino-French Laboratory of Deep-Sea Microbiology, LM2E, Plouzané, France, 3 Université de Lyon, INSA Lyon, CNRS UMR 5240, Villeurbanne, France, 4 Key Laboratory of Marine Biogenetic Resources, The Third Institute of Oceanography SOA, Xiamen, China

INTRODUCTION

Deep-sea ecosystems share a common physical parameter, namely high hydrostatic pressure (HHP). Some of the microorganisms isolated at great depths have a high physiological plasticity to face pressure variations. The adaptive strategies by which deep-sea microorganisms cope with HHP variations remain to be elucidated, especially considering the extent of their biotopes on Earth. Herein, we investigated the gene expression patterns of Thermococcus piezophilus, a piezohyperthermophilic archaeon isolated from the deepest hydrothermal vent known to date, under sub-optimal, optimal and supra-optimal pressures (0.1, 50, and 90 MPa, respectively). At stressful pressures [sub-optimal (0.1 MPa) and supra-optimal (90 MPa) conditions], no classical stress response was observed. Instead, we observed an unexpected transcriptional modulation of more than a hundred gene clusters, under the putative control of the master transcriptional regulator SurR, some of which are described as being involved in energy metabolism. This suggests a fine-tuning effect of HHP on the SurR regulon. Pressure could act on gene regulation, in addition to modulating their expression.

Keywords: pressure, piezophile, thermococci, hydrothermal, transcriptomics
in gene expression, switch in metabolism to counteract the loss of biological activity, structural adaptations of macromolecules in order to face HHP and specific stress responses (Simonato et al., 2006; Oger and Jebbar, 2010; Picard and Daniel, 2013; Jebbar et al., 2015).

Although deep oceanic hydrothermal vents are characterized by high temperatures and hydrostatic pressures, very few (hyper)thermophilic piezophilic prokaryotes have been isolated from these ecosystems, and of the available isolates few have been tested for pressure tolerance, mostly due to the inherent technical constraints. Thus, the adaptation mechanisms of (hyper)thermophilic piezophiles remain unwell documented. Compared to psychrophiles, in which pressure and cold adaptation mechanisms may be overlapping, it is easier to independently sift through pressure adaptation mechanisms within (hyper)thermophilic models where high pressures and high temperatures have antagonistic effects (Siliakus et al., 2017). So far, only few (hyper)thermophilic piezophilic or piezosensitive microorganisms have been studied with respect to their response to HHP and most studies have focused on the order Thermococcales, a lineage of hyperthermophilic, heterotrophic, sulfur-reducing archaea, some of which are ubiquitous in the hot areas of deep-sea hydrothermal vents. Many studies focused particularly on the piezophiles Thermococcus barophilus and Pyrococcus yayanosii (Cario et al., 2015b; Vannier et al., 2015; Michoud and Jebbar, 2016). The transcriptomic response to supra-optimal pressures for growth of Thermococcus kodakarensis, a piezosensitive archaeon belonging to Thermococcales, resembles a classical stress response. It involves an up-regulation of genes involved in replication, repair and defense mechanisms, and a down-regulation of genes involved in energy production/conversion and in amino acid transport and metabolism. In contrast, the piezophilic taxon T. barophilus responds to changes in hydrostatic pressure by some variations in the metabolic pathways used and notably by variations of expression levels of genes coding for transporters, hydrogenases and oxidoreductases (Vannier et al., 2015). At stressful sub-optimal pressures, it also accumulates mannosylglycerate, a compatible solute, probably to help maintain cell volume and homeostasis (Cario et al., 2016). Adaptations at the membrane level (variations of the ratio archaeol/caldarchaeol and of lycopene unsaturations) were also observed in T. barophilus to maintain membrane fluidity as a result of pressure changes (Cario et al., 2015a). In addition, proteome adaptations (flexibility linked to reduced hydration water dynamics) of piezophilic species were also reported (Martinez et al., 2016). The obligate piezophile P. yayanosii responds to supra- and sub-optimal pressure changes by a major metabolic shift, namely a repression of the H2 metabolism and an increase in activity of sulfur-dependent hydrogenases, in addition to changes in the chemotaxis pathway, translation and CRISPR-cas gene expressions (Michoud and Jebbar, 2016). Overall, Thermococcales respond to pressure variations through structural flexibility and modulation of gene expression.

The hyperthermophilic piezophilic archaeon Thermococcus piezophilus strain CDGS\textsuperscript{T} was isolated from the world’s deepest known hydrothermal vent field, at nearly 5,000 m water depth, at the Mid-Cayman rise (Dalmasso et al., 2016b). It has an optimal pressure for growth of 50 MPa and holds the current record of pressure range for growth, growing effectively from atmospheric pressure to at least 120 MPa, and, though with difficulty up to 130 MPa (Dalmasso et al., 2016b). Due to its wide range of growth pressure, it appeared as a good model organism to perform transcriptome-level studies of adaptation to various pressure conditions, as for T. barophilus and P. yayanosii (Vannier et al., 2015; Michoud and Jebbar, 2016). To address the question of the effects of HHP variations on the transcriptome of T. piezophilus, RNA-seq analyses were performed on cells grown at sub-optimal (0.1 MPa), optimal (50 MPa), and supra-optimal (90 MPa) pressures for growth. The selection of these pressures was experimentally determined (growth yields identical to 0.1 and 90 MPa and sufficient biomass to perform transcriptomic analysis) (Dalmasso et al., 2016b). Here, by highlighting key gene clusters and the associated metabolic pathways engaged at optimal, sub-optimal and supra-optimal pressures for growth of this hyperthermophilic piezophilic archaean model, we provide new insights into the molecular responses of piezophiles to pressure variations.

**MATERIALS AND METHODS**

**Medium Culture and Growth Conditions**

Thermococcus piezophilus strain CDGS\textsuperscript{T} (UBOCC 3296\textsuperscript{T} = ATCC TSD-33\textsuperscript{T}) was isolated from a hydrothermal chimney sample from the Mid Cayman rise Beebe vent field, at 4,964 m water depth (Dalmasso et al., 2016b). The strain was grown at 75°C under anaerobic conditions, in modified Ravot medium prepared without maltose, with polysulfur (1% of polysulfide from a 0.05 M stock solution), as described elsewhere (Dalmasso et al., 2016b). Cells were grown until the mid-exponential growth phase (11 h at 0.1 MPa, 9 h at 50 MPa, and 12 h at 90 MPa) into 20 mL sterile plastic syringes incubated at 0.1, 50, and 90 MPa of hydrostatic pressure (Thermostated HP/HT incubators Top Industrie, France), from independent startup cultures grown at 50 MPa. The 3 biological replicates per pressure were incubated in the same stainless-steel pressure vessel (pressurized by pumping water into them), at the same time, together with negative controls. Cells were then harvested by centrifugation.

**Determination of Cell Numbers**

Cell counts were performed on a Thoma chamber (Preciss, France; surface area: 0.0025 mm\textsuperscript{2}, depth: 0.100 mm) and by phase contrast microscopy (Olympus BX60), to verify that the cell density was similar to that obtained at the end of the exponential growth phase during growth kinetics performed previously under exactly the same conditions.

**RNA Extraction and Purification**

As soon as the cell pellets (from 20 mL cultures) were obtained, TRIzol\textsuperscript{R} reagent (Ambion, Life Technologies) was added to them in order to extract total RNA. DNase I treatment (Promega\textsuperscript{R}) was then performed at 37°C for 30 min to remove genomic
DNA from the isolated total RNA. Total RNA was further purified onto RNA spin columns (RNasey Mini kit Qiagen®, RNA cleanup protocol), and stored at −80°C after verification of non-degradation with an Agilent 2100 Bioanalyzer (RNA 6000 Pico chip kit) and before sequencing. RNA concentrations were measured with a NanoDrop ND 1000® spectrophotometer (Thermo Fisher Scientific Inc.).

RNA-Seq
A Ribo-Zero rRNA Removal Bacteria kit (Epicenter, Madison, WI) used with manufacturer’s protocol was applied to deplete a maximum of rRNAs and enrich total RNA in mRNA. As it is often the case with Archaea, ribodepletion attempts were poorly efficient and the RNA-seq dataset still contained 73–94% ribosomal sequences. Ion Total RNA-Seq Kit v2 was used to make cDNA, to add barcodes, and to amplify the library. All cleanup steps were performed using Agencourt Ampure XP beads (Thermo Fisher Scientific). Each step was validated by a bioanalyzer quality control on Agilent RNA chips and/or with the “Qubit” fluorometer (Life Technologies) to determine the quality, size of fragments and quantity of produced material. RNA libraries were sequenced on P1v2 chips using the Ion S5™ System (Thermo Fisher Scientific). Sequencing was completed by the GeT-Genotoul platform in Toulouse (France). The RNA-seq data have been deposited into the NIH Sequence Read Archive (SRA) under the BioProject ID PRJNA739722 (accession numbers SRX11191114–11191122).

Bioinformatic Data Analysis
RNA-seq reads were mapped on all genes with Bowtie v2.3.1 (Langmead et al., 2009). Bowtie2 was run with default parameters and —no-unal. The result was converted from SAM to BAM format using Samtools v1.3.1 (Li et al., 2009). The reads counts were normalized to Transcripts Per Kilobase Million (TPM) (Supplementary Table 1). Then, TPM were mapped with Anvi’o (v7) (Eren et al., 2015) onto the T. piezophilus genome sequence deposited in the NCBI database under the accession number GCA_001647085.1 ASM164708v1. First, a contig database was generated (anvi-gen-contigs-database) with the genome sequence available on NCBI and a table containing all gene positions (—external-gene-calls and —ignore-internal-stop-codons arguments). Next, BAM files were processed individually with anvi-init-bam to get indexed alignment files. The latter were used to generate individual anvi’o profile to link the information available in the alignment file to the contigs databases (anvi-profile with required parameters and —min-contig-length 70). Last, all three profiles were merged with anvi-merge (default parameters). The command anvi-interactive was used to visualize the mapping results over the three conditions.

Differential Gene Expression Analysis
Data normalization and differential gene expression analysis were done with R/bioconductor packages edgeR (Robinson et al., 2010) and DESeq2 (Love et al., 2014). For each pressure condition, the data obtained from three independent biological replicate cDNA libraries were analyzed. Principal component analysis (PCA) was used to detect potential outlier on the samples. No outlier was detected. Regarding the filtration of transcripts, EdgeR’s RLE (“Relative Log Expression”) standardization was used, with filtering of weakly expressed genes, and then correction of tests with the BH method (Benjamini-Hochberg procedure for False Discovery Rate; alpha threshold = 0.05). The DEseq2 package was used to investigate the differential expression of genes between the different pressure conditions. Transcripts that were considered statistically significant were those with an adjusted p-value < 1%.

Gene Ontology Analysis
Functional annotation clustering of genes differentially expressed was carried out with the online software DAVID (Huang da et al., 2009),1 with default parameters.

Motifs Binding Sites Screening
The search for SurR and TrmB-like binding motifs within T. piezophilus genome was carried out using in-house Perl scripts. Gene promoters were defined as the 200 pb zones upstream of the start codon of the locus tag.

RESULTS AND DISCUSSION

General Response of Thermococcus piezophilus to Hydrostatic Pressure
Nine independent biological replicates, each inoculated with a different startup culture [generated at the optimal growth pressure (50 MPa)] were incubated in high-temperature stainless steel vessels at the three selected pressures (3 × 0.1 MPa; 3 × 50 MPa; 3 × 90 MPa). Their transcriptomes were generated after RNA extraction with TRizol®, then ribodepletion. After sequence filtration, there were 87,940,548 non-ribosomal usable total reads that were mapped as transcripts per kilobase million (TPM) on the 2,126 concatenated coding DNA sequence (CDS) of T. piezophilus (Figure 1A). After the differential expression validation step, it appeared that 1,373 of the 2,126 genes were regulated by pressure changes (Figure 1C but see Supplementary Table 2 for details) but, at most, the expression of 540 genes varied significantly when shifting from a tested pressure to another (Figure 1B and Supplementary Figure 1). At 0.1 MPa, 540 genes were overexpressed and 532 were underexpressed, while at 90 MPa, 220 genes were overexpressed and 224 genes were underexpressed compared to 50 MPa. There were more genes differentially regulated by pressure that were shared between high pressures (50 or 90 MPa) and atmospheric condition (0.1 MPa) (270 or 339 shared genes, respectively) than genes differentially regulated by pressure that were shared when comparing non-optimal pressures (0.1 or 90 MPa) to optimal growth condition (50 MPa) (109 or 125, respectively). The hierarchical clustering on z-scores highlighted a closer gene expression response between 0.1 and 90 MPa (Figure 1C).

It is therefore possible that the regulatory response to pressure variation compared to optimal growth condition (50 MPa) may
be broader and more specific at atmospheric pressure than at 90 MPa, even though the growth rate is almost identical at 0.1 and 90 MPa (Dalmasso et al., 2016b).

In order to gain a better understanding of the molecular tuning that takes place into the cell, we explored more precisely (i) the expression level of the genes regulated by two-well characterized transcriptional regulators (SurR and regulators of the TrmB family), (ii) the categories of genes regulated by pressure and (iii) the function of gene clusters whose expression is impacted by pressure, with emphasis on those involved in energy conversion and conservation.

**Is Pressure a Physical Parameter Able to Modify the Transcriptional Regulation Mechanism?**

The adaptive response to fluctuating environmental conditions has been less studied in *Archaea* than in other life domains.
(Karr, 2014). In *Thermococcales*, the regulators of the TrmB family and the SurR regulator are the two best-characterized transcriptional regulators. Regulators in the TrmB family control transcription according to sugar availability (Lee et al., 2003, 2005, 2008). These regulators bind to DNA in promoter regions to activate or repress the transcription of genes, thanks to TrmB-like binding motifs which have been published elsewhere (Gindner et al., 2014). SurR is the master regulator of primary electron flow pathways that responds directly to the presence or absence of sulfur in the environment (Lipscomb et al., 2009, 2017; Hidese et al., 2017). This ArsR-type master-regulator, which is unique to *Thermococcales* and conserved, is able to activate or repress the expression of the genes involved in the response to sulfur thanks to the presence of SurR binding motifs (GTTnAAC or GTTnGTT) in their promoters (Lipscomb et al., 2009). Its sequence-specific DNA binding activity is driven by a redox-active displacement of cysteine residues within a CxxC motif. In the presence of sulfur, a disulfide bond is formed and this oxidized form of SurR inhibits its binding activity. As a result, it loses its transcriptional modulation activities which are strongly correlated with the position of its binding sites, facilitating the recruitment of transcriptional machinery or blocking its progression (Yang et al., 2010).

It is interesting to note that while this study focused on the effect of variation in hydrostatic pressure (with no change in medium composition between experiments), we observed an increase in transcripts for the three TrmB-like genes (A7C91_RS03095, A7C91_RS05650, and A7C91_RS07665) and SurR gene (A7C91_RS07565) at 0.1 MPa vs. 50 and 90 MPa (Supplementary Table 2). In order to assess the potential overall effect of this regulation, the presence of SurR and TrmB-like binding motifs was examined in *T. piezophilus* gene promoters. TrmB-like binding motifs were detected in the promoter of only nine genes, of which only two were found to be regulated by pressure. This result seems to point to a limited role of TrmB and its targets in the response to pressure. However, due to evolutionary constraints, it is likely that the binding motifs used in our investigation in this archaeal species may be slightly different from those described in other species. In fact, these motifs were not detected in the promoter of the maltodextrin operon (MD) which is also overexpressed at 0.1 MPa vs. 50 and 90 MPa, whereas they were expected to be found there.

With regard to SurR, many genes under the control of this regulator have their expression modulated by pressure (Table 1; Figure 2, and Supplementary Figure 2; see next chapter). Thus, 170 promoters bearing at least one binding motif were detected among the 2,126 total CDS of *T. piezophilus*. In addition, among the 1373 CDS whose expression was regulated by pressure, 119 harbored a SurR binding motif. These values represent 8 and 8.7% of the total and overexpressed CDS, respectively, highlighting no significant increase in the expression of genes potentially under the control of SurR at different pressure conditions. However, given that 70% of genes potentially regulated by SurR are also pressure-regulated, this could be a clue of a high plasticity in adaptive response in *T. piezophilus*, but also within *Thermococcales* in general. Indeed, there is an effective variation in gene expression of SurR-regulated clusters. This could be the consequence of a pressure effect on the conformational state of SurR, modifying its binding capacity even in presence of sulfur, as has been shown for TrmB in *P. furiosus* (Krug et al., 2013). Moreover, a detailed analysis of the motif position in the promoters, revealed the presence of several potential binding sites, which, according to the BRE/TATA box, could provide some flexibility in terms of repression or activation, as

### Table 1: Variations in the expression of genes or gene clusters, known to be involved in hydrogen and sulfur metabolism.

| Product | Locus tag | Total CDS number | 0.1 > 50 MPa | 90 MPa | 0.1 > 90 MPa | 50 > 90 MPa |
|---------|-----------|------------------|--------------|--------|--------------|-------------|
| Mrp-Mbh | A7C91_RS04230-04165 | 14 | 8 | nd | nd | 13 | 14 |
| Fdh1-Mfh1-Mnh1 | A7C91_RS06945-07020 | 16 | nd | nd | 6 | 7 | nd |
| Fdh2-Mfh2-Mnh2 | A7C91_RS04320-04240 | 17 | 8 | 12 | nd | 7 | nd |
| F420-reducing hydrogenase | A7C91_RS_04340-04330 | 3 | 3 | 3 | nd | nd | nd |
| SHI | A7C91_RS08740-08725 | 4 | nd | nd | nd | 4 | 4 |
| SHIII | A7C91_RS06200-06185 | 4 | 4 | 3 | nd | nd | 4 |
| Mrp-Mbs | A7C91_RS08510-08450 | 13 | 12 | 13 | nd | 4 | nd |
| Nar | A7C91_RS06865 | 1 | nd | nd | nd | nd | nd |
| SurR | A7C91_RS07565 | 1 | 1 | 1 | nd | nd | nd |
| Pdo | A7C91_RS07570 | 1 | nd | nd | 1 | nd | nd |
| Nif II | A7C91_RS06205-6210 | 2 | nd | nd | 1 | nd | 2 |
| Nif II/Xfn | A7C91_RS02315-2310 | 2 | nd | nd | 2 | 1 | 1 |
| Nif III | A7C91_RS02425-2430 | 2 | nd | nd | 1 | nd | 2 |
| Fdh3 | A7C91_RS08745-08770 | 6 | nd | nd | 2 | 1 | nd |
| RNR | A7C91_RS02975 | 1 | nd | nd | 1 | nd | 1 |
| ATP synthase | A7C91_RS03355-03315 | 9 | nd | nd | 6 | nd | 7 |
| TnxR | A7C91_RS04130 | 1 | nd | nd | 1 | nd | 1 |
for the Mbs cluster (Table 2). In addition, some of the genes described as being regulated by SurR in other Thermococcales bear mutated motifs, such as for the ones encoding SHI (Table 2), but this will need to be further investigated by an EMSA assay to see if this affects the binding affinity of SurR. One other hypothesis would be the involvement of another transcriptional regulator. This hypothesis is supported by the presence of 37 other regulators among the genes regulated by pressure (Supplementary Table 2). Their respective occurrences in terms of over-expression according to pressure, i.e., 0.1, 50, and 90 MPa, are 23, 14, and 11, respectively (Supplementary Table 2). In the light of these results, the sub-optimal condition (0.1 MPa) seems to require a higher number of regulator overexpressions than the supra-optimal growth condition (90 MPa), respectively 21 vs. 4. This correlates with the greater number of total genes regulated in these conditions (540 and 220, respectively, Supplementary Figure 1). Overall, these data support the hypothesis that atmospheric pressure requires a stronger adaptive response than higher hydrostatic pressure conditions in T. piezophilus.

Categories of Genes Regulated by Pressure

To uncover the molecular mechanisms behind the pressure adaptive responses, the differentially expressed genes (DEG) were analyzed with the functional annotation tool DAVID which allows identifying functional categories and ranking them based on gene-enrichments in annotation terms (EASE score) (Huang da et al., 2009; Supplementary Table 3). Gene clusters characterized by a gene-enrichment (in this instance having an EASE score superior to 1) were analyzed in details. Results are presented in Supplementary Figure 1. The broadest functional response was predicted for 90 MPa vs. 0.1 MPa with five different classes of genes that were enriched (iron, hydrogen ion transport, ligase, ribosomal protein and nucleotidyltransferase) vs. one or two for the other conditions. These results show that the number of regulated genes can be twice as high in one comparison of conditions as in another without necessarily being more functionally diverse (540 vs. 220 overexpressed genes at 0.1 and 90 MPa, respectively against 50 MPa). As observed in other Thermococcus species, the adaptive response to pressure variations in Thermococcales appears to be more balanced, global and integrative than a classical response to a physical stress (Vannier et al., 2015; Michoud and Jebbar, 2016).

Functions of the Gene Clusters Regulated by Pressure

Main Functions Impacted by Pressure

The vast majority of gene expression variation was related to a global metabolic response, as described for T. barophilus and P. yayanosii (Vannier et al., 2015; Michoud and Jebbar, 2016). Variations in the level of transcription occurred globally at the level of gene clusters (Figure 1A). T. piezophilus appeared to adapt to pressure changes by modulating amino
acid and vitamin synthesis, production of potential compatibles solutes, and regulating the chemotaxis, S-layer and CRISPR-Cas systems. All these results are described in details in Supplementary Data 1. In summary, the strain does not possess the genetic potential to synthesize the three aromatic amino acids L-phenylalanine, L-tyrosine, and L-tryptophan, and underexpresses genes for the synthesis of a.a histidine while overexpressing genes encoding different families of transporters of amino acids, of oligopeptides and of dipeptides under non-optimal conditions. In the same fashion, few genes of amino acids L-phenylalanine, L-tyrosine, and L-tryptophan, and possess the genetic potential to synthesize the three aromatic.

TABLE 2 | SurR binding motifs positions upstream from the start codon of CDS involved in the electron flow.

| Product                  | Locus tag                      | First CDS of the product | Motif occurrence | Distance (size) of the binding motif from start codon of the first CDS |
|--------------------------|--------------------------------|--------------------------|------------------|---------------------------------------------------------------------|
| Mrp-Mbh                  | A7C91_RS04230-04165            | A7C91_RS04230            | 2                | 21 (short)                                                          |
| Fdh1-Mth1-Mnh1           | A7C91_RS06945-07020            | A7C91_RS06945            | 1                | 137 (short with mutation: gttacaat)                                 |
| Fdh2-Mth2-Mnh2           | A7C91_RS04320-04240            | A7C91_RS04320            | 1                | 123 (short)                                                        |
| Fd20-reducing hydrogenase | A7C91_RS_04340-04330           | A7C91_RS04340            | 1                | 71 (long)                                                          |
| SII                      | A7C91_RS08740-08725            | A7C91_RS08740            | 1                | 105 (long with mutation: gtttaaacctttggtt deletion)                |
| SHII                     | A7C91_RS06200-06185            | A7C91_RS06200            | 1                | 69 (long)                                                          |
| Mrp-Mbs                  | A7C91_RS08510-08450            | A7C91_RS08510            | 3                | 29 (short)                                                        |
| Nsr                      | A7C91_RS06865                  | A7C91_RS06865            | 1                | 17 (short)                                                        |
| SurR                     | A7C91_RS07565                  | A7C91_RS07565            | 2                | 49 (long)                                                         |
| Pdo                      | A7C91_RS07570                  | A7C91_RS07570            | 2                | 23 (short)                                                        |
| NfII/NfXII               | A7C91_RS02315-2310             | A7C91_RS02315            | 2                | 44 (short)                                                        |
| Fdh3                     | A7C91_RS08745-08770            | A7C91_RS08745            | 1                | 71 (long)                                                          |
| RNR                      | A7C91_RS02975                  | A7C91_RS02975            | 0                | /                                                                 |
| ATP synthase             | A7C91_RS03355-03315            | A7C91_RS03355            | 0                | /                                                                 |
| TnxR                     | A7C91_RS04130                  | A7C91_RS04130            | 0                | /                                                                 |

Effect of Pressure on Hydrogenases and Energy Conversion Circuits

Effect of Hydrostatic Pressure on Membrane-Bound Hydrogenases MBH and MBS

Members of the Thermococcales order grow heterotrophically by fermentation of peptides or sugars used as carbon and energy sources. This leads to the production of CO₂, acetate, and gas, whose composition, respectively H₂S or H₂, depends in particular on the presence or absence of elemental sulfur in the medium (Verhees et al., 2003; Sakuraba et al., 2004). The metabolism generates reduced ferredoxins that are used by membrane-bound oxidoreductases to create an ion gradient with the extracellular environment (Kengen et al., 1996; Verhees et al., 2003; Sakuraba et al., 2004). This ion gradient can further be used by an ATP synthase to produce ATP (Figure 3; Pisa et al., 2007). In Thermococcales in general and T. piezophilus in particular, two main systems of membrane bound oxidoreductases exist (Schut et al., 2013). The first one is the simple respiratory system MBH (membrane-bound hydrogenase) that is generally activated in the absence of S⁰ (Sapra et al., 2003) and links the oxidation of reduced ferredoxin generated during glycolysis to the formation of H₂ (Kanai et al., 2011; Schut et al., 2012). When sulfur is provided in the medium, a second type of membrane oxidoreductase, is active: the MBS (Membrane-bound sulfane reductase) system (Wu et al., 2018), highly homologous to MBH (13 vs. 14-genes cluster). Recent studies have shown the generation of H₂S from spontaneous breakdown of tri- and
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FIGURE 3 | Schematic representation of the main transcriptional metabolic regulations observed at sub-optimal (0.1 MPa) and supra-optimal (90 MPa) pressures for growth. This figure is centered on the general metabolism and focuses notably on hydrogenases and energy conversion processes and on their transcriptional regulator SurR, in the context of adaptive response to hydrostatic pressure. Blue and red arrows represent the overexpression (up direction) and subexpression (down direction) of genes at sub-optimal (0.1 MPa), and supra-optimal (90 MPa) pressures for growth, respectively, compared to the optimal pressure for growth (50 MPa).

Our transcriptomic study has shown that the level of expression of both MBH and MBS gene clusters is modulated by pressure (Table 1). This result is surprising, as *T. piezophilus* has been grown with polysulfur in all these experiments. Indeed, numerous studies on *Thermococcales* carried out at atmospheric pressure have shown that the genes coding for the MBS complex have their level of transcription increased when sulfur is present in the culture medium. On the contrary, it has been shown that the level of expression of the genes coding for the MBH complex decreases sharply in the presence of sulfur (still at atmospheric pressure) (Chou et al., 2007; Jager et al., 2014). Here, our results show that the expression level of the *Mbh* gene cluster varies despite the presence of sulfur. At the sub-optimal pressure of 0.1 MPa, the gene clusters of the MBS and MBH complexes are both overexpressed (Table 1 and Figures 2, 3), compared to 50 MPa. The genes of the *Mbh* cluster were considerably overexpressed at the supra-optimal pressure of 90 MPa compared to 50 and 0.1 MPa. It was also slightly more transcribed at 0.1 MPa compared to 50 MPa (Table 1 and Figures 2, 3). This suggests a pressure-effect overtaking a sulfur-effect on the regulation of gene expression.

**Effect of Hydrostatic Pressure on Cytosolic Hydrogenases, Oxidoreductases, and Glycolytic Enzymes**

Other enzymes involved in these $H_2$/sulfur metabolic pathways (Figure 3) include the cytosolic hydrogenases and oxidoreductases SHII/II, NfnI/II/III, Pdo, and TrxR, which have been shown to be under the regulation of SurR (Lipscomb et al., 2017). SHI and SHII are soluble hydrogenases, that could couple the synthesis of protons from an $H_2$ uptake to the regeneration of NADPH from NADP$^+$ (Schut et al., 2013). The ferredoxin:NADP$^+$ oxidoreductases (Nfn) of *Thermococcales* are involved in the electron transfer from ferredoxin to NADP$^+$, under high concentrations of $H_2$ (Verhaart et al., 2010). The subsequent NADPH can be then reoxidized by a glutamate dehydrogenase and an alanine amino-transferase in the alanine biosynthetic pathway. Pdo is another glutaredoxin-like protein disulfide oxidoreductase whose roles are not totally understood. It is notably regulated by SurR and steps in the overall cellular redox balance and in the hydrogen metabolism. It could receive electrons from NADPH through a thioredoxin reductase (TrxR), and then serve as an electron carrier itself for ribonucleotide reductase (RNR), allowing links to deoxyribonucleotide synthesis (Schut et al., 2013; Demmer et al., 2015).

Here, an important overexpression of the gene cluster encoding the soluble hydrogenase SHII was observed at 0.1 and 90 MPa (Figure 2) compared to 50 MPa. This overexpression was more pronounced at 0.1 MPa (Table 1). Genes encoding the
cytoplasmic SHI hydrogenase subunits were overexpressed but only at 90 MPa compared to 50 and 0.1 MPa (Figure 2). These soluble hydrogenases could thus be mobilized under stressful pressure conditions to regenerate NADPH. An overexpression of the NADH-dependent ferrodoxin:NADP + oxidoreductase (NfnI) was observed at 50 and 90 MPa, but not at atmospheric pressure. As with “Thermococcus onnurineus,” T. piezophilus has 2 genes coding for homologs of the S and L subunits of NfnI (A7C91_RS06205-6210) of P. furiosus (Nguyen et al., 2017), and also possesses NfnII (A7C91_RS02315-2310) and NfnIII (A7C91_RS02425-2430). NfnI was overexpressed at 90 MPa vs. 50 MPa, while NfnII was overexpressed at 50 MPa (vs. 0.1 MPa). NfnIII was overexpressed at 90 MPa vs. 0.1 MPa. In addition, genes encoding a glutaredoxin-like protein (A7C91_RS07570) and the disulfide oxidoreductase Pdo (A7C91_RS07570) were overexpressed at 50 MPa. Genes coding for the thioredoxin reductase TrxR (A5C91_RS04130) and for the ribonucleotide reductase RNR (AC91_RS02975) were overexpressed at 50 and 90 MPa vs. 0.1 MPa. Altogether, these observations tend to indicate that under stressful conditions, H2 production (with Mbh) is activated. In addition, at 0.1 MPa, the Mbs, and Mbh gene clusters are overexpressed, suggesting a cumulative effect to cope with pressure. Soundly, several genes coding for glycolytic enzymes, a pathway which regenerates reduced ferrodoxin, were overexpressed at the reference pressure of 50 MPa [i.e., ADP-dependent glucokinase (A7C91_RS07115), phosphoglycerate kinase (A7C91_RS05540) and glyceraldehyde-3-phosphate dehydrogenase (A7C91_RS09215)], and also at 90 MPa [glyceraldehyde-3-phosphate dehydrogenase, glyceraldehyde-3-phosphate ferrodoxin oxidoreductase (A7C91_RS04780) (GAPOR)].

Effect of Hydrostatic Pressure on Other Mrp-Mbh Oxidoreductases

In T. piezophilus, other types of Mrp-Mbh oxidoreductases clusters were also regulated by pressure. These clusters code for formate dehydrogenases (FDH) which are found only in some of the Thermococcales, such as P. yayanosii and “T. onnurineus” (Schut et al., 2013). Thus, three formate dehydrogenase gene clusters were found in T. piezophilus genome, similar to those of “T. onnurineus,” namely the membrane-bound hydrogenase clusters Fdh1-Mfh1-Mnh1 (A7C91_RS06945-7020) and Fdh2-Mfh2-Mnh2 (A7C91_RS04320-04240), and the cytoplasmic hydrogenase cluster Fdh3 (A7C91_RS08745-08770). The synteny of these clusters is conserved between both organisms. The presence of the Fdh2-Mfh2-Mnh2 gene cluster together with the one encoding a formate transporter suggests that T. piezophilus could metabolize formate. This was not observed experimentally at atmospheric pressure (Dalmasso et al., 2016a), likely because growth on formate was tested at 75°C, under conditions where the reaction was probably not exergonic (Kim et al., 2010).

At 50 MPa, genes of the clusters Fdh1-Mfh1-Mnh1 and Fdh3 were overexpressed, as well as the associated ATP synthase clusters. Genes of the cluster Fdh2-Mfh2-Mnh2 were also more transcribed at 50 MPa and even more again at 0.1 MPa. This might indicate that growth on formate could be stimulated under optimal and sub-optimal pressure conditions.

A cytoplasmic homolog of the F420-reducing hydrogenase (A7C91_RS04340-04330) present in methanogenic archaea (frhAGB-encoding hydrogenase) whose expression is linked to various functions in Thermococcus sp., including carbon monoxide metabolism, chemotactic signal transduction and Fdh3 regulation (Jeon et al., 2015; Lee et al., 2017), was also overexpressed at 0.1 MPa compared to 50 and to 90 MPa. This hydrogenase is involved in various cellular processes in “T. onnurineus,” which could also be the case in T. piezophilus. This observation is also consistent with the fact that the cell undergoes a broad metabolic change to cope with pressure variations.

Effect of Pressure on SurR-Regulated Genes: A Synthesis

In summary, with the exception of genes of the Mrp-Mbs cluster, genes encoding enzymes which are known to be under SurR negative control, namely Pdo and Nfn, were all overexpressed at the reference pressure of 50 MPa vs. 0.1 MPa (in presence of sulfur). Interestingly, the genes and gene clusters that were under SurR positive control were overexpressed at supra- and sometimes sub-optimal pressures for growth (see above). This suggests that, in T. piezophilus, the expression patterns of genes known to be under the control of the SurR regulon appear to follow a classical response/behavior to sulfur only under optimal pressure conditions. Sub-optimal pressures and, to a lesser extent, supra-optimal pressures are likely to influence the expression of these genes in a new way of regulation (Figure 3). As an illustration, a cooperation between the membrane complexes MBH and MBS appears to occur at atmospheric pressure regardless of the presence of sulfur.

CONCLUSION

This study presents the transcriptional adaptive strategy implemented by T. piezophilus, the piezophilic microorganism with the widest range of tolerance to HHP (0.1–130 MPa) known to date. The overall differential gene expression revealed by RNA-seq confirms the results previously obtained with T. barophilus and P. yayanosii (Vannier et al., 2015; Michoud and Jebbar, 2016) that there is an overall metabolic change rather than a classical stress response. A thorough knowledge of the process of energy-conservation and its regulation in other Thermococcales models, has enabled us to highlight the involvement of these processes in the adaptive response, at least in the regulation of the expression of genes coding for the proteins involved in this process. These results are changing our knowledge of the mechanism of regulation of this energy-conservation process under conditions of non-optimal pressures and of the role of SurR under these conditions.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.
**AUTHOR CONTRIBUTIONS**

KA and CD designed the experiment. CD and MG carried out the experiments. YM and DC performed the bioinformatics analyses. KA, YM, and JH interpreted the data and wrote the manuscript. PO, MJ, and ZS helped supervise the project. All authors reviewed the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.730231/full#supplementary-material

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