Thermophilic microbial cellulose decomposition and methanogenesis pathways recharacterized by metatranscriptomic and metagenomic analysis

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The metatranscriptomic recharacterization in the present study captured microbial enzymes at the unprecedented scale of 40,000 active genes belonged to 2,269 KEGG functions were identified. The novel information obtained herein revealed interesting patterns and provides an initial transcriptional insight into the thermophilic cellulose methanization process. Synergistic beta-sugar consumption by Thermotogales is crucial for cellulose hydrolysis in the thermophilic cellulose-degrading consortium because the primary cellulose degraders Clostridiales showed metabolic incompetence in subsequent beta-sugar pathways. Additionally, comparable transcription of putative Sus-like polysaccharide utilization loci (PULs) was observed in an unclassified order of Bacteroidetes suggesting the importance of PULs mechanism for polysaccharides breakdown in thermophilic systems. Despite the abundance of acetate as a fermentation product, the acetate-utilizing Methanosarcinales were less prevalent by 60% than the hydrogenotrophic Methanobacteriales. Whereas the aceticlastic methanogenesis pathway was markedly more active in terms of transcriptional activities in key genes, indicating that the less dominant Methanosarcinales are more active than their hydrogenotrophic counterparts in methane metabolism. These findings suggest that the minority of aceticlastic methanogens are not necessarily associated with repressed metabolism, in a pattern that was commonly observed in the cellulose-based methanization consortium, and thus challenge the causal likelihood proposed by previous studies.

Cellulose is Earth’s most abundant biomass, and it is gaining worldwide attention as a renewable resource for bioenergy production1. In both natural and engineered systems, the bioconversion of lignocellulosic biomass benefits from synergistic reactions among microorganisms within a microbiome2. Our understanding of community dynamics and the ecological roles of microorganisms living in various cellulose-degrading communities has been revolutionized by the application of whole genome shotgun sequencing based on Next Generation Sequencing (NGS), which is also referred to as metagenomic sequencing. The NGS-based metagenomes of cellulose deconstruction microorganisms have revealed an unexpectedly high diversity of genes related to polysaccharide hydrolysis3,4 and were later shown to be an extensive resource for the discovery of novel glycoside hydrolases5. However, these excellent frontier explorations of genes embedded in genomes, which outline the broad genomic potential of a cellulolytic system, lack the ability to identify the pathways that are actually expressed or the key genes that are differentially transcribed.

Metaproteomics and metatranscriptomics (also known as RNA-seq) are both capable of providing useful insight into this expression issue. Nevertheless, the relatively low protein separation throughput of metaproteomics ranges from hundreds to thousands of identified proteins6–8 and, in some cases, fails to favor sufficient gene/protein coverage for reliable data interpretation. Some contradictory results were reported in these metaproteomic studies; for instance, Hanreich, A. et al. identified an unusually high expression of enzymes related to the methanogenesis process9, and other studies suggested that genes involved in carbohydrate metabolism are much more active10.
Relative to proteomic-based approaches, NGS-based metatranscriptomic sequencing has the advantage of providing massive gene identification, and the expression of these genes enables a comprehensive observation of functional microbes and genes involved in microbial processes. Thus far, this method has been applied most extensively in human-related medical disciplines to analyze the transcription profiles of genes related to certain symptoms and, to a lesser extent, to study gene expression in environmental samples, especially polysaccharide-active microbiota. The aim of the present study was to gain insight into the transcriptional activities of the key genes and microbial populations involved in thermophilic cellulose deconstruction and to evaluate metatranscriptomics as a potentially suitable technology for this purpose. To this end, total RNA and DNA extracts were subjected to metatranscriptomic and metagenomic profiling, respectively, with an Illumina platform. The technical reproducibility of the nucleotide isolation method (DNA or RNA), Illumina library preparation and subsequent sequencing were quantified.

This first metatranscriptomic attempt to disclose the expression activity of genes that are involved in thermophilic cellulose decomposition may provide novel insights into the following subjects: (1) the differences between transcriptional activities and the genetic potential of carbohydrate-active genes (CAGs) in thermophilic cellulose deconstruction and (2) the active roles of various microbial populations in thermophilic cellulose deconstruction and methanogenesis. Additionally, the analytic procedure established here will serve as a reference for fundamental issues in the transcriptional quantification of gene activity in a metatranscriptome with unevenly distributed microbial populations.

Results
Reproducibility of RNA and DNA libraries. The analysis workflow combining replicated metatranscriptomic and metagenomic datasets is illustrated in Figure 1. First, the technical reproducibility of metagenomic and metatranscriptomic sample preparation (including RNA/DNA extraction, library construction and Illumina sequencing) was investigated using DNA and RNA library replicates, respectively. The reliable reproducibility of the metagenome preparation was confirmed by the strong consistency ($R^2 > 0.9$, Table S1) of both the taxa composition and functional annotation between metagenomic replicates. However, the mRNA proportion of metatranscriptomics showed noticeable variations (with an $R^2$ value of approximately 0.7,

![Figure 1 | Schematic of the experimental and analytic workflow. An electrophoresis gel of two replicates of extracted DNA (left) and total RNA (right) are shown.](image-url)
Table S1) when only two RNA samples from a single environmental sample were analyzed.

Community structure. Approximately 0.06% of total DNA reads were designated as rDNA sequences in the DNA dataset (Table S2). Of the total rDNA reads, 87% could be aligned to the reference 16S sequences in the Greengenes database12. Based on clustering analysis, the thermophilic cellulose-degrading consortium is dominated by four bacterial orders, *Anaerolineales* (14.6%), *Clostridiales* (13.9%), *Bacteroidales* (12.4%) and *Thermotogales* (9.7%), in addition to the two archaeal populations, *Methanobacteriales* (7.8%) and *Methanosarcinales* (5.9%) (Figure 2).

Metagenomic gene recovery and annotation. The rarefaction curve indicated that the sequencing depth was sufficient for de novo assembly (Figure S1). Ninety percent of the 50 million total DNA sequences were included in the de novo assembly by IDBA_UD. Assembled scaffolds longer than 300 bp exhibited an N50 of 10,963 bp (see Table S3 for detailed information of the metagenomic assembly). From the assembled scaffolds, 155,454 ORFs were predicted. Among these ORFs, 26.3% showed transcriptional activity (assembly). From the assembled scaffolds, 155,454 ORFs were predicted. Among these ORFs, 26.3% showed transcriptional activity (assembly).

**Transcriptional activities of major populations in the community.** A comparison of transcriptional activities across different populations was initially based on the ratio of mRNA reads to coding DNA reads as proposed by Yu, K. et al.13 As shown in Figure S2, a significant annotation bias towards *Proteobacteria* was observed when using this approach in which the short reads were used directly as search queries. This bias mirrored the uneven representation of different taxa in the NCBI nr database, the effect of which was significantly enlarged by the low classification ratio of short reads (less than 40%, Table S4). Given the inherent difficulty of finding homologies for short NGS reads, the reliability of results built on this quantification approach is questionable14.

To circumvent this limitation, the transcriptional activities of protein-coding genes were investigated in an alternative manner based on the MRPKM value (the ratio of RPKM-RNA to RPKM-DNA as defined in Equations S1, S2, and S3) of the ORFs, which tended to have a much higher classification ratios (63.4% in present study, Table S4). As shown in Figure S3, species in *Synergistetes* and *Nitrospirae* showed the highest average MRPKM values of 1.5 and 1.2, respectively, compared to the other microbial groups in the community. Genes encoding enzymes related to cell mobility via bacterial flagella showed extraordinarily high expression (with MRPKM values larger than 100) in both of these phyla, suggesting their superior motility within the community.

Global function of the community. Several interesting phenomena were observed in the transcriptional pattern of SEED subsystems. First, the two subsystems Motility and Chemotaxis and Stress Response showed substantial transcription (Figure 3, right). Two microbial groups played major roles in these two subsystems, *Thermotogae* in Motility and Chemotaxis and *Bacteroidetes* in Stress Response (Figure 3, right). *Thermotogae* is also actively involved in cellulose hydrolysis through the expression of a wide range of beta-glycosidases (Figure 4). Extraordinary transcriptional activities (with MRPKM > 100) were observed for genes (alkyl hydroperoxide reductase [AhpC] and ferric ion binding domain) involved in cell protection against reactive oxygen species in *Bacteroidetes*.

Expression of CAGs. Among the expressed CAGs, metabolic acclimation towards cellulose substrate deconstruction was evident when enzyme families related to cellulose decomposition showed 1.5 times higher transcriptional activity than those related to other steps in carbohydrate metabolism. Furthermore, enzymes catalyzing cellulose hydrolysis (the blue-labeled cellulase families in Figure 4, right) are highly active despite their comparably lower prevalence in the metagenome (Figure 4 left). In contrast to the simplicity of cellulosomes, the thermophilic populations maintain the substantial genetic redundancy of oligosaccharide-degrading enzymes from 8 families (Figure 4 left). The excessive genetic diversity of oligosaccharide-degrading genes is regarded as crucial for cellulosytic consortia to maintain metabolic flexibility15.

As expected, *Clostridiales* play an essential role in cellulose chain breakdown by expressing a large proportion of the active exocellulases (GH05) and all of the endocellulases (GH48 and GH09). The most active endocellulases in *Clostridiales* (GH09 and GH48) are adjacent to genes encoding the heat shock proteins of the sigma factor and HSP60, respectively (Figure S4). Additionally, the overrepresentation of cohesin and dockerin, proteins that are commonly involved in the formation of lignocellulosic multi-enzyme complexes (cellulosomes)16.
(Figure S5), indicated the vital role of cellulase proximity via cellulosome complexes of Clostridiales in initializing cellulose hydrolysis in this thermophilic consortium. Unlike active cellulase and hemicellulase expression, a transcriptional and genetic deficit of oligosaccharide-degrading enzymes, especially the typical beta-glycosidases of GH01, GH02 and GH03, were noted in this Clostridiales population (Figure 4 left). Spirochaetales and Thermotogales species are the other two active cellulose degraders in the thermophilic consortium that produce GH05 exocellulases and a wide range of beta-glycosidases. Another interesting transcriptional phenomenon occurred despite the general absence of hemicellulose substrates in this consortium, which is solely fed hexose-based polysaccharides (microcrystalline cellulose); hemicellulases (red-labeled GH families in Figure 4) of both Clostridiales and Spirochaetales exhibited noticeable transcriptional activities.

Expression of genes involved in methanogenesis. The transcriptional activities of genes involved in the thermophilic methanogenesis process were investigated in a similar manner based on RPKM-RNA and MRPKM. Genes involved in methanogenesis belong to the two archaeal orders Methanobacteriales and Methanosarcinales, with Methanobacteriales being the most prevalent by 60% (Figure 5, left). However, genes from the aceticlastic Methanosarcinales showed remarkably higher transcriptional activities (a six-fold higher MRPKM value) compared to those of the hydrogenotrophic Methanobacteriales. The overall aceticlastic pathway was three times more active than the hydrogenotrophic methanogenesis pathway in terms of the transcription of characteristic functional enzymes for each pathway (excluding mutual enzymes shared by hydrogenotrophic and aceticlastic pathways as represented by steps 3, 4 and 5 in Figure 5).

Discussion

The technical reproducibility of NGS-based metatranscriptomic sequencing is a topic lacking wide recognition. At this early stage of applying RNA-seq to metatranscriptomes, it is common to see frontier research without an emphasis on replication, especially for technical replicates. After an extensive literature survey, we found only one previous work that addressed the issue of technical replicates. Tsementzi, D. et al. noticed that the variability in technical replicates was almost as large as it was in the biological replicates. Their findings highlighted the presence of noticeable variation between mRNA technical replicates. In contrast, extensive studies have shown that the NGS-based transcriptome of a single organ (stem cells, liver and kidneys) or single species (yeast) is highly replicable with little technical variation. Therefore, we speculate that inadequate sequencing coverage other than RNA extraction contributes primarily to the variation between metatranscriptomic technical replicates as observed in this study (Table S1), and the enormous amount of mRNA molecules, at approximately $8 \times 10^{23}$ microbial mRNA molecules per liter of reactor sludge with volatile suspended solids of 800 mg/L ($\sim 200$ mRNA molecules per bacterial cell) makes it economically difficult to obtain the sequencing depth.
required to ensure the representation of the entire metatranscriptional profile\(^25\). The scale of variation observed in this study was comparable to that reported by Tsementzi, D. \textit{et al.}\(^{20}\). Therefore even the large NGS-based metatranscriptional libraries constructed here (3.8 million mRNA sequences after filtering ribosomal and small RNA content out of 29.5 million Illumina reads for each replicated RNA library, Figure S2) could at best provide a snapshot of the major activities of the community at a particular time point. Additionally, we picked the peak of biogas production to ensure active microbial metabolism in the microbiota because based on the long-term monitoring the peak biogas formation occurred almost simultaneously with the highest cellulose uptake rate in the sequencing batch reactor (SBR) cycle.

Because of the annotation difficulty caused by the short reads of Illumina sequencing, we compared the distribution and function of different phylotypes within the community at the order level. The adaptation of the thermophilic microbial consortium to cellulose yielded simplified communities in which members of \textit{Anaerolineae}, \textit{Clostridiales}, \textit{Bacteroidales} and \textit{Thermotogales} (listed in order of dominance) were the most prevalent populations. The consortium showed notable microbial diversity with 700 species (Shannon Index of 6.7) (Figure S6), which was comparable to that of the cow rumen (approximately 1000 OTUs)\(^5\) and the termite hindgut (800 OTUs and Shannon Index of 5.05\(^{11}\)). Compared to our previous community analysis at 120 days\(^{26}\), a significant decline in the \textit{Clostridiales} population and the consequent increase in \textit{Bacteroidales} and \textit{Anaerolineae} was noted in the SBR long-term operation at 545 days (this study). Eichorst, M. \textit{et al.}\(^{27}\) observed a similar community shift from \textit{Firmicutes} to a novel \textit{Bacteroidetes} population in an aerobic thermophilic microbiome that was adapted to microcrystalline cellulose, and they argued that the gradual accumulation of solubilized cellulose after initial hydrolysis was the reason for this trend. In contrast, our annotation-based comprehensive protein database (NCBI \textit{nr} database) did not disclose the direct metabolism advantage of \textit{Bacteroidales} or \textit{Anaerolineae} growing on soluble oligosaccharides (Figure 4). However, since unknown proteins (proteins that cannot be taxonomically classified), especially the beta-glycosidase of GH02 and GH03, played important roles in the oligosaccharide metabolism of the community, we cannot conclude that \textit{Bacteroidales} or \textit{Anaerolineae} are unimportant to oligosaccharide consumption. Instead, we observed the strong transcription of putative Sus-like polysaccharide utilization loci (PULs) (with RPKM-RNA comparable to GH09 observed in \textit{Clostridiales}) in an unclassified order of the \textit{Bacteroidetes} phylum following the identification protocol proposed by Rosewarne, C. \textit{et al.}\(^{28}\). This phenomenon not only consolidated the involvement of Sus-like PULs from \textit{Bacteroidetes} in cellulose-hydrolysis\(^{26,29}\) but also helped, in part, to reveal that the accumulation of the \textit{Bacteroidetes} population in this

Figure 4 | Distribution of various orders in the expression and genomic profiles of GH families related to lignocellulose decomposition in the thermophilic cellulose-degrading consortium. Left figure: The relative abundance of various GH families as presented by RPKM-DNA. Right figure: The overall transcriptional activities of different GH families as indicated by RPKM-RNA. The GH families were sorted according to their corresponding transcriptional activities with labels colored according their functions in cellulose deconstruction, as defined by Pope \textit{et al.}\(^3\): blue, Cellulases; orange, Endohemicellulases; green, Oligosaccharide-degrading enzymes; pink, Debranching enzymes. RPKM-RNA and RPKM-DNA values were compared as the percentage of all cellulosic degradation related genes. Based on the attached-hydrolysis model of cellulosic biomass as proposed by Lynd, L. \textit{et al.}\(^1\), the anaerobic hydrolysis process of microcrystalline cellulose was illustrated in the flowchart below, with the involved enzymes colored accordingly.
cellulolytic consortia may actually be related to its special cellulose hydrolysis capability. Additionally, we noticed the strong transcription of genes involved in cell protection against oxygen species in 

Clostridiales (Figure 3), indicating the ability of this population to grow in a facultative manner. We speculate that temporary oxygen exposure during sample preparation might cause the strong oxidative stress resistance observed in 

Bacteroidetes.

Clostridiales play vital roles in cellulose hydrolysis via cellulosome complexes. The proximity of endocellulases to heat shock protein explains the transcriptional advantage of these cellulases in response to high temperatures (Figure S4). The consistency of this genetic arrangement with that of 

Clostridium clairevum DSM 19732 circumcribed the phylogenetic origin of the active 

Clostridiales as a branching strain of this species. 

Thermotogales only exhibited an exo-cutting capacity towards the exposed chains produced by endocellulases of 

Clostridiales, suggesting their dependency on 

Clostridiales for carbohydrate metabolism. In return, by expressing beta-glycosidases, both 

Spirochaetales and 

Thermotogales could facilitate the microbial uptake of tetrasacharides and cellulose, the accumulation of which will otherwise cause strong inhibition on 

Clostridiales cellulases such as GH48. This synergistic mechanism could explain the earlier observed cellulose degradation enhancement in 

Clostridium thermocellum when the 

Sporochaeta phylotypes were present within its environment and help to shed light on the ubiquitous presence of symbiotic 

Spirochaetes in the gut of diverse termites.

Because the consortium was absolutely predominated by 

Clostridiales at 120 days of enrichment, which constituted up to 70% of the community, it is reasonable to speculate that the 

Clostridiales populations contributed the majority of both the hydrolysis and beta-sugar metabolism of the community at this earlier time point. In contrast, a major contribution of beta-glycosidase activity was observed in 

Thermotogales and 

Spirochaetales populations within 545 days in the metatranscriptome, suggesting that synergistic effects between cellulose-hydrolyzing 

Clostridiales and beta-sugar-consuming 

Thermotogales and 

Spirochaetales play a critical role over the long-term in the SBR. However, further validation of this hypothesis is required to reveal the dynamics of population involvement in cellulose bioconversion. Additionally, we observed a strong mobility via bacterial flagella in 

Thermotogales that was consistent with the general cell motility reported in the cellulosytic members of this order. This increase in the transcription of genes involved in cell motility highlights the importance of physical cell movement in facilitating the capture and breakdown of beta-sugars in 

Thermotogales.

Despite the general absence of hemicellulose substrate, a moderate transcription level of hemicellulases (labeled red in Figure 4) in 

Clostridiales and 

Spirochaetales was observed. Aside from the rare possibility of being housekeeping genes with expressions that are unaffected by experimental conditions, co-transcription with cellulases was more likely to be the machinery coordinating the hemicellulase regulation of 

Clostridiales and 

Spirochaetales populations. The lack of gene clusters for cellulases and hemicellulases had long been regarded as the genetic barrier for these two enzymes to co-transcribe in 

Clostridiales until the recent discovery of celC-glyR3-licA co-transcription in 

Clostridium thermocellum. However, this co-transcriptional regulation has never been observed for strains in 

Spirochaetales before. In the present study, we did not observe genetic clusters consisting of cellulase and hemicellulase on the scaffolds of either 

Clostridiales or 

Spirochaetales. Thus, the observed co-transcription pattern of hemicellulases and cellulases in these two populations could serve to consolidate the co-transcription machinery that coordinates hemicellulase activities in these populations.

Despite the fact that acetate is always the major intermediate product, the prevalence of exclusively hydrogenotrophic 

Methanobacteriales over aceticlastic methanogens (primarily 

Methanosarcinales) is common in the cellulose-based methanization system. Previously, the presence of over-competing hydrogenotrophic populations was interpreted as the influence of the inhibitory effect of environmental factors on the activity of the aceticlastic methanogens. Such factors include a high level of ammonia or volatile fatty acids (VFAs), extreme pH values or elevated temperatures. However, our results may overturn this hypothesis because the less prevalent aceticlastic population showed significantly higher overall transcriptional activity in methanogenesis than its hydrogenotrophic counterpart. These findings suggested that the minority of aceticlastic methanogens is not necessarily associated with a repressed metabolism. Instead, we speculate that other overlooked factors, such as a slower growth rate of aceticlastic methanogens, may actually shape the 

Methanobacteriales-dominated distribution of methanogens in the thermophilic cellulose methanization consortium. This finding also indicates the weakness of studying methanogenesis pathways based on the phylogenetic prevalence of representative methanogens.

Hydrogenotrophic 

Methanobacteriales often co-exist with syntrophic acetate-oxidizing bacteria (SAOB), which facilitate the fermentation of acetate to hydrogen and carbon dioxide. Some researchers claimed that enhanced acetate oxidation by SAOB is crucial for maintaining the effectiveness of this hydrogen-utilizing methanogenesis pathway; however, the attempt to enhance hydrogenotrophic methanogenesis via the bio-augmentation of the SAOB population was unsuccessful. Therefore, the synergistic mechanism between 

Methanobacteriales and syntrophic bacteria remains unclear. Unfortunately, owing to the limited identification of thermophilic SAOB and the lack of known enzymes that are specific to its acetate oxidizing pathway, active SAOB populations could not be clearly identified. However, the active symbiotic involvement of 

Nitrospirales in methanogenesis suggested that the population actively transcribed CoM-S-S-CoB heterodisulfide reductase, which regenerated coenzyme M and coenzyme B after the final methanogenesis reaction (Step 4 in Figure 5). This population also exhibited visible activity towards sulfate reduction, which consumes hydrogen for electrons and thus provides a thermodynamically favorable environment for acetate oxidation to take place.

By combining NGS-based metatranscriptomics and metagenomics, the present study provides initial transcriptional insights into the expressed biological functions during thermophilic cellulosic biomass methanization. Novel information on phylogeny and the functions of the 40,000 active genes identified in the metatranscriptome highlight the importance of complementarity interactions between microbial groups (Thermotogales, 

Spirochaetales, and unclassified order of 

Bacteroidetes and 

Clostridiales) for efficient cellulose hydrolysis. More importantly, we observed stronger transcriptional activities in genes that were involved in aceticlastic methanogenesis pathways when the aceticlastic 

Methanosarcinales are less dominant than their hydrogenotrophic counterparts, 

Methanobacteriales. This finding contradicts the earlier hypothesis on the repressed activity of aceticlastic methanogens and suggests that the less prevalent aceticlastic populations could play more important roles in cellulose methanogenesis than previously expected. More intensive biological and technical replication is required to reveal whether this is a general pattern in similar systems. Further metatranscriptomic investigation of aceticlastic methanogenesis activity in lignocellulosic biomass methanogenesis systems, especially in the large-scale digesters, could help to better explain the ecological contributions of different methanogens during these processes and eventually provide practical guidelines for microbial manipulation in cellulose decomposition.

Methods

Thermophilic cellulose methanization consortium. Anaerobic digestion sludge (ADS) collected from a local wastewater treatment plant (Shake Wu Hui Sewage Treatment Plant, Hong Kong) was enriched in a sequential batch reactor (SBR, working volume of 800 ml) at 55°C, with microcrystalline cellulose (20 μm, Sigma, USA) as the primary substrate at a loading concentration of 2 g/L, and glucose was
used as a co-substrate at a COD ratio of 10:1. Glucose was added to maintain the effectiveness of cellulose conversion in long-term operation; this effect from the monosugar co-substrate was discussed in detail in our previous study. The pH of the SBR at the beginning was approximately pH 7.0 and was automatically maintained above pH 6.0 throughout the fermentation. Each batch was suspended when no more gas was produced. After 545 days of enrichment, the batch cycle time was stabilized at approximately 47 h. The adapted culture showed a cellulose uptake rate of 1.2 g/(L d). Based on the continuous monitoring of SBR performance, we noticed that the peak biogas formation also corresponded to the maximum cellulose uptake speed in the SBR cycle. To ensure the active transcriptional status of genes encoding reactive enzymes that were involved in both cellulose hydrolysis and methanogenesis processes, sludge samples were collected from the SBR at peak gas formation, which occurred approximately 24 h after cellulose dosing.

**DNA extraction.** Two replicates of genomic DNA were individually extracted from 4 ml of sludge slurry (equal to an approximately 500-mg dry weight sample) from the thermophilic consortium with a FastDNA SPIN Kit for Soil (MP Biomedicals, LLC, Illkirch, France). The concentrations of extracted DNA replicates were 214.2 ng/ml with a 260/280 ratio of 1.88 and 203.7 ng/ml with a 260/280 ratio of 1.87 as quantified by a Nanodrop spectrophotometer (ND-1000, USA). Additionally, the quality of the extracted DNA and RNA was verified by electrophoresis (Figure S7).

**Total RNA isolation and cDNA synthesis.** At the same time point as the above DNA samples, total RNA isolation was performed with a TRIzol Plus RNA Purification Kit (Life Technologies, USA) immediately after sampling. In brief, a 12-ml (roughly equal to a 1500-μg dry weight sample) consortium sample was centrifuged at 13,000 × g for 2 min at 4 °C. Replicates of the biomass pellets were subjected to independent cell lysis and total RNA precipitation according to the manufacturer’s instructions. Next, genomic DNA was removed with an Amplification Grade DNase I Kit (Sigma, USA). RNA qualification was performed by electrophoresis (Figure S7). The integrity of the extracted RNA was checked with a Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). The extracted RNA replicates had RNA integrity numbers of 6.7 and 6.6, respectively, on a scale of 1–10, with 10 representing no degradation (Figure S8). First-strand and second-strand cDNA were synthesized with the Superscript III First-Strand Synthesis SuperMix (Invitrogen, CA, USA) and the Second-Strand cDNA Synthesis Kit, respectively (BeyoTime, Jiangsu, China).

**DNA, cDNA library construction and sequencing.** Independent libraries of ~180 bp were prepared for replicated DNA and cDNA samples by following the manufacturer’s instructions (Illumina). A sequencing depth of 2.8 G was applied to each metagenomic and metatranscriptomic replicate (Table S5).

**Quality control (QC) of metagenomic and metatranscriptomic datasets.** Metagenomic and metatranscriptomic raw reads derived from the Illumina Hiseq2000 platform were filtered for quality by discarding any read with 0.1% unknown nucleotides or with 50% nucleotides with a quality score lower than 20. Next, reads in the DNA dataset were checked for artificial duplicates that were produced during sequencing. The quality control for artificial duplicate reads were removed by a self-written script following the MG-RAST47 artifact identification protocol (See Table S5 for detailed dataset information after QC).

**Bioinformatic analysis.** Reproducibility analysis. Replicated libraries in the metagenomic dataset (named DNA datasets) and a metatranscriptomic dataset (named RNA datasets) were individually submitted to the MG-RAST server (Table S5 for accession number). Taxonomy annotation was performed using the Best Hit Classification Algorithm against the GenBank and SSU databases provided by the server. The cutoffs for searching against the SSU database had an E-value of 1E-20, a similarity of 60% and an alignment length of 15 amino acids. Functional annotation was conducted as follows: 1) a E-value of 1E-20, 2) a similarity of 60%, and 3) an alignment length of 15 amino acids. Functional annotation was conducted as follows: 1) a E-value of 1E-20, 2) a similarity of 60%, and 3) an alignment length of 15 amino acids.

**Figure 5** | Transcriptional activities (right) and genomic prevalence (left) of genes in the methanogenesis pathway. Genes were classified into various phylogenetic orders, as shown in the top figure, and colored according to their functions in the methanogenesis process (as adopted from the KEGG Methane Metabolism pathway), as shown in the flowchart (bottom figure).
against SEED subsystems using an E-value cutoff of 1E-5 and a hierarchical classification algorithm. The annotation based on the MG-RAST server was used for the reproducibility test.

**rRNA subtraction and annotation.** The analysis workflow is illustrated in Figure 1. First, rRNA sequences were separated from mRNA sequences in the post-QC RNA dataset by rRNAPicker. Sequences encoding rRNA genes within the DNA dataset (named rDNA sequences) were simultaneously isolated from the protein-coding sequences (named coding DNA sequences) in the DNA dataset by BLAST against the Silva SSU (version 11.1) database with an E-value cutoff of 1E-20.

To reveal the community structure, rDNA sequences were aligned to reference ribosomal RNA sequences in the Green Gene 13.5 database. The aligned rDNA reads (87% of all rDNA reads) were clustered using the Uclust_ref algorithm adopted by QIME 1.7.0 into operational taxonomic units (OTUs), with a similarity cutoff of 0.97.

**Assembly of metagenomic and metatranscriptomic datasets.** The artifact-filtered sequences of two replicated DNA datasets were assembled together using IDBA_UD with combined kmer sizes of 20, 40, 60, 80, and 100. Scaffolds longer than 300 bp were subjected to open reading frame (ORFs) prediction by MetaGeneMark. To ensure the annotation accuracy of the derived ORFs, only ORFs longer than 300 bp were kept for expression analysis.

**ORF annotation.** ORFs were searched against the nr database using Rapshear at an E-value cutoff of 1E-5. The search results were parsed by MEGAN4 to assign taxa (with the LCA algorithm) and KEGG/SEED-subsystem functions. Next, the amino acid sequences of the predicted ORFs were screened against the Pfam database version 26.0 by PfamScan (E-value cutoff of 1E-15) for particular glycoside hydrolase (GH) families and carbohydrate binding modules (CBM) as classified by the Carbohydrate Active Enzyme (CAZy) database.

**Quantifying transcriptional activity.** For quantification, the coding DNA and mRNA sequences were mapped back to ORFs with Bowtie2 allowing two mismatches over the whole sequence length. Based on the Bowtie alignment, RSEM was used to compute the RPKM (Reads Per Kilobase of transcript per Million mapped reads) values for DNA and RNA datasets (hereafter, RPKM-DNA and RPKM-RNA as defined in Equation S1 and S2). These two values were used as measurements for the relative abundance of a gene in the metagenome and the overall transcriptional activity of a gene in the metatranscriptome. Given the uneven richness of various genes in the metagenome, the overall transcriptional activity of each gene must be normalized by its relative abundance in the microbial community to reveal its absolute transcriptional activity. As a result, MRPKM (short for Metatranscriptomic RPKM) was proposed in the present study to evaluate the absolute transcriptional activity (which is equivalent to the expression level) of each gene in the metatranscriptome. MRRKM could be calculated as the ratio of RPKM-RNA to RPKM-DNA.

In the present study, the RPKM-DNA value sums for genes assigned to any taxa were used to reveal the prevalence of that population in the community, and the transcriptional activities of different populations were compared on the basis of the average MRPKM values of genes assigned. Additionally, the overall transcriptional activities of metabolic pathways were evaluated based on the sum of RPKM-RNA values for the genes involved.
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**Author contributions**

Y.X. conducted the experiments, analyzed the data and drafted the manuscript. Y.W., T.J. and H.Z. conducted the experiments. T.Z. and H.F. designed the experiments and contributed to the manuscript editing. All authors reviewed the manuscript.

**Additional information**

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