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Responses of *Methanosarcina barkeri* to acetate stress

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Abstract

Background: Anaerobic digestion of easily degradable biowaste can lead to the accumulation of volatile fatty acids, which will cause environmental stress to the sensitive methanogens consequently. The metabolic characteristics of methanogens under acetate stress can affect the overall performance of mixed consortia. Nevertheless, there exist huge gaps in understanding the responses of the dominant methanogens to the stress, e.g., Methanosarcinaceae. Such methanogens are resistant to environmental deterioration and able to utilize multiple carbon sources. In this study, transcriptomic and proteomic analyses were conducted to explore the responses of *Methanosarcina barkeri* strain MS at different acetate concentrations of 10, 25, and 50 mM.

Results: The trend of OD600 and the regulation of the specific genes in 50 mM acetate, indicated that high concentration of acetate promoted the acclimation of *M. barkeri* to acetate stress. Acetate stress hindered the regulation of quorum sensing and thereby eliminated the advantages of cell aggregation, which was beneficial to resist stress. Under acetate stress, *M. barkeri* allocated more resources to enhance the uptake of iron to maintain the integrities of electron-transport chains and other essential biological processes. Comparing with the initial stages of different acetate concentrations, most of the genes participating in acetoclastic methanogenesis did not show significantly different expressions except *hdrB1C1*, an electron-bifurcating heterodisulfide reductase participating in energy conversion and improving thermodynamic efficiency. Meanwhile, *vnfDGHK* and *nifDHK* participating in nitrogen fixation pathway were upregulated.

Conclusion: In this work, transcriptomic and proteomic analyses are combined to reveal the responses of *M. barkeri* to acetate stress in terms of central metabolic pathways, which provides basic clues for exploring the responses of other specific methanogens under high organics load. Moreover, the results can also be used to gain insights into the complex interactions and geochemical cycles among natural or engineered populations. Furthermore, these findings also provide the potential for designing effective and robust anaerobic digesters with high organic loads.

Keywords: Methanosarcina barkeri MS, Transcriptomic, Proteomic, Acetate stress, Anaerobic digestion

Background

Anaerobic digestion (AD) has been highlighted as an important organic waste treatment technology for bioenergy production. Acetate is a main intermediate during AD processes and can be accumulated to high concentrations, due to the rapid fermentation of easily biodegradable waste [1]. The high acetate concentration is identified as a significant cause of inhibition in AD processes [2–4]. And it leads AD to a dilemma called "inhibited pseudo-steady state" along with ammonium and pH, where AD processes maintain stable performances under neutral conditions but with lower methane yields [5, 6]. Nevertheless, few studies provide direct insights into the responses of specific methanogens to acetate stress, which frequently occurs in engineered AD processes.

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He et al. Biotechnol Biofuels (2019) 12:289 Page 2 of 14

Methanosarcina can be dominant during AD processes [7–9] and are identified as the key functional microbes to alleviate acid inhibition [10]. They prefer a higher acetate concentration [11], maintain a stable acetate utilization rate [12, 13] and even survive at a pH value of approximately 4.5 [14]. Moreover, *Methanosarcina* can form complex multicellular structures to resist stress [15].

In addition to the dominant niche in AD processes, Methanosarcina has more advantages over other methanogens. It can yield methane through all four known methanogenesis pathways [16], and shift pathway according to the levels of stress [17–19]. Meanwhile, it has two disparate energy-conserving systems [20] and can alter electron transport chains [21, 22]. Additionally, M. barkeri has a much higher growth yield on H2 and CO2 than the methanogens without cytochromes [23]. Previous experiments indicated that Methanosarcinaceae in mixed consortia shifted methanogenesis pathways gradually. It was shifted from acetoclastic methanogenesis to hydrogenotrophic methanogenesis, when acetate concentrations increased from 50 to 150 and 250 mM [5]. All of these features determine the widespread distribution of Methanosarcina in AD processes and its high tolerance to acetate stress. Nevertheless, to the best of our knowledge, the responses of Methanosarcina to acetate stress are still unexplored.

Currently, omics have been widely implemented in the studies of AD processes to search for novel microorganisms [24], elucidate key metabolic activities [25] and other objectives [26]. Indeed, extensive studies on AD processes have made use of cultivation-independent biotechnologies to document the variations of microbial populations under acetate stress. But it is still difficult to gain the insights into the responses of specific methanogens, due to environmental complexities and microbial interactions in mixed consortia. In contrast, transcriptomic or proteomic analysis on pure cultivation can exclude interferences from other microbes, which therefore has been applied to explore the stress responses of specific microbes [27-29]. Nevertheless, the effective analytical method of combining pure cultivation with transcriptomic and proteomic analyses, has not yet to be used to explore the responses of specific methanogens to acetate stress.

Therefore, this analytical method was used in the present study to explore the responses of *M. barkeri* strain MS, a model methanogen, to different levels of organic load stresses formed by different total acetate concentrations. Regulations of the genes participating in stress response, signal transduction, element translocation, acetoclastic methanogenesis and nitrogen fixation, show the responses of *M. barkeri* to acetate stress, which provides bases to understand the responses of other methanogens.

These findings are conductive to explore the complex interactions in mixed consortia during AD processes and develop strategies to alleviate inhibitory factors and optimize AD processes.

Results

The biochemical data and micrographs

The sampling points are marked in Fig. 1a. The 10-, 25-, and 50-group are labeled as "10", "25", and "50", respectively. "I" and "T" represent initial and terminal sampling points, respectively. The methanogenesis started to proceed rapidly from the initial sampling points. The terminal sampling points can be used to verify under the condition that the total acetate concentration is approximately equal, whether the gene regulations of *M. barkeri* will be different after long-term inhibition. As shown in Fig. 1a, the total acetate concentrations of 25-T and 10-I, 50-T and 25-I were approximately equal, respectively.

The previous work suggested that free acetic acid functioned not only as the substrate for acetoclastic methanogenesis but also as an inhibitor [12], while present work was mainly to focus on the inhibitory effects of different acetate concentrations on M. barkeri. Both of the total acetate and free acetic acid concentrations showed significant differences among the initial sampling points (P < 0.05) (Additional file 1: Table S1), which indicated that the M. barkeri in three groups were undergoing different levels of acetate stress.

The trend of each biochemical data was fitted using the Boltzmann function (Additional file 2: Fig. S1 and Additional file 3: Table S2). The rates at each sampling point were calculated by the fitted curves (Table 1). During a lag phase of 10 days (Fig. 1a), the increase in acetate concentrations might be due to the gradual release of the acetate absorbed by cell aggregations. After the lag phase, acetate in three groups began to be degraded and was close to the utilization threshold, which is consistent with previous studies [30]. Methane was yielded simultaneously in the three groups. The accumulative yields of methane were roughly close to the theoretical yield of acetoclastic methanogenesis, with a maximum deviation of 12.9% (Fig. 1b). Along the degradation of acetate, the pH value in each group constantly increased from moderately acidic to neutral (Fig. 1c). The OD600 values decreased apparently and showed coherent descending trends in three groups during the first 30 days. But the descending trend in the 50-group was more gradual than that in the 10- and 25-group after the turning point (Fig. 1d and Table 1). Even though M. barkeri was no longer in exponential phase, the stress from stationary phase could not make OD600 values decrease so apparently, and the descending trend would not slow down or even cease. These characteristics indicated that acetate

He et al. Biotechnol Biofuels (2019) 12:289 Page 3 of 14

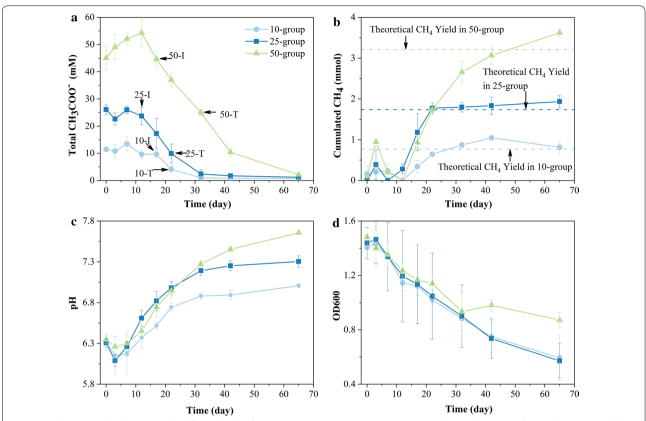


Fig. 1 The biochemical indictors under different levels of acetate stress. **a** CH_3COO^- concentration and sampling points. **b** Cumulated CH_4 yield. **c** pH values in culture media. **d** OD600. Error bars indicate \pm sd of biological triplicates; invisible error bars indicate the sd values are very small

Table 1 The biochemical data at the initial and terminal sampling points of three groups

Group	Sample point	OD600	OD600 descending rate (day ⁻¹)	Total acetate (mM)	Acetate consumption rate (mM/day)	CH ₄ production rate (mmol/day)	pН	pH rising rate (day ⁻¹)
10-group	10-l	1.122	0.018	9.66	0.73	0.049	6.47	0.046
	10-T	1.022	0.017	4.15	0.94	0.073	6.69	0.037
25-group	25-I	1.194	0.020	23.70	0.59	0.075	6.56	0.051
	25-T	1.048	0.018	9.92	1.57	0.034	6.93	0.036
50-group	50-l	1.168	0.015	44.59	0.74	0.095	6.69	0.040
	50-T	0.934	0.0079	24.93	1.78	0.092	7.22	0.030

stress was absolutely the major stress and the M. barkeri in 50-T might have acclimated to the acetate stress. The latter was further proved by the higher minimal acetate consumption rate and minimal $\mathrm{CH_4}$ production rate in the 50-group than those in the 10- and 25-group (Additional file 4: Table S3).

As acetate concentration increased, cell aggregations disintegrated, with larger distances between cells, and irregular cell features instead of plump sphericity appeared (Additional file 5: Fig. S2a-c). It was difficult to

find denser cell aggregations in 50-I, but cell aggregations of approximately 50 μ m in length were observed in 10-I and 25-I (Additional file 5: Fig. S2m, n). It indicated that the high level of acetate stress interfered with the formation and stability of cell aggregations. The decrease in fluorescence signal of coenzyme F₄₂₀ indicated that the activities of *M. barkeri* were weakened with the increase in acetate stress (Additional file 5: Fig. S2d–f). At the terminal sampling points, not only the size of cells themselves, but also the size of cell aggregations decreased,

He et al. Biotechnol Biofuels (2019) 12:289 Page 4 of 14

and many separate cells appeared. Cell features were generally irregular at the terminal sampling points (Additional file 5: Fig. S2g–i). The fluorescent signals, which were not observed in 10-T and 25-T, but were observed in 50-T. It indicated the active biological processes in *M. barkeri* (Additional file 5: Fig. S2j–l) and confirmed the OD600 trend in the 50-group (Fig. 1d).

Transcriptomic analysis

The transcriptomic protocol yielded an average of 18,130,832 filtered non-rRNA reads with an average length of 145 bp per library (Additional file 6: Table S4a). Approximately 85.26–99.46% of the 3494 annotated open reading frames were recovered by transcriptomic analysis. As the reactions progressed, the proportions of high-abundance transcripts decreased with the increasing proportions of low-abundance transcripts in each group, indicating that the number of highly expressed genes decreased (Fig. 2).

The detailed information of all differently expressed genes (DEGs) is listed in Additional file 7: Dataset S1. There were more DEGs in "10-T_10-I", "25-T_25-I", and "50-T_50-I" than in "25-I_10-I", "50-I_10-I", and "50-I_25-I" (Additional file 8: Fig. S3), which illustrated that the regulation mechanisms during the consumption of acetate were more complex than these between the initial sampling points. The ratio of the identical DEGs among "10-T_10-I", "25-T_25-I", and "50-T_50-I" was up to 30%, and most of them were identified as hypothetical proteins, indicating that hypothetical proteins might be essential to resist acetate stress. The results of RT-qPCR

indicated that the RNA-Seq data were reliable (Additional file 6: Table S4b).

The KEGG module enrichment analysis showed that nitrogen fixation pathway was upregulated significantly in "50-I_10-I" and "50-I_25-I". With the consumption of acetate, nitrogen fixation pathway was downregulated in "25-T_25-I" and "50-T_50-I" (Fig. 3). Nevertheless, there was no significant up- or downregulation in acetoclastic methanogenesis pathway, which was the main methane metabolism using acetate as substrate.

For the comparisons between the initial sampling points, the GO enrichment analysis also showed that the terms related to nitrogen fixation and nitrogen utilization were upregulated in "50-I_10-I" and "50-I_25-I" (Fig. 4), which were consistent with the results from above KEGG module enrichment analysis. The activities related to quorum sensing (QS) in "50-I_25-I", such as response to stimulus and cell communication, were downregulated (Additional file 9: Fig. S4). With the consumption of acetate, the nitrogenase activities in "25-T_25-I" and "50-T 50-I" were downregulated (Fig. 5).

WGCNA identified 13 modules of genes and the minimum module size was 32 transcripts. These modules were distinguished by colors (Additional file 10: Dataset S2). 100 and 115 genes (labeled as "N_Ac" and "N_pH" groups, respectively) were negatively correlated with acetate consumption rates and pH values, respectively, of which 56 genes were identical (Additional file 10: Dataset S2). These genes might play essential roles at the initial sampling points where inhibitory factors were the strongest. Gene MSBRM_0367 including both the "N_Ac"

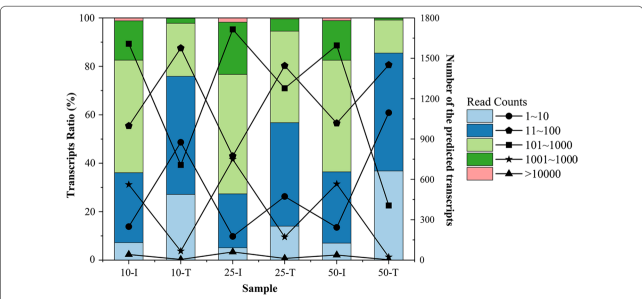


Fig. 2 Ratio and number of predicted transcripts with different read counts in each sample. The bar chart represents the ratio of the transcripts with different read counts in the specified sample, and the line chart represents the number of these transcripts

He et al. Biotechnol Biofuels (2019) 12:289 Page 5 of 14

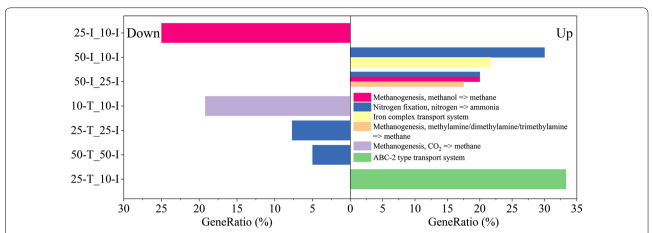


Fig. 3 KEGG module enrichment analyses. The downregulated modules are shown on the left, and the upregulated modules are shown on the right. The filter criterion for KEGG module enrichment was P < 0.05 (Benjamini and Hochberg methods). Gene ratio is calculated by dividing k by n, where k is the number of DEGs assigned to the specific KEGG module and n is the number of DEGs annotated by KO identifiers

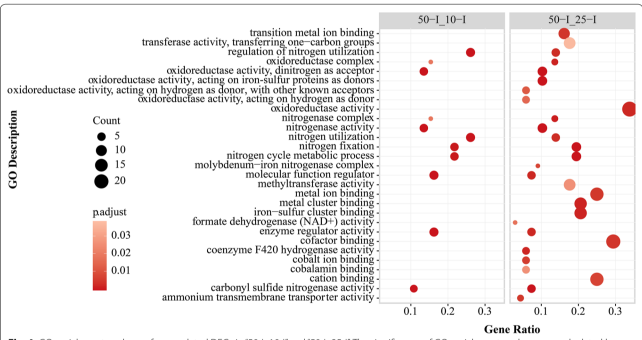


Fig. 4 GO enrichment analyses of upregulated DEGs in "50-I_10-I" and "50-I_25-I". The significance of GO enrichment analyses was calculated by Benjamini and Hochberg method

and "N_pH" groups encodes transcriptional regulators, associated with the genes participating in nitrogen metabolism, QS, methane metabolism and other processes (Additional file 11: Fig. S5a). In addition to gene MSBRM_0367, gene MSBRM_0203 and MSBRM_2051 were also included in the "N_pH" group, and these two genes had relationships with the genes participating in energy synthesis, transcriptional regulation, sensory transduction, iron uptake and other processes, indicating

that these processes were essential for resisting stress at the initial sampling points (Additional file 11: Fig. S5c, d). The detailed results of WGCNA are described in Additional file 12: Text S1.

Proteomic analysis

Proteomic analysis recovered 1813 of 3494 predicted proteins. In "50-I_10-I", there were 29 and 54 proteins showing upregulation and downregulation, respectively

He et al. Biotechnol Biofuels (2019) 12:289 Page 6 of 14

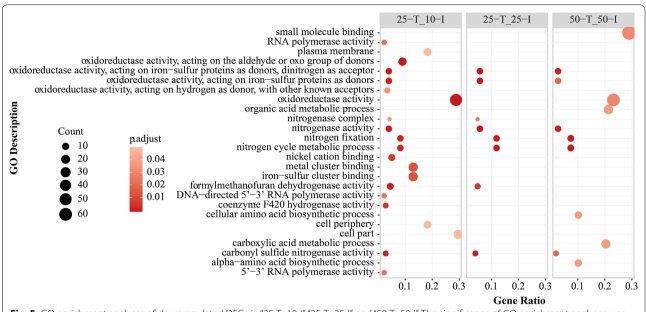


Fig. 5 GO enrichment analyses of downregulated DEGs in "25-T_10-I", "25-T_25-I", and "50-T_50-I". The significance of GO enrichment analyses was calculated by Benjamini and Hochberg method

(Additional file 13: Dataset S3). In "50-T_50-I", 144 and 68 proteins showed upregulation and downregulation, respectively (Additional file 13: Dataset S3). The fold changes in mRNAs and their corresponding proteins in "50-I_10-I" or "50-T_50-I" did not show coherence (Fig. 6), indicating that transcription and translation were

uncoupled for most proteins. In "50-T_50-I", the proteins with the largest fold changes were hypothetical proteins, and approximately 33% of the differently expressed proteins (DEPs) were hypothetical proteins. The high proportion of hypothetical proteins revealed huge gaps in the knowledge of the responses to acetate stress. In

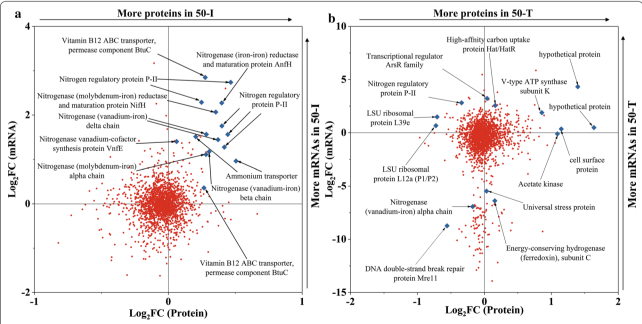


Fig. 6 The fold changes in mRNAs and their corresponding proteins. a "50-I_10-I; b "50-T_50-I". Red dots or blue diamond points represent genes detected in both transcriptomic and proteomic

He et al. Biotechnol Biofuels (2019) 12:289 Page 7 of 14

"50-I_10-I", the mRNAs and their corresponding proteins participating in nitrogen fixation, were all upregulated, which highlighted the importance of bioavailable nitrogen sources for resisting acetate stress (Fig. 6a).

In "50-I_10-I" and "50-T_50-I", most of the DEPs that were identified as hypothetical proteins were cytoplasmic proteins, indicating they rarely participated in transmembrane processes like substance transport. These proteins might primarily participate in cytoplasmic activities to maintain the internal stabilities of cells. For the hypothetical proteins that were identified as signal peptides, most of them could not be located (Additional file 13: Dataset S3), showing complex signaling networks under acetate stress. Even if the above results showed the relationships of these hypothetical proteins to acetate stress, it was problematic to speculate their detailed functions.

Discussion

Stress proteins under different levels of acetate stress

The genes encoding heat shock proteins (Hsp 60) and DNA double-strand break (Dsb) repair proteins were upregulated in "50-I_25-I" and "50-I_10-I", respectively. These proteins participated in mitochondrial protein import or macromolecular assembly and restrained cell death or comprehensive genetic variation in cells. The ATP-dependent DNA ligase gene (*lig1*) was also upregulated in "50-I_10-I" and "50-I_25-I", which has an active role in DNA replication, recombination and repair [31]. This indicated that the DNA structure was unstable at the initial sampling point under high level of acetate stress, although *M. barkeri* was capable of surviving in 50 mM acetate [12, 32].

Along with the consumption of acetate, most of the DEGs encoding stress proteins and Lig1 were upregulated in "10-T_10-I", indicating that M. barkeri still did not acclimatize to the acetate stress. Nevertheless, in "50-T_50-I", most of the genes encoding stress proteins were downregulated according to the transcriptomic analysis, and proteomic analysis showed that all detected Hsp 60 and Dsb proteins were downregulated. In "50-T_25-I", the *dsb* was downregulated. Furthermore, most of the DEGs encoding Hsp, Dsb and universal stress proteins were upregulated in "25-T_10-I". Considering the OD600 trend in each group (Fig. 1d) and the regulation trends of DEGs encoding stress proteins, it was revealed that a period of exposure to a high level of acetate stress might promote M. barkeri to acclimatize to acetate stress. This presumption was consistent with the phenomenon in our previous study where the abundance of Methanosarcinaceae started to be detected or increase in the later stages of the reactions [5].

Signal transduction in cell aggregations

The genes encoding cell division proteins (FtsEX complex) which were essential for assembling or stabilizing the septal ring [33], were downregulated in "10-T_10-I" and "25-T_25-I" (6.41-fold and 13.27-fold, respectively). Nevertheless, these genes showed a slightly decreased fold change in "50-T_50-I" (2.06-fold). These regulation trends roughly coincided with the OD600 trends (Fig. 1d). *ftsH* participates not only in cell division and cell control, but also in membrane functions and gene expression, which was upregulated in "50-T_25-I" and downregulated in "25-T_10-I". Logically speaking, a low level of stress had fewer negative impacts on cells, but the present work indicated that a high level of acetate stress could stimulate the greater "potential" for *M. barkeri* to resist stress.

Cell proliferation could facilitate the formation of cell aggregations. The micrographs and the previous study found that *M. barkeri* resisted stress by forming multicellular aggregations (Additional file 5: Fig. S2) [34]. In the cell aggregations, the gradients of acetate concentrations were formed to alleviate the acetate stress on the inner cells and prevent more H₂ from diffusing [35]. H₂ cycling was essential for acetoclastic methanogenesis [36].

Cell aggregations exchanged information of cell densities by quorum sensing (QS) system using peptide signals, such as oligopeptides and dipeptides. In "25-I_10-I", "50-I 10-I", and "50-I 25-I", the genes encoding periplasmic oligopeptide-binding protein (OppA) or oligopeptide/dipeptide transport system permease proteins (OppC and DppB) were upregulated. Nevertheless, the genes encoding ATP-binding proteins (OppD and DppF) were downregulated in "25-I_10-I" and "50-I_10-I". ATP hydrolysis was essential for the normal functions of ABC transport system [37], therefore, the downregulation of oppD and dppF indicated that acetate stress could block the information exchange on a community-wide scale. The blocked information exchange offset the advantages of cell aggregations, and M. barkeri might not make proper responses to acetate stress. Because the activation of the Type II secretion system proteins was under the control of QS system [38], the downregulation of the DEGs encoding Type II secretion system proteins also indicated the blocked information exchange. In other comparisons, *opp/dppBCDF* were all downregulated, but opp/dppA did not show up- or downregulation. It indicated that *opp/dppA* was more resistant to acetate stress, which was helpful to maintain the normal functions of the QS system in cell aggregations.

Approximately half of the DEGs encoding LSU and SSU ribosomal proteins were upregulated in "50-T_50-I" and

He et al. Biotechnol Biofuels (2019) 12:289 Page 8 of 14

"50-T_25-I" (41.4% and 55.6%, respectively), but 88.9% of the DEGs encoding the same proteins were downregulated in "25-T_10-I". The expression of ribosomal proteins usually represented protein synthesis, which was an energy-consuming process accounting for 75% of energy expenditure in cells [39]. It was revealed that *M. barkeri* gradually recovered energy synthesis to support the synthesis of some proteins in the 50-group, which helped *M. barkeri* acclimatize to acetate stress.

Membrane and ABC transporters for translocating elements

Transmembrane transfer activities were related to membrane and ABC transporters. Except for the comparisons between the initial sampling points, the fluctuant regulation trends of the DEGs encoding cell surface proteins in other comparisons indicated that the responses of cell membranes were complicated. And in "10-T_10-I", "25-T_25-I", "50-T_50-I", and "50-T_25-I", the genes encoding mannose-6-phosphate isomerase were upregulated, which was related to polysaccharide biosynthesis and the modification of cell wall carbohydrates. It indicated that *M. barkeri* modified its cell surface structures according to the levels of acetate stress, which was also observed in the previous study [40].

The phosphate transport system (Pst) was a more efficient system for Pi translocation in *M. barkeri*. In "10-T_10-I" and "25-T_25-I", *pstB* was downregulated which provided energy to free Pi [41]. It indicated that acetate stress could result in an insufficient energy supply for translocating elements and thereby cause a series of negative effects on physiological functions. But *pstB* did not show significant regulation in "50-T_50-I" and "50-T_25-I", therefore the stable expression of *pstB* might promote the acclimation of *M. barkeri* to the acetate stress in 50-group.

Iron is essential for maintaining stable methanogenesis [42, 43], where iron was a main component in redox enzymes, such as ferredoxin and Hdr. The previous studies demonstrated that iron deficiency hindered nucleotide synthesis and ATP production [44, 45]. For more iron uptake, the genes encoding ferrous iron transport protein B, which were essential for iron supply in anaerobic conditions, were upregulated in "50-I_10-I" and "50-I_25-I". And the proteomic analysis also showed the upregulation of iron complex transport system permease proteins in "50-I_10-I" (Additional file 13: Dataset S3). Furthermore, the iron complex transport activity was enriched in "50-I_10-I" (Fig. 3). It was suggested that the reason why exogenous iron could maintain stable methanogenesis was not only it could enhance the interspecies electron transfer between species and reduce oxidative-reductive potential [46, 47], but also it could maintain the integrities of some key enzymatic activities. Compared with the gene regulation related to other elements' uptake, the upregulation of iron uptake indicated the inhibited *M. barkeri* would allocate more resources to enhance the uptake of more essential elements, especially under high level of acetate stress.

The inappropriate transmembrane ion gradients formed under acetate stress could cause the imbalance of osmotic pressures. The glycine betaine transport system (Pro) could provide osmo-protection [48]. In "25-I_10-I" and "50-I_10-I", proV encoding ATP-binding protein was downregulated. The insufficient energy supply would hinder the normal functions of Pro under acetate stress, which thereby induced cell death. The upregulation of proV in "50-T_25-I" also might promote the acclimatization of M. barkeri to the acetate stress in 50-group. As expected, proV was downregulated in "25-T_10-I", where the M. barkeri did not acclimatize to the acetate stress.

Acetoclastic methanogenesis for energy synthesis

As the terminal of electron transfer, methane was yielded through acetoclastic methanogenesis with energy synthesis. In "50-I_25-I", the genes encoding heterodisulfide reductase subunits (HdrB1C1) were upregulated, but the hdrDE did not show significant regulations. HdrA1B1C1 had been speculated to be an electron-bifurcating enzyme in energy conversion as shown in Fig. 7. One possible explanation for why multitype Hdr expression was that HdrABC could help cells to acclimatize to the fluctuations in substrate concentrations and conserve energy more efficiently [23, 49]. The available free energy from acetoclastic methanogenesis under standard condition ($\Delta G^{o'} = -36 \text{ kJ/CH}_{A}$) provided only a small amount of energy for growth, because of the endergonic reaction where acetate was activated to acetyl-CoA. The upregulation of hdrB1C1 could be a complementary mechanism for *M. barkeri* to improve the thermodynamic efficiency under the high level of acetate stress.

Methanosarcina barkeri was one of the few methanogens that could participate in direct interspecies electron transfer [50], and it could shift methanogenesis pathways according to stress levels. Furthermore, Methanosarcina could produce H₂ and act as a syntrophic acetate oxidizer during growth on acetate [32]. On the other hand, HdrA1B1C1 and homologous HdrA2B2C2, which had similar functions [49, 51], were suggested to be essential for direct interspecies electron transfer [52]. These characteristics indicated there might be a preliminary hypothesis that in the cell aggregations formed under high level of acetate stress, some M. barkeri might act as syntrophic acetate oxidizers, and another M. barkeri might perform hydrogenotrophic methanogenesis to yield more energy

He et al. Biotechnol Biofuels (2019) 12:289 Page 9 of 14

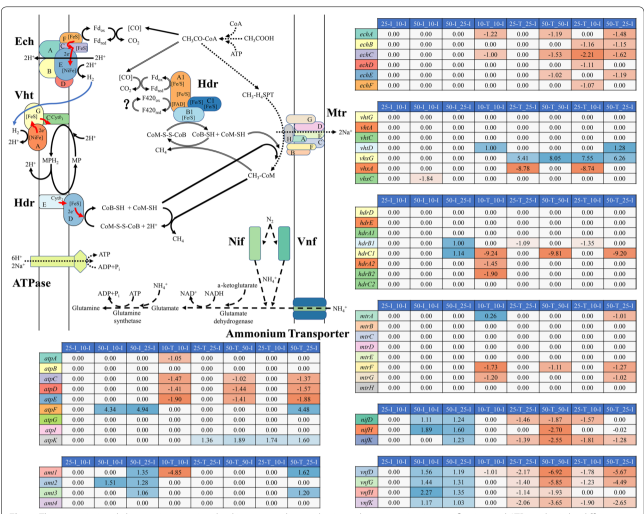


Fig. 7 The transcriptional changes in genes involved in proposed acetoclastic methanogenesis, nitrogen fixation and ATP synthesis by different comparisons. The values in the heatmaps show the average Log_2 fold changes in transcripts translating the same subunit of each enzyme in different comparisons. Red represents downregulation, blue represents upregulation, and white represents no significant regulation. The solid lines indicate the conventional acetoclastic methanogenesis. The double bond lines indicate the postulated acetoclastic methanogenesis. The dotted lines indicate the overlap of conventional acetoclastic methanogenesis and postulated acetoclastic methanogenesis. The dashed lines indicate the nitrogen fixation pathway. Red solid lines indicate electron flow in each enzyme and blue solid lines indicate H_2 flow in acetoclastic methanogenesis. The question mark represents the enzyme re-oxidizing $F_{420}H_2$

for cellular activities. Therefore, electron transfer would exist in one species with distinct pathways. Moreover, the DEGs encoding coenzyme F_{420} hydrogenase subunit alpha, beta, and gamma (FrhABG), which participated in hydrogenotrophic methanogenesis, were upregulated in "50-I_25-I". The upregulation of frhABG was part of the evidence for the electron transfer between $M.\ barkeri$ under high level of acetate stress. And the hypothetical proteins occupying most of the proteome might also participate in this electron transfer. Nevertheless, this hypothesis needs to be further verified.

The other key genes participating in acetoclastic methanogenesis did not show significant regulation at

the initial sampling points (Fig. 7). It indicated that *M. barkeri* metabolized acetate stably under different levels of acetate stress within its tolerance range. The stable acetate metabolism could help *M. barkeri* to outcompete other methanogens for acetate and thereby dominating mixed consortia. The gene encoding V-type ATP synthase subunit F (AtpF), which was likely a regulatory subunit for controlling pH homeostasis, was upregulated in "50-I_10-I" and "50-I_25-I" (Fig. 7). Therefore, maintaining pH homeostasis was a mechanism of *M. barkeri* to hold the intracellular environment suitable under acetate stress.

He et al. Biotechnol Biofuels (2019) 12:289 Page 10 of 14

"10-T 10-I", with the reactions proceeding, hdrA2B2C1 were downregulated (Fig. 7). And the genes encoding energy-conserving hydrogenase subunits A and C (EchA and EchC, respectively), which are essential for forming transmembrane proton gradients and keeping H₂ cycling during acetoclastic methanogenesis, were also downregulated (Fig. 7). Furthermore, the gene encoding tetrahydromethanopterin S-methyltransferase subunit A (MtrA) was downregulated, which could form a transmembrane sodium gradient for ATP production [16, 21]. Meanwhile, atpAECD were downregulated. These results indicated that an insufficient acetate concentration had negative impacts on the energy synthesis of M. barkeri, whose affinity for acetate was low. It might be one of the reasons why M. barkeri were eliminated in low acetate concentrations. Although mtrFG were also downregulated, they appeared to be the nonessential components in mtr operon in terms of transferring methyl [53]. The same regulation trends of mtrAFG indicated that there might have potential interactions among them.

In "25-T_25-I", methanophenazine-reducing hydrogenase (VhxAG) with unknown functions, showed opposite regulation trends (up- and downregulation, respectively). Furthermore, vhx in "10-T_10-I", "25-T_25-I", and "50-T_50-I" showed significant regulation. It was suggested that the deletion of *vhxD* in *vhx* operon caused Vhx to have no analogous functions of another methanophenazine-reducing hydrogenase (Vht) [20, 54], which was essential for Hdr to conserve energy in acetoclastic methanogenesis [36]. Nevertheless, vhx operon could encode the main functional subunits homologous to Vht, and VhtD might process Vhx under the controls of the DEGs encoding mobile element proteins or transposons. Combining the gene regulations with the operon structure, it indicated that Vhx might participate in acetoclastic methanogenesis as a hydrogenase. In "50-T_50-I", the downregulation of ech and atp might be caused by the less acetate concentration at "50-T" point where M. barkeri had acclimatized to the acetate stress as discussed above.

Methanogenic N_2 fixation for bioavailable nitrogen resources

Methanogenic N_2 fixation is proposed as an essential way to provide bioavailable nitrogen resources [55], and the methanogens being capable of fixing N_2 such as M. barkeri, Methanobacterium bryantii and so on, distribute in diverse habitats [56–58]. The DEGs encoding nitrogenase (molybdenum–iron) subunits (NifHDK), and alternative nitrogenase (vanadium–iron) subunits (VnfHDK), which has lower catalytic efficiency in Methanosarcina species [59], were upregulated in

"50-I_10-I" and "50-I_25-I" (Fig. 7). It indicated that ammonia was indispensable for *M. barkeri* to resist acetate stress, which could also be confirmed by the upregulation of the DEGs encoding ammonium transporters and nitrogen regulatory protein p II. In "50-I_10-I", the regulation trends of the nitrogen regulatory proteins, nitrogenases and ammonium transporters, showed positive correlation between the transcriptomic and proteomic analyses (Fig. 6a). Ammonia can be assimilated into cellular nitrogen by glutamine synthetase (Gs) and glutamate dehydrogenase (Gdh) (Fig. 7). Gs was identified as a DEP in "50-I_10-I", and *gs* was upregulated in "50-I_25-I".

Ammonia is a biologically useful reduced form incorporated into amino acids and other vital compounds. Therefore, the enhanced nitrogen fixation pathway could help M. barkeri maintain the normal functions of key biological processes in the initial stages of acetate stress. And it could also help the more sensitive M. barkeri to outcompete other methanogens under acetate stress, as observed in the previous studies [5, 34]. Although M. barkeri could grow under nitrogen fixation conditions with acetate, the growth rate was considerably slower than with ammonium as nitrogen source [60]. Therefore, ammonium was used as the nitrogen source in the present work and its concentration was about 9.35 mM as the recommended cultivation condition, which could support the normal growth of M. barkeri. The enhanced nitrogen fixation pathway indicated that more bioavailable nitrogen sources were essential to resist acetate stress. However, as another major problem of anaerobic digestion, ammonia stress is caused by the high concentration of ammonia. Therefore, it is necessary to find a "best of both worlds" ammonia concentration, which still needs further explorations.

In addition to maintaining the normal functions of key biological processes, the enriched nitrogen fixation pathway might also keep the proton balance in cells. Under the same pH values, acetate stress produces relatively more free acetic acid, which can diffuse into cells. The intracellular is a near-neutral environment, so free acetic acid will be dissociated and then increase the intracellular proton concentration. The excessive intracellular protons could uncouple proton force on cytomembrane and hinder energy synthesis [61, 62]. Nevertheless, the ammonia synthesized by the enriched nitrogen fixation pathway could bind the excessive protons in cytoplasm, and thereby alleviated the imbalance of protons, which was toxic to *M. barkeri*.

Except for "25-I_10-I", "50-I_10-I", and "50-I_25-I", the DEGs encoding nitrogenase subunits were all

He et al. Biotechnol Biofuels (2019) 12:289 Page 11 of 14

downregulated in other comparisons. For "10-T_10-I", "25-T_25-I", and "25-T_10-I", the downregulation might be caused by the insufficient energy synthesis as discussed above, because N_2 fixation required abundant reducing power and ATP [55, 63]. For "50-T_50-I" and "50-T_25-I", $M.\ barkeri$ had acclimatized to the acetate stress at the "50-T" point, indicating more nitrogen sources were not needed.

The differences between proteomic and transcriptomic analyses

The regulations of some mRNAs and their corresponding proteins showed poor correlation, which was well known and observed in previous studies [64, 65]. Proteins could remain stable even beyond the entire cell cycle relative to mRNAs, whose half-lives averaged only a few min in E. coli [66]. The different temporal scales of transcriptome and proteome could be used to explain the poor correlation between some mRNAs and proteins. In addition to the differences in temporal scales, posttranscriptional or translational regulations were another explanation. The low proportion of coding regions in the genome of M. barkeri (79.2%) indicated that a large number of antisense RNAs and sRNAs existed, which participated in posttranscriptional regulation. The modulation effects of sRNAs played a prominent role in archaea by influencing the evolution and stability of mRNAs and translation efficiency [65, 67]. Translation initiation also affected mRNA-protein correlation [67], therefore the DEGs encoding translation initiation factors in "50-T 50-I" could explain the poor mRNA-protein correlation in part.

Conclusions

The present work describes the gene regulations of *M. barkeri* in terms of the main metabolism pathways under acetate stress using transcriptomic and proteomic analyses. The findings provide bases to understand responses of other methanogens, and are conductive to explore the complex interactions in mixed consortia. As the sole conductor of methanogenesis, methanogens have multifaceted impacts on mixed consortia.

The 50 mM acetate had the most severe inhibition on *M. barkeri* between the initial sampling points, while the high level of acetate stress (50 mM) promoted the acclimation of *M. barkeri* to stress. The information exchange by QS system was hindered by insufficient ATP hydrolysis, which offset the advantages of cell aggregations. Enriched iron uptake suggested that *M. barkeri* could allocate more resources to enhance the uptake of key elements relative to other elements. Therefore, the addition

of external iron might be a useful approach to alleviate acetate stress in engineered anaerobic digestors.

As the sole pathway to metabolize acetate, the aceto-clastic methanogenesis pathway did not show significant regulation of conventional enzymes between the initial sampling points. The high tolerance of acetoclastic methanogenesis to acetate stress made Methanosarcinaceae the dominant methanogens in mixed consortia, where the other microbes in the outer layer of granule sludge could further provide protection. The expressions of hdrABC were complementary mechanisms to keep $\rm H_2$ cycling and improve thermodynamic efficiency under acetate stress. It was widely considered that Vhx could not function as homologous Vht due to the lack of maturation protease, while the significant regulation of vhx revealed that Vhx might be a complement to Vht, which needs to be further explored.

The enriched in methanogenic N_2 fixation pathway revealed that sufficient bioavailable nitrogen sources were essential for sensitive M. barkeri to resist acetate stress. However, the mechanisms where N_2 fixation alleviated acetate stress needed to be further explored.

The genes encoding the different subunits of one complex did not show the same regulation trends. On the other hand, the genes encoding the same subunit had distinct regulation trends. Furthermore, a considerable proportion of the DEGs were annotated as hypothetic proteins, and many novel transcripts were assembled by the software. This indicated that comprehensive gene regulations existed in *M. barkeri* under acetate stress, and some genes were activated, which were not expressed under suitable conditions. Future studies can focus on the functions of the DEGs encoding hypothetic proteins, which may prevent the disorder of physiological activities and adjust pathways dynamically. Meanwhile, the complex interactions in mixed consortia under acetate stress are unclear and need to be further explored.

Methods

All materials and methods are described in detail in Additional file 12: Text S1.

Pure cultivation and stress conditions

Methanosarcina barkeri MS (DSM 800) was incubated in 250-mL sealed serum bottles with a 160-mL medium volume at 35 °C. These serum bottles were divided into three groups, named as 10-group, 25-group, and 50-group, in which the total acetate concentrations were set as 10, 25, and 50 mM, respectively (Additional file 14: Fig. S6). Although the highest acetate concentration used in

He et al. Biotechnol Biofuels (2019) 12:289 Page 12 of 14

the previous study was 250 mM [12], it far exceeded the actual acetate concentration in engineered AD processes (Additional file 15: Table S5), and also far exceeded the acetate concentration in some studies on pure methanogens [32, 68, 69]. Therefore, the highest acetate concentration in the present work was set as 50 mM, which was used in the previous study [32]. The present work mainly focused on the responses of *M. barkeri* to different total acetate concentrations, therefore the initial pH values of three groups were controlled at about 6. Acetoclastic methanogenesis could not proceed well below this pH value.

Transcriptomic analysis

The raw reads were filtered and trimmed using fastp [70]. Trimmed reads were aligned to the reference genome from Ensembl Genomes database using HISAT2 [71]. Stringtie [72] was applied to quantify the abundances of mRNAs. DEGs were estimated by edgeR [73]. The selection criteria for DEGs was a fold change>2 and false discovery rate<0.05. The DEGs were verified by RT-qPCR analyses. The GO and KEGG module enrichment analyses of DEGs were conducted using clusterProfiler [74]. The weighted correlation network analysis was conducted using WGCNA [75]. The raw data have been deposited to NCBI in SRA (PRJNA528099).

The comparisons are labeled in the format of "sampling point 1_sampling point 2", where sampling point 1 represents the treatment sample and sampling point 2 represents the control sample. The comparisons included "25-I_10-I", "50-I_10-I", "50-I_25-I", "10-T_10-I", "25-T_25-I", "50-T_50-I", "25-T_10-I", and "50-T_25-I".

Proteomic analysis

The cell pellets collected from 10-I, 50-I, and 50-T were used for proteomic analysis. The digested peptides were labeled using iTRAQ reagents (AB Sciex). The labeled peptides were analyzed on an EASY-nLC 1200 system coupled to a Q-Exactive mass spectrometer (Thermo-Fisher). All raw data were collected using Thermo Xcalibur and analyzed using Sequest HT (Thermo-Fisher). Proteomic data have been deposited to the ProteomeX-change Consortium via the iProx partner repository with the dataset identifier PXD013207. The prediction of transmembrane helices were conducted by TMHMM [76]. PSORTb [77] was used to predict the proteins' subcellular localization, and SignalP [78] was used to predict the presence of signal peptides.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s13068-019-1630-5.

Additional file 1: Table S1. The concentration of total acetate and free acetic acid in each group.

Additional file 2: Fig. S1. Fitted curves of biochemical data. (a) Total CH₃COO⁻ concentration; (b) Cumulated CH₄ yield; (c) pH values in culture media; (d) OD600 values. The curves were fitted using the Boltzmann function on the average values of the data at each sampling point.

Additional file 3: Table S2. The parameter values of each fitted curve.

Additional file 4: Table S3. Maximum and minimum rates of 10-, 25- and 50-group.

Additional file 5: Fig. S2. The brightfield and epifluorescence micrographs of *M. barkeri* MS from different sampling points. (a), (d), and (m) 10-l; (b), (e), and (n) 25-l; (c) and (f) 50-l; (g) and (j) 10-T; (h) and (k) 25-T; (i) and (l) 50-T.

Additional file 6: Table S4. The transcriptomic sequencing data analysis. (a) The counts of filtered non-rRNA reads mapped to the reference genome. (b) Fold changes of differentially expressed genes from RNA-seq and RT-qPCR in different comparisons.

Additional file 7: Dataset S1. The detailed information of all DEGs in each comparison.

Additional file 8: Fig. S3. The regulation of genes in each comparison. (a) "25-I_10-I"; (b) "50-I_10-I"; (c) "50-I_25-I"; (d) "10-T_10-I"; (e) "25-T_25-I"; (f) "50-T_50-I"; (g) "25-T_10-I"; (h) "50-T_25-I". The number of DEGs doesn't include the novel transcripts assembled by Stringtie.

Additional file 9: Fig. S4. GO enrichment analysis of downregulated DEGs in "25-l_10-l" and "50-l_25-l".

Additional file 10: Dataset S2. The detailed information of WGCNA analyses.

Additional file 11: Fig. S5. The transcriptional networks of MSBRM_0367, MSBRM_0968, MSBRM_0203, and MSBRM_2051. (a) The transcriptional network of MSBRM_0367. (b) The transcriptional network of MSBRM_0968. (c) The transcriptional network of MSBRM_0203. (d) The transcriptional network of MSBRM_2051. Nodes in the network represent genes, and gray lines link genes with the top 50 pairwise TOM values; thicker lines indicate higher TOM values. The background colors of nodes indicate which module they belong to (the detailed information of nodes is listed in the Additional file 10: Dataset S2).

Additional file 12: Text S1. The supplementary notes about materials, methods and WGCNA analyses.

Additional file 13: Dataset S3. The detailed information of proteomic analysis.

Additional file 14: Fig. S6. Flow chart of setting up different levels of acetate stress.

Additional file 15: Table S5. Acetate concentration in full-scale anaerobic digestors.

Abbreviations

AD: anaerobic digestion; AtpF: V-type ATP synthase subunit F; DEGs: different expressed genes; DEPs: different expressed proteins; Dsb: DNA double-strand break; DppB: dipeptide transport system permease protein; DppF: dipeptide transport system ATP-binding protein; Ech: energy-conserving hydrogenase; Fts: cell division proteins; FrhA: coenzyme F420 hydrogenase subunit alpha; FrhB: coenzyme F420 hydrogenase subunit beta; FrhG: coenzyme F420 hydrogenase subunit gamma; Gs: glutamine synthetase;

He et al. Biotechnol Biofuels (2019) 12:289 Page 13 of 14

Gdh: glutamate dehydrogenase; Hdr: heterodisulfide reductase; Hsp: heat shock protein; lig1: ATP-dependent DNA ligase; Mtr: tetrahydromethanopterin S-methyltransferase; Nif: nitrogenase molybdenum-iron protein; OppA: periplasmic oligopeptide-binding protein; OppC: oligopeptide transport system permease protein; OppD: oligopeptide transport system ATP-binding protein; Pst: phosphate transport system; Pro: glycine betaine transport system; QS: quorum sensing system; Vhx: methanophenazine-reducing hydrogenase; Vht: methanophenazine-reducing hydrogenase vanadium-iron protein.

Authors' contributions

PH raised the concept and drafted the manuscript. FL developed the general research question, discussed the results and revised the manuscript. HD carried out all of the experiments, analyzed the data and drafted the manuscript. WH contributed to data analysis. YL contributed to experiment design. LS contributed to manuscript revision. All authors read and approved the final manuscript.

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Availability of data and materials

The sequence datasets generated during the current work are available in the NCBI Sequence Read Archive and iProX, accession numbers provided in the text. The datasets supporting the conclusions of this article are uploaded as additional materials.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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He et al. Biotechnol Biofuels (2019) 12:289 Page 14 of 14

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