Genomics and prevalence of bacterial and archaean isolates from biogas-producing microbiomes

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Abstract

Background: To elucidate biogas microbial communities and processes, the application of high-throughput DNA analysis approaches is becoming increasingly important. Unfortunately, generated data can only partially be interpreted rudimentarily since databases lack reference sequences.

Results: Novel cellulolytic, hydrolytic, and acidogenic acetogenic Bacteria as well as methanogenic Archaea originating from different anaerobic digestion communities were analyzed on the genomic level to assess their role in biomass decomposition and biogas production. Some of the analyzed bacterial strains were recently described as new species and even genera, namely Herbinix hemicellulosilytica T3/55T, Herbinix luporum SD1DT, Clostridium bornimense M2/40T, Proteiniphilum saccharofermentans M3/6T, Fermentimonas caenicola ING2-ESB5, and Petrimonas mucosa ING2-ESA. High-throughput genome sequencing of 22 anaerobic digestion isolates enabled functional genome interpretation, metabolic reconstruction, and prediction of microbial traits regarding their abilities to utilize complex bio-polymers and to perform specific fermentation pathways. To determine the prevalence of the isolates included in this study in different biogas systems, corresponding metagenome fragment mappings were done. Methanoculleus bourgensis was found to be abundant in three mesophilic biogas plants studied and slightly less abundant in a thermophilic biogas plant, whereas Defluviitoga tunisiensis was only prominent in the thermophilic system. Moreover, several of the analyzed species were clearly detectable in the mesophilic biogas plants, but appeared to be only moderately abundant. Among the species for which genome sequence information was publicly available prior to this study, only the species Amphibacillus xylanus, Clostridium clariflavum, and Lactobacillus acidophilus are of importance for the biogas microbiomes analyzed, but did not reach the level of abundance as determined for M. bourgensis and D. tunisiensis.

Conclusions: Isolation of key anaerobic digestion microorganisms and their functional interpretation was achieved by application of elaborated cultivation techniques and subsequent genome analyses. New isolates and their genome information extend the repository covering anaerobic digestion community members.

Keywords: Anaerobic digestion, Biomethanation, Genome sequencing, Fragment recruitment, Defluviitoga tunisiensis, Methanoculleus bourgensis

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Background

Anaerobic digestion (AD) and biomethanation are commonly applied for the treatment and decomposition of organic material and bio-waste, finally yielding methane (CH₄)-rich biogas. The whole AD process can be divided into four phases: hydrolysis, acidogenesis, acetogenesis, and methanogenesis. Organic polymers are hydrolyzed into sugar molecules, fatty acids, and amino acids by hydrolytic enzymes. These metabolites are further degraded into the intermediate volatile fatty acids (VFA), acetate, alcohols, carbon dioxide (CO₂), and hydrogen (H₂) during acidogenesis and acetogenesis. Finally, CH₄ is produced either from acetate or from H₂ and CO₂. The challenges in each of these steps are reflected within the complexity of the microbial community converting biomass to biogas. Community compositions and dynamics were frequently investigated using different molecular biological methods. Among these, quantitative ‘real-time’ polymerase chain reaction (qPCR), e.g., [1–5], terminal restriction fragment length polymorphism (TRFLP) [6–8], and the 16S rRNA gene amplicon [9, 10] as well as metagenome sequencing approaches [9, 11–14] applying high-throughput (HT) technologies are the most commonly used methods. In these studies, bacterial members belonging to the classes Clostridia and Bacteroidia were identified to dominate the biogas microbial communities, followed by Proteobacteria, Bacilli, Flavobacteria, Spirochaetes, and Erysipelotrichi. Within the domain Archaea, members from the orders Methanomicrobiales, Methanospirillales, and Methanobacteriales were described to be abundant in biogas systems.

However, all recently published metagenome and metatranscriptome studies addressing elucidation of the biogas microbiology reported on a huge fraction of unassignable sequences suggesting that most of the microorganisms in biogas communities are so far unknown [15–18]. This is due to the limiting availability of reference strains and their corresponding genome sequences in public databases. Moreover, reference sequences are often derived from only distantly related strains isolated from different environments. For a better understanding of the microbial trophic networks in AD and any further biotechnological optimization of the biomethanation process, extension of public databases regarding relevant sequence information seems to be an indispensable prerequisite.

Recently, studies on the isolation, sequencing, and physiological characterization of novel microbial strains from various mesophilic and thermophilic biogas reactors were published, e.g., [18–29]. However, only few of these studies addressed the question of whether the described strain played a dominant role within the analyzed microbial community. Accordingly, the objective of this work was to sequence and analyze a collection of recently described as well as newly isolated bacterial and archaean strains from different biogas microbial communities to provide insights into their metabolic potential and life-style, and to estimate their prevalence in selected agricultural biogas reactors. In total, 22 different strains originating from meso- and thermophilic anaerobic digesters utilizing renewable primary products and/or organic wastes were analyzed. Based on genome analyses, isolates were functionally classified and assigned to functional roles within the AD process. Moreover, refinement of the metagenome fragment recruitment approach was used for the evaluation of an isolate’s prominence in different biogas communities. Overall the aim of this study was the considerable complementation of the reference repository by new genome information regarding AD communities.

Methods

Microbial strains used in this study and isolation of novel strains

In this study, 22 bacterial and archaean strains were studied from eight meso- and thermophilic, laboratory-scale and agricultural biogas plants (BGPs) utilizing renewable primary products as well as from three further AD sources (detailed information listed in Table 1). The strains Methanoculleus chikugoensis L21-II-0 and Sporanaerobacter sp. PP17-6a were isolated within this study as follows.

Methanoculleus chikugoensis L21-II-0 Reactor material was diluted fivefold in DSMZ medium 287 [30] containing 20 mM acetate and H₂/CO₂ as the only carbon and energy sources. Initial incubation occurred at 37 °C for 4 weeks without antibiotics. Subsequent cultivation was performed by successive transfer of culture aliquots after incubation periods of 4 weeks into the same medium supplemented with different combinations of the antibiotics tetracycline HCl (15 µg ml⁻¹), vancomycin HCl (50 µg ml⁻¹), ampicillin (100 µg ml⁻¹), and bacitracin (15 µg ml⁻¹) or with penicillin (350 µg ml⁻¹). After a total of 12 cultivation cycles, purity of the culture was confirmed by microscopic inspection and by denaturing gradient gel electrophoresis (DGGE) fingerprint analysis. Strain M. chikugoensis L21-II-0 is available from the Leibniz Institute German Collection of Microorganisms and Cell Cultures (DSMZ, Braunschweig, Germany) under the Accession No. DSM 100195. Sporanaerobacter sp. PP17-6a: Reactor material was diluted 5 × 10⁶-fold in DSMZ medium 120 [31]. After 4 weeks of incubation at 37 °C, an aliquot of the culture was transferred into the same medium supplemented with penicillin (350 µg ml⁻¹). Transfer and incubation in the same medium were repeated four times. Subsequently,
| Species and strain | Family | Origin | Reference for the isolation strategy or strain origin | Closest related NCBI GenBank entry with a validly published taxonomic affiliation | Similarity of 16S rRNA gene between isolate and GenBank entry (%) | NCBI GenBank entry of closest relative |
|-------------------|--------|--------|-----------------------------------------------------|---------------------------------------------------------------------------------|---------------------------------------------------------------------|----------------------------------------|
| Bacteria          |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Clostridium       | Clostridiaceae | 51.255499 6.396524 | Liquid pump/wet fermentation | Maize, pig manure, grass | 54 | ![8]( clostridium cellulosi AS1.1777 ) | 98.8 | LN881577 |
| cellulosi DG5     |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Clostridium sp.   |        |        |                                                     |                                                                                  |                                                                    |                                        |
| NC3               |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Clostridium       | Clostridiaceae | 51.255499 6.396524 | Liquid pump/wet fermentation | Maize, pig manure, grass | 54 | ![8]( clostridium putrefaciens DSM 1291 ) | 93.0 | NR113324 |
| bornimense M2/40T |        |        |                                                     |                                                                                  |                                                                    |                                        |
| O. thermocellum   |        |        |                                                     |                                                                                  |                                                                    |                                        |
| BC1               |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Proteinobacter sp. | Clostridales incertae sedis | 49.512893 7.083068 | CSTR, wet fermentation | Maize silage, grass, cattle manure | 39 | ![1]( proteinobacter ethanoligenes GW ) | 96.0 | NR044093 |
| DW1               |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Sporanaerobacter  |        |        |                                                     |                                                                                  |                                                                    |                                        |
| sp. PP17-6a       |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Herbinix          | Lachnospiraceae | 51.255499 6.396524 | Liquid pump/wet fermentation | Maize, pig manure, grass | 54 | ![8]( herbinix hemicellulosilytica T3/551 ) | 100 | LN626355 |
| hemicellulosilytica T3/55T |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Herbinix luppenum |        |        |                                                     |                                                                                  |                                                                    |                                        |
| SD1T              |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Propionispora     | Veillonellaceae | 48.3924 11.7569 | CSTR, wet fermentation | Maize silage, grass | 38 | ![8]( propionispora hippel KS1 ) | 95.0 | NR036875 |
| sp. 2/2-37        |        |        |                                                     |                                                                                  |                                                                    |                                        |
| Bacillus          | Bacillaceae | 48.3924 11.7569 | CSTR, wet fermentation | Maize silage, pig manure | 52 | ![8]( bacillus thermoamylovorans DK1T ) | 99.0 | NR029151 |
Table 1 continued

| Species and strain | Family       | Origin Location of BGP | Type of reactor | Fed substrate | T (°C) of reactor | Reference for the isolation strategy or strain origin | Closest related NCBI GenBank entry with a validly published taxonomic affiliation | Similarity of 16S rRNA gene between isolate and GenBank entry (%) | NCBI GenBank entry of closest relative |
|--------------------|--------------|-------------------------|-----------------|---------------|-------------------|------------------------------------------------------|---------------------------------------------------------------------------------|---------------------------------------------------------------------------------|-------------------------------------|
| **Proteiniphilum** | **Porphyromonadaceae** | 52.3871 13.0993 | Lab-scale UASS/wet fermentation | Maize silage, wheat straw | 37 | [26] | Proteiniphilum saccharofermentans M3/6<sup>T</sup> | 100 | KP233809 |
| **Porphyromonas**  | **Porphyromonadaceae** | 51.255499 6.396524 | Lab-scale CSTR/wet fermentation | Maize silage, pig manure, cattle manure | 37 | | Fermentimonas caenicola ING2-ES<sup>T</sup> | 100 | KP233810 |
| **Petrimonas**     | **Porphyromonadaceae** | 51.255499 6.396524 | Lab-scale CSTR/wet fermentation | Maize silage, pig manure, cattle manure | 37 | | Petrimonas mucosa ING2-ES<sup>T</sup> | 100 | KP233808 |
| **Defluviitoga**   | **Petrotogaceae** | 51.255499 6.396524 | Liquid pump/wet fermentation | Maize, pig manure, grass | 54 | [27] | Defluviitoga tunisiensis SulfLac<sup>T</sup> | 99.9 | NR122085 |
| **Methanobacterium** | **Methanobacteriaceae** | DSMZ<sup>α</sup> | | | 37 | [50] | Methanobacterium formicicum MF<sup>T</sup> | 100 | NR115168 |
| **Methanobacterium** | **Methanobacteriaceae** | 49.873559 6.481390 | CSTR, wet fermentation | Maize silage, grass, cattle manure | 40 | [21] | Methanobacterium formicicum MF<sup>T</sup> | 100 | NR115168 |
| **Methanobacterium** | **Methanobacteriaceae** | 49.512893 7.083068 | CSTR, wet fermentation | Maize silage, grass, cattle manure | 39 | | Methanobacterium formicicum MF<sup>T</sup> | 98.0 | NR115168 |
| **Methanobacterium** | **Methanobacteriaceae** | 53.736687 10.083949 | CSTR, dry fermentation | Household garbage | 37 | [18]<sup>α</sup> | Methanobacterium congolense<sup>C</sup> | 99.0 | NR028175 |
| **Methanothermobacter** | **Methanobacteriaceae** | 51.255499 6.396524 | Liquid pump/wet fermentation | Maize, pig manure, grass | 54 | [18]<sup>α</sup> | Methanothermobacter wolfeii VKM B-1829<sup>T</sup> | 100 | NR040964.1 |
### Table 1 continued

| Species and strain | Family | Origin | Location of BGP | Type of reactor | Fed substrate | T (°C) of reactor | Reference for the isolation strategy or strain origin | Closest related NCBI GenBank entry with a validly published taxonomic affiliation | Similarity of 16S rRNA gene between isolate and GenBank entry (%), NCBI GenBank entry of closest relative |
|--------------------|--------|--------|-----------------|-----------------|---------------|------------------|------------------------------------------------------|---------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Methanoculleus bourgensis MS2 | Methanomicrobiaceae | DSMZ | 51.255499 | Lab-scale CSTR/wet fermentation | Maize silage, pig manure, cattle manure | 37 | [49] Methanoculleus bourgensis MS2 | 100, NR042786 |
| Methanoculleus chikugoensis L21-II-0 | | | 6.396524 | | | 37 | This study Methanoculleus chikugoensis MG62 | 99.0, NR028152 |

CSTR, continuously stirred tank reactor; UASS, upflow anaerobic solid-state reactor

a DSMZ, Leibniz Institute DSMZ-German Collection of Microorganisms and Cell Cultures, Braunschweig, Germany
b Isolation strategy number four described in more detail by [18]
c Isolation strategy number eight (a) published in [18]
d Isolation strategy number five published in [18]
e Isolation strategy number seven published in [18]
f Isolation strategy number two published in [18]
g Isolation strategy number ten published in [18]
h Isolation strategy number eleven published in [18]
cultivation occurred by successive transfer of culture aliquots after incubation periods of 4 weeks into fresh medium supplemented with different combinations of antibiotics as mentioned above for isolation of the strain L21-II-0. After 14 cultivation cycles, isolation of the bacterial strain was performed by plating of the culture material on BBL™ Columbia Agar Base medium (Th. Geyer, Germany) supplemented with 5% laked horse blood (Oxoid, Germany). For purification, single colonies were picked and re-streaked, and incubation occurred at 37 °C.

**Phylogenetic classification of the analyzed bacterial and archaeal strains**

To determine the phylogenetic relationship between the different strains and closely related type strains, a phylogenetic tree was constructed. For this, the 16S rRNA gene sequences retrieved from the genome sequences of the analyzed strains were aligned using the SINA alignment service v.1.2.11, which is provided online [32]. Subsequently, the SINA alignment and the All-Species Living Tree LTPs123 [33] from the SILVA ribosomal RNA project [34], only consisting of the 16S rRNA gene sequences of validly described type strains, were loaded into the ARB program [35]. Finally, the SINA alignment was placed into the existing LTP tree using ARB’s parsimony method. Only type strains closely related to the corresponding isolate analyzed within this study are shown in the tree, whereas the remaining type strains were hidden manually applying “remove species from the tree” function implemented in ARB.

**Genomic DNA extraction, sequencing, and bioinformatic analyses of biogas community members**

Whole genome sequences of 13 strains, which were used in this study, were published previously (references given in Table 2). Genome sequencing of the following strains was performed within this study: *Proteiniborus* sp. DW1, *Clostridium* sp. N3C (DSM 100067), *Sporanaerobacter* sp. PP17-6a, *Proteiniphilum* saccharofermentans M3/6Ts, *Petrimonas mucosa* ING2-E5AT, *Methanobacterium formicicum* Mb9, *Methanobacterium congelense* Buetzberg, [36] *Methanothermobacter wolfeii* SIV6, and *M. chikugoensis* L21-II-0. In the case of *Clostridium* sp. N3C, *Sporanaerobacter* sp. PP17-6a, and *P. saccharofermentans* M3/6Ts, genomic DNA was extracted applying the innuPREP Bacteria DNA Kit (Analytik Jena, Germany). Genomic DNA of *P. mucosa* ING2-E5AT and *M. chikugoensis* L21-II-0 was extracted as described previously [37]. Genomic DNA of the strain *Proteiniborus* sp. DW1 was obtained applying the protocol published previously [19] and genomic DNA from *M. congelense* Buetzberg was extracted from 10 × 10 ml of a liquid culture using the Gene Matrix stool DNA purification kit (Roboklon, Germany). DNA of strain *M. wolfeii* SIV6 was obtained applying the FastDNA Spin Kit for Soil (MP Biomedicals).

For bacterial strains mentioned above, 4 μg of purified chromosomal DNA was used to construct an 8-k mate-pair sequencing library (Nextera Mate Pair Sample Preparation Kit, Illumina Inc., Eindhoven, Netherlands) and sequenced applying the mate-pair protocol on an Illumina MiSeq system. Sequencing libraries of the archaeal strains *M. chikugoensis* L21-II-0 and *M. wolfeii* SIV6 were made from 2 μg of chromosomal DNA using the TruSeq DNA PCR-Free Library Preparation Kit (Illumina Inc., Eindhoven, Netherlands) and sequenced applying the paired-end protocol on an Illumina MiSeq system.

The obtained sequences were de novo assembled using the GS de novo Assembler Software (version 2.8, Roche). An in silico gap closure approach was performed [38], which resulted in a draft genome sequence or in a circular chromosome. Gene prediction and annotation of the genomes were performed within the GenDB 2.0 annotation system [39]. Manual metabolic pathway reconstruction was carried out by means of the KEGG pathway mapping implemented in GenDB that compares gene sequences with the corresponding gene product sequences of the NCBI database, with pairwise protein sequence identity being at least 30%. To predict genes encoding carbohydrate-active enzymes, the carbohydrate-active enzyme database (CAZy) annotation webserver dbCAN [40] was used.

**Prevalence of the investigated strains within microbial communities of four different agricultural biogas plants applying the metagenome fragment recruitment approach**

To evaluate the prevalence of the 22 analyzed strains within the microbial communities of the four different BGPs described previously [41], the corresponding metagenome sequences available for these BGPs (metagenome Accession Nos. at the NCBI database: SRA357208-09, SRA357211, SRA357213-14, SRA357221-23) were mapped on the genome sequences of these isolates with FR-HIT (v0.7; [42]) to sensitively recruit also metagenomic reads with lower sequence identity (global alignment down to 75% nucleotide sequence identity; Additional file 1).

As a baseline to compare against, four known and abundant metagenome-assembled genomes (MAGs) published previously [41] were included (the fifth genome bin 206_Thermotogae matching *Defluviitoga tunisiensis* L3 was excluded, because it is contained in the isolate collection; Table 1).

Furthermore, Mash (v1.1; [43]) was used to quickly identify potentially abundant and publicly available genome sequences in RefSeq (as of June 14, 2016; [44]).
| Species and strain | Assembly status | Genome size (bp) | GC content (%) | No. of genes | No. of rRNA operons | No. of tRNA genes | No. of protein coding genes | EBI accession no. | References |
|-------------------|----------------|-----------------|----------------|--------------|---------------------|-------------------|--------------------------|-----------------|------------|
| **Bacteria**       |                |                 |                |              |                     |                   |                          |                 |            |
| *Clostridium cel- | CCC n.a.       | 2,229,578       | 44.15          | 2088         | 6                   | 59                | 2017                     | ERP006074       | [53]       |
| lulosi* DG5        |                |                 |                |              |                     |                   |                          |                 |            |
| *Clostridium sp.  | Draft genome   | 3,037,440       | 32.43          | 2880         | 3                   | 66                | 2880                     | FMJL01000001–   | This study  |
| N8C                |                |                 |                |              |                     |                   |                          | FMJL01000109    |            |
| *Clostridium borni- | CCC n.a.      | 2,917,864       | 29.78          | 2694         | 8                   | 56                | 2613                     | HG917868        | [37]       |
| *merense M2405    | Chromid        | 699,161         | 28.09          | 680          | 0                   | 0                 | 680                      | HG917869        |            |
| *Clostridium therm- | Draft genome   | 3,454,918       | 39.10          | 3094         | 4                   | 52                | 3095                     | CBQ0010000001–  | [61]       |
| *ocellum BC1       |                |                 |                |              |                     |                   |                          | CBQ0010000139   |            |
| *Proteiniborus sp. | Draft genome   | 3,121,392       | 32.44          | 2795         | 3                   | 40                | 1793                     | FMDO01000001–   | This study  |
| DWV1a              |                |                 |                |              |                     |                   |                          | FMDO01000162    |            |
| *Sporanaerobacter* | Draft genome   | 3,296,762       | 33.45          | 3148         | 1                   | 46                | 3148                     | FMIF01000001–   | This study  |
| sp. PP17-6a        |                |                 |                |              |                     |                   |                          | FMIF01000053    |            |
| *Herbinix hemi-   | Draft genome   | 3,037,031       | 36.69          | 2681         | 4                   | 35                | 1726                     | CVTD20000001–   | [24]       |
| *cellulosilytica*  |                |                 |                |              |                     |                   |                          | CVTD20000035    |            |
| T3/SS3T            |                |                 |                |              |                     |                   |                          |                 |            |
| *Herbinix lupu-   | CCC n.a.       | 2,609,352       | 35.25          | 2362         | 4                   | 53                | 1517                     | LN879430        | [78]       |
| *romum SD104      |                |                 |                |              |                     |                   |                          |                 |            |
| *Peptoniphilaceae* | CCC n.a.       | 1,601,846       | 34.85          | 1541         | 4                   | 53                | 1476                     | LM997412        | [22]       |
| bacterium str.     |                |                 |                |              |                     |                   |                          |                 |            |
| ING2-D1G           |                |                 |                |              |                     |                   |                          |                 |            |
| *Propionispora*    | Draft genome   | 4,122,013       | 45.58          | 3690         | 1                   | 76                | 2685                     | CYSPO1000001–   | [29]       |
| sp. 2/2–37         |                |                 |                |              |                     |                   |                          | CYSPO1000043    |            |
| *Bacillus*         | Draft genome   | 3,708,331       | 37.28          | 3472         | 10                  | 59                | 2957                     | CCRF01000001–   | [79]       |
| *thermoamyloly-  |                |                 |                |              |                     |                   |                          | CCRF01000106    |            |
| vorans 1A1         |                |                 |                |              |                     |                   |                          |                 |            |
| *Proteiniphilium*  | CCC n.a.       | 4,414,963       | 43.63          | 3450         | 3                   | 48                | 3447                     | LT605205        | This study  |
| *saccharofermen-  |                |                 |                |              |                     |                   |                          |                 |            |
| tans M3/65         |                |                 |                |              |                     |                   |                          |                 |            |
| *Fermentimonas*    | CCC n.a.       | 2,808,926       | 37.30          | 2455         | 2                   | 44                | 2405                     | LN515532        | [25]       |
| *caenicola ING2-ESB5|                |                 |                |              |                     |                   |                          |                 |            |
| **Petrimonas**     | CCC n.a.       | 3,362,317       | 48.24          | 2693         | 2                   | 46                | 2693                     | ERS1319466      | This study  |
| *mucosa ING2-ESA1 |                |                 |                |              |                     |                   |                          |                 |            |
Table 2 continued

| Species and strain                                      | Assembly status | Genome structure | Genome size (bp) | GC content (%) | No. of genes | No. of rRNA operons | No. of tRNA genes | No. of protein coding genes | EBI accession no. | References |
|--------------------------------------------------------|-----------------|------------------|------------------|----------------|--------------|---------------------|-------------------|--------------------------|--------------------|------------|
| *Defluvitoga tunisiensis* L3                             | CCC             | n.a.             | 2,053,097        | 31.38          | 1881         | 3                   | 47                | 1815                     | LN824141           | [23]       |
| Archaea                                                 |                 |                  |                  |                |              |                     |                   |                          |                    |            |
| Methanobacterium formiciculum *MS1*                      | CCC             | n.a.             | 2,478,074        | 41.23          | 2409         | 2                   | 44                | 2100                     | LN515531           | [80]       |
| Methanobacterium formicicum *Mb9*                        | CCC             | n.a.             | 2,494,510        | 41.14          | 2416         | 2                   | 43                | 2126                     | ERS549551          | This study  |
| Methanobacterium sp. *Mb1*                               | CCC             | n.a.             | 2,029,766        | 39.74          | 2021         | 2                   | 41                | 1689                     | HG425166           | [19]       |
| Methanobacterium congolense *Buetzberg*                  | CCC             | n.a.             | 2,459,553        | 38.48          | 2351         | 3                   | 41                | 2351                     | LT607756           | This study  |
| Plasmid                                                 |                 |                  | 18,118           | 36.05          | 24           | 0                   | 0                 | 24                       | LT607757           |            |
| Methanothermobacter wolfei *SIV6*                        | CCC             | n.a.             | 1,686,891        | 48.89          | 1793         | 2                   | 36                | 1444                     | ERS1319767          | This study  |
| Methanoculleus bourgensis *MS2*                          | CCC             | n.a.             | 2,789,773        | 60.64          | 2586         | 1                   | 45                | 2586                     | HE964772           | [81]       |
| Methanoculleus chikugoensis *L21-11-0*                   | Draft genome    | 70               | 2,649,997        | 61.83          | 2671         | 1                   | 45                | 2671                     | FMID01000001–FMID01000070 | This study  |

CCC, circulary closed chromosome; n.a., not applicable

The strain *Protoniborus sp. DW1* was cultivated together with *Methanobacterium sp. Mb1*; the DW1 genome sequence was recovered from sequencing of a mixed culture consisting of strains DW1 and Mb1.
The meaning of abundance in this context refers exclusively to the number of metagenome sequences mapped to the genome sequence. For a sketch size of 1,000,000 and a k-mer size of 21, pairwise distances between the metagenomic read sets and all 5061 genomes in RefSeq (plus, as a control, the 22 strains from this study) were calculated. Requiring a minimum of 20 k-mer hits not only confirmed the potential relevance of the selected 22 strains, but additionally identified 46 publicly available strains from RefSeq for further analyses.

All metagenome sequences available for the four BGP were mapped on the genome sequences of these isolates, the four MAGs, and the 46 reference strains with Kallisto [45] (v0.43.1). For each genome, the GPM (genomes per million) values were calculated using the TPM (transcripts per million) values reported by Kallisto (see Additional file 3).

**Results and discussion**

**Selection of a set of microbial isolates from different biogas-producing communities**

Limited availability of genome sequence information in public databases for AD community members generally constrains the interpretation of metagenomic and metatranscriptomic data of such data leading to large amounts of non-classifiable metagenome sequences from AD habitats [15–18, 46, 47]. Accordingly, parallel application of both traditional culturomics [48] as well as molecular analysis combined with HT sequencing techniques is necessary for detailed studies of complex microbial biogas consortia. Applying 16 different isolation strategies, bacterial and archaeal isolates were obtained from different mesophilic and thermophilic production- and laboratory-scale BGP (Table 1). Furthermore, two archaeal members, namely *M. bourgensis* MS2T [49] and *M. formicicum* MF [50], were obtained from the DSMZ and included in this study as the reference strains for methanogenic *Archaea* since they were also isolated from AD communities. German BGP sampled for this study differed in utilized substrates ranging from maize silage, grass, and wheat straw to cattle and/or pig manure. Moreover, one digester analyzed was fed with organic residues and waste material as substrate. Additionally, a bio-waste compost treatment site close to the city of Munich (Germany) was sampled to isolate cellulolytic bacteria. Besides different renewable biomass sources utilized for the AD process, the biogas reactors differed regarding digester design, fermentation technology, and the applied temperature regime ranging from 37 to 54 °C.

This study comprises the analysis of 15 bacterial strains classified as belonging to the phyla *Firmicutes*, *Thermotogae*, and *Bacteroidetes* and seven archaeal isolates of the phylum *Euryarchaeota*. Details on all isolates of this study, their taxonomy, their origin, and the respective isolation strategy applied are provided in Table 1.

**Phylogenetic classification of the microbial isolates selected from different biogas communities**

To determine the taxonomic position of the strains analyzed, their 16S rRNA gene sequences were compared to the corresponding sequences from closely related type strains deposited in the SILVA database (Fig. 1). The calculated phylogenetic tree comprises four main groups representing the phyla *Bacteroidetes*, *Firmicutes*, *Thermotogae*, and *Euryarchaeota*. Among the *Bacteroidetes* members, the strains *P. saccharofermentans M3/6T*, *P. mucosa* ING2-E5AT, and *Fermentimonas caenicola* ING2-E5BT were recently described as novel species and were suggested to participate in hydrolysis and acidogenesis of the AD process [26].

Most of the bacterial strains analyzed were allocated to the phylum *Firmicutes*, and within this taxon to the classes *Clostridia*, *Bacilli*, *Tissierellia*, and *Negativicutes*. A diverse group of isolates belong to the class *Clostridia*. They are related to characterized species such as *Clostridium cellulosis* (also denominated as ‘Ruminiclostridium’ cellulosis), *Clostridium thermocellum* (also denominated as ‘Ruminicolosporidium’ thermodendrum) [51], *Clostridium cellulovorans*, and *Clostridium bornimense*. The latter one was recently described as novel species [20]. All mentioned species represent lignocellulosic biomass degraders [20, 52, 53]. Two other *Clostridia* isolates, namely T3/55T and SD1D2T, were recently assigned to the species *Herbinix hemicellulositica* [54] and *Herbinix luporum* [55], respectively, of the new genus *Herbinix*. Both strains are distantly related to the type strain *Mobilitalea sibirica* P3M-3T [56] and were described to be involved in thermophilic degradation of lignocellulosic biomass.

The isolates 1A1, ING2-D1G, and 2/2-37 are closely related to the species *Bacillus thermoanviylovorans* (class *Bacilli*), *Peptoniphilus indolicus* (class *Tissierellia*), and *Propionispora hippie* (class *Negativicutes*), respectively. The corresponding reference strains were described to perform hydrolytic and acidogenic functions in the AD process [57–59].

Another isolate from a thermophilic BGP was classified as *D. tunisiensis* (phylum *Thermotogae*, class *Thermotogae*) representing an isolated branch of the bacterial part of the tree (Fig. 1). The strain *D. tunisiensis* L3 was described to be adapted to high temperatures and able to utilize different complex carbohydrates to produce ethanol, acetate, H2, and CO2 [27, 28]. The latter three metabolites represent substrates for methanogenic *Archaea*.

The strains *Sporanaerobacter* sp. PP17-6a and *Peptoniphilaceae* bacterium str. ING2-D1G are only distantly related to known bacterial species of the family...
Clostridiales incertae sedis and Peptoniphilaceae (90–91% identity), respectively, suggesting that they represent new species.

The fourth group of the phylogenetic tree represents methanogenic Archaea classified as members of the classes Methanomicrobia and Methanobacteria (both belonging to the phylum Euryarchaeota). Members of these classes were described to perform hydrogenotrophic methanogenesis utilizing CO₂ and H₂ as substrates for CH₄ synthesis [18, 21].

Genome sequence analyses of the whole set of microbial isolates selected

To gain insights into the functional potential of all strains listed in Table 1, their genomes were completely sequenced by application of HT sequencing technologies. Genome sequence information provides the basis for metabolic reconstruction and assignment of functional roles within the AD process, thus enabling biotechnological exploitation of genome features involved in fermentation processes utilizing renewable primary products.

Out of 22 genome sequences, nine, namely those of Proteiniborus sp. DW1, Clostridium sp. N3C, Sporanaerobacter sp. PP17-6a, P. saccharofermentans M3/6⁸T, P. mucosa ING2-E5A⁸T, M. formicicum Mb9, M. congolense Buettelberg, M. wolfei SIV6, and M. chikugoensis L21-II-0, were newly established in this study. Genome sequences of the remaining 13 strains were published previously mainly in the form of Genome Announcements (for references, refer to Table 2). The genome sequences of the microorganisms analyzed were established on an Illumina MiSeq system. In silico and PCR-based gap closure strategies resulted in 13 finished and nine draft genome sequences. General genome features, e.g., genome structure, assembly status, size, GC content, and numbers of predicted genes, are summarized in Table 2. Established genomes range in size from 1.6 to 4.4 Mb and feature GC contents from 28.09 to 61.83%. Moreover, C. bornimense M2/40⁸T, in addition to the chromosome, harbors a 699,161-bp chromid (secondary replicon) in its genome containing 680 coding sequences [37]. The methanogen M. congolense Buettelberg also harbors an accessory genetic element, namely a plasmid featuring a size of 18,118 bp. Genome annotation applying the GenDB 2.0 platform enabled functional interpretation of genes and reconstruction of metabolic pathways involved in the AD process. Genome analyses provided insights into the lifestyle and functional roles of bacterial and archaeal strains.

Fig. 1 Phylogenetic diversity of archaeal and bacterial strains analyzed in this study in relation to the corresponding type species. The program ARB [35] was applied to construct the phylogenetic tree based on the full-length 16S rRNA gene sequences obtained from the strain’s genome sequences and in the case of closely related type species from the SILVA database [34]. The scale bar represents 1% sequence divergence.
Screening of the subset of bacterial genomes to identify genes encoding carbohydrate-active enzymes potentially involved in biomass degradation

To elucidate genes encoding carbohydrate-active enzymes, functional genome annotation applying the HMM-based carbohydrate-active enzyme annotation database dbCAN [40] was performed (Fig. 2). Between 71 and 358 genes encoding enzymes or modules with predicted activity on carbohydrates were identified in each of the bacterial strains analyzed. Among them are dockerin-containing glycoside hydrolases (GH), representing putative cellulosomal enzymes, corresponding cohesin-containing scaffolds, enzymes acting on large carbohydrate molecules, and carbohydrate-binding motifs involved in sugar binding. The obtained results separate the analyzed strains into two groups: group I strains were predicted to degrade cellulose and hemicellulose, whereas group II strains represent secondary fermentative bacteria relying on metabolites (mainly mono-, di-, and oligosaccharides) produced by group I members (as obvious presence of cellulolytic genes). The Clostridiaceae strains DG5, T3/55T, SD1DT, M2/40T, and BC1 harbor a more diverse repertoire of genes involved in the degradation of complex polysaccharides such as cellulose (GH5, GH8, GH9, GH48), xylan (GH10, GH11), and cellobiose- or celloextrin-phosphorylase genes (GH94). Furthermore, genes for cohesin-containing putative scaffolds and the corresponding dockerin-containing glycoside hydrolases with a potential for cellulosome formation were also identified in the genomes of these strains. Previous studies reported on the importance of the phylum Firmicutes for hydrolysis of cellulosic material in biogas digesters [12, 60]. In particular, Clostridiaceae and Ruminococcaceae members are involved in this first step of biomass digestion [11, 18]. Clostridiaceae strains Proteiniborus sp. DW1 and Clostridium sp. N3C were predicted to represent non-cellulolytic isolates (Fig. 2), whereas the cellulolytic strain C. thermocellum BC1 [61] is known to be a very efficient cellulose degrader since it encodes cellulosome components and is able to degrade hemicelluloses and pectins [60]. In contrast to the cellulolytic Clostridiaceae, the Porphyromonadaceae members, namely P. saccharofermentans M3/6T, P. mucosa ING2-E5A T, and F. caenicola ING2-E58T, encode enzymes predicted to degrade pectins and a variety of hemicelluloses (GH16, GH26, GH30, GH53, GH74). These strains do not seem to be able to hydrolyze arabininoxylan (lack of GH10, GH11) and crystalline cellulose (lack of GH48). Likewise, D. tunisiensis L3 (Petrogactaceae family) also possesses a large set of genes predicted to facilitate cleavage of a variety of sugars including cellobiose, arabinosides (GH27), chitin (GH18), pullulan and starch (GH13), and lichenan (GH16) [28].

Another strain supposed to represent a secondary fermentative bacterium, namely B. thermoamylovorans 1A1 (Bacillaceae family), may contribute to oligosaccharide degradation with genes for GH1, GH2, GH3, or GH43 enzymes. In addition, genes required for growth on cellobiose are present in its genome. Considering the fact that strain 1A1 originally was isolated from a co-culture also containing C. thermocellum [61], it is assumed that B. thermoamylovorans 1A1 further metabolizes cellobiose produced by cellulolytic Clostridia.

![Fig. 2](image_url) Diversity of genes encoding carbohydrate-active enzymes (CAZymes) predicted to be involved in hydrolysis and/or rearrangement of glycosidic bonds for each bacterial isolate studied. The screening for the presence of CAZymes was accomplished applying the HMM-based (Hidden-Markov-Model-based) carbohydrate-active enzyme annotation database dbCAN [40]. The numbers of bacterial genes belonging to a corresponding glycosidic hydrolase (GH) family are given in the fields.
Members of the genus *Propionispora* (*Veillonellaceae*) previously were identified in AD communities [62] and predicted to utilize mostly sugars and sugar alcohols, e.g., glucose, fructose, xylitol, or mannitol for growth [59]. The strain *Propionispora* sp. 2/2–37 analyzed in this study additionally harbors genes encoding enzymes participating in cellubiose, starch, and chitin degradation as determined by means of the CAzy analysis.

In contrast, the results obtained for *Peptoniphilaceae* bacterium str. ING2-D1G showed that this bacterium does not encode enzymes involved in the degradation of complex carbohydrates. However, the strain ING2-D1G encodes all enzymes needed to utilize amino acids and monomeric carbohydrates as a carbon source [22]. Its function in the anaerobic digestion process can be hypothesized to be associated with acidogenesis, which was supported by reconstruction of corresponding metabolic pathways.

**Prediction of fermentation pathways based on sequence information for the subset of bacterial genomes**

Bacteria involved in AD perform a number of different fermentation pathways to recycle reduction equivalents that are produced in the course of metabolite utilization. To determine the fermentation type and the functional role of a given isolate within the biogas process, enzymes encoded in its genome were assigned to selected fermentation pathways represented in the KEGG database (Table 3, Additional file 2 and Fig. 3). Pathways leading to propionate, ethanol, formate, butyrate, acetate, and lactate synthesis were considered in this approach.

Certain bacteria are able to convert sugars, acids, alcohols, or amino acids to propionic acid under anaerobic conditions utilizing the methylmalonyl-CoA or the acrylyl-CoA pathways of the propanoate metabolism [27]. Among the analyzed bacteria, the strains *Propionispora* sp. 2/2–37, *P. saccharofermentans* M3/6T, *P. mucosa* ING2-E5AT, and *F. caenicola* ING2-E5BT encode all enzymes of the methylmalonyl-CoA pathway for the production of propionic acid from pyruvate. Only the strain *Proteinibacterium* sp. DW1 was predicted to utilize lactate for propionic acid production via the acrylyl-CoA pathway. Since the enrichment of propionic acid was described as an indicator for process imbalance [26, 27], data on the physiology of propionic acid-producing bacteria can be valuable for the optimization of the biogas plants.

Butyric acid-forming bacteria in biogas systems have been insuffciently characterized so far [27]. Genes encoding enzymes required for butyric acid formation via the butanoate pathway were found in the genomes of the strains *Propionispora* sp. PP16-6a, *Peptoniphilaceae* bacterium str. ING2-D1G, *C. borinense* M2/40T, *P. saccharofermentans* M3/6T, *Clostridium* sp. N3C, *P. mucosa* ING2-E5AT, *F. caenicola* ING2-E5BT, and *B. thermoamylovorans* 1A1. Butanoate production was recently described for the strains *H. luporum* SD1DT [55] and *H. hemicellulosilytica* T3/55T [54]. However, the genomes of these bacteria only encode the last two enzymes of the butanoate pathway, namely the phosphate butyryl transferase Ptb and butyrate kinase Buk, predicted to be responsible for butanoate synthesis in these strains.

During acidogenesis, volatile organic compounds such as ethanol, acetate, and formate are produced in the course of the AD process. The latter two metabolites are substrates for methanogenic *Archaea*. Analysis of pathways involved in ethanol, acetate, and formate synthesis, i.e., the mixed-acid fermentation, revealed that all analyzed bacteria harbor genes encoding enzymes of this pathway (see Additional file 2). With the exception of the *Peptoniphilaceae* bacterium str. ING2-D1G, in all other isolates the necessary genes to produce ethanol from pyruvate were identified. Moreover, genes encoding enzymes participating in formate production were found in the *C. cellulosi* DG5, *C. borinense* M2/40T, *D. tunisiensis* L3, *C. thermocellum* BC1, and *B. thermoamylovorans* 1A1 genomes. Furthermore, all analyzed bacteria were predicted to be able to produce acetate from acetyl-CoA. Genes encoding the enzymes phosphate acetyltransferase Pta (EC: 2.3.1.8) and acetate kinase Ack (EC: 2.7.2.1), converting acetyl-CoA to acetyl phosphate and subsequently to acetate, were found. In addition, genes encoding the enzymes pyruvate decarboxylase Pdc (EC: 4.1.1.1) and alcohol dehydrogenase Adh (EC: 1.1.1.1), converting pyruvate to acetaldehyde and finally to ethanol, were found in all genomes with the exception of the strain *Peptoniphilaceae* bacterium str. ING2-D1G, which does not possess an *adl* gene. Surprisingly, in the case of the strains *P. mucosa* ING2-E5AT, *F. caenicola* ING2-E5BT, and *P. saccharofermentans* M3/6T, no ethanol production was observed in growth experiments [26]. Possibly, the growth conditions tested might not be favorable to support ethanol synthesis.

Many bacterial species produce 2,3-butanediol under anaerobic conditions from glucose, with *Klebsiella oxytoca* and *Bacillus licheniformis* described as efficient 2,3-butanediol producers [64]. Among the bacteria analyzed, only *Propionispora* sp. 2/2–37 harbors a full set of genes encoding all necessary enzymes (refer to Additional file 2).

Lactic acid was found to be the main fermentation product from household waste digestion [65]. Members of the genera *Bacillus*, *Lactobacillus*, *Leuconostoc*, *Pediococcus*, and *Streptococcus* were previously described to produce lactic acid from several types of sugars [12, 47, 66]. To determine whether the analyzed bacteria have the potential to produce lactic acid, the genomes were
Table 3  Prediction of bacterial fermentation pathways as deduced from genome sequence information

| Pathway analyzed                        | Predicted product after fermentation | Clostridium cellulosi DG5 | Clostridium sp. N3C | Clostridium bornimense M2/40T | Clostridium thermocellum BC1 | Proteinibacter sp. PP17-6a | Sporanaerobacter sp. PP17-6a | Herbinix hemi-cellulosilytica T3/S5T | Herbinix luporum SD1D T |
|----------------------------------------|--------------------------------------|---------------------------|---------------------|-------------------------------|-----------------------------|----------------------------|-------------------------------|------------------------------------|-------------------------|
|                                        |                                      | GP EP<sup>a</sup>         | GP EP<sup>b</sup>   | GP EP<sup>b</sup>             | GP EP<sup>b</sup>           | GP EP<sup>b</sup>           | GP EP<sup>b</sup>           | GP EP<sup)b</sup>                  | GP EP<sup>b</sup>          |
| Propionic acid fermentation<sup>c</sup>|                                      |                           |                     |                               |                             |                           |                               |                                     |                         |
| Acrylyl-CoA pathway                    | Propionic acid                       | –                         | ND                  | – NC (D)                      | –                           | – NA                       | – NA                          | – NA                               | – NC (D)                 |
| Methylmalonyl-CoA pathway              |                                      |                           |                     |                               |                             |                           |                               |                                     |                         |
| Ethanol fermentation                   | Ethanol                              | +                         | D                   | +                             | +                           | +                          | +                             | + D                                | + D                     |
| Formic acid fermentation               |                                      |                           |                     |                               |                             |                           |                               |                                     |                         |
| 2,3-Butanediol fermentation            | 2,3-Butanediol                       | –                         | ND                  | – ND                          | –                           | –                          | – ND                          | – ND                               |                         |
| Formic acid                            |                                      |                           |                     |                               |                             |                           |                               |                                     |                         |
| CO<sub>2</sub> and H<sub>2</sub>       |                                      | –                         | –                   | +                             | +                           | +                          | +                             | +                                  |                         |
| Mixed-acid fermentation                | Ethanol                              | +                         | D                   | +                             | +                           | +                          | +                             | + D                                | + D                     |
|                                        | Acetate                              | +                         | +                   | + ND                          | +                           | +                          | +                             | +                                  | +                      |
|                                        | Lactate                              | +                         | ND                  | +                             | +                           | +                          | –                             | + ND                               | + ND                   |
|                                        | Succinate                            | –                         | –                   | +                             | –                           | –                          | –                             | –                                  | +                      |
| Butyric acid fermentation              | Butyrate                             | –                         | +                   | +                             | –                           | +                          | +                             | + D                                | + D                     |
|                                        | Acetate                              | +                         | D                   | +                             | +                           | +                          | +                             | +                                  | +                      |
|                                        | Lactic acid fermentation              |                                        |                    |                               |                             |                           |                               |                                     |                         |
| Homolactic acid fermentation           | Lactate                              | +                         | ND                  | +                             | +                           | –                          | –                             | + ND                               | + ND                   |
| Heterolactic acid fermentation         |                                      |                           |                     |                               |                             |                           |                               |                                     |                         |
|                                        | Acetate                              | –                         | –                   | – D                           | –                           | –                          | –                             | –                                  | –                      |
|                                        | Lactate                              | +                         | D                   | +                             | +                           | +                          | +                             | + D                                | + D                     |
|                                        | Ethanol                              | +                         | +                   | +                             | +                           | +                          | +                             | +                                  | +                      |
Table 3 continued

| Pathway analyzed                | Predicted product after fermentation | Peptoniphilaceae bacterium str. ING2-D1G | Propionispora sp. 2/2-37 | Bacillus thermoaerophilus 1A1 | Proteiniphilium saccharofermentans M3/6 | Fermenimonas caenicola ING2-ESA | Petrimonas mucosa ING2-ESAf | Defluvitoga tunisiensis L3 |
|---------------------------------|--------------------------------------|------------------------------------------|--------------------------|-------------------------------|----------------------------------------|-------------------------------|---------------------------|--------------------------|
|                                 |                                      | GP | EP | GP | EP | GP | EP | GP | EP | GP | EP | GP | EP |
| Propionic acid fermentation      |                                      |    |    |    |    |    |    |    |    |    |    |    |    |
| Acrylyl-CoA pathway              | Propionic acid                       | −  | ND | −  | D  | −  | ND | −  | D  | −  | D  | −  | D  | ND |
| Methylmalonyl-CoA pathway        |                                      |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Ethanol fermentation             | Ethanol                              | −  | +  | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  |
| Formic acid fermentation         |                                      |    |    |    |    |    |    |    |    |    |    |    |    |    |
| 2,3-Butanediol fermentation      | 2,3-Butanediol                       | −  | +  | ND | +  | ND | −  | −  | −  | −  | −  | −  | −  | −  |
|                                   | Formic acid                          | −  | +  | +  | +  | +  | +  | −  | +  | +  | +  | +  | +  | −  |
|                                   | CO₂ and H₂                           | −  | −  | −  | −  | −  | −  | −  | +  | +  | +  | +  | +  | +  |
| Mixed-acid fermentation          | Ethanol                              | −  | +  | D  | +  | D  | +  | +  | +  | +  | +  | +  | +  | +  |
|                                   | Acetate                              | +  | D  | +  | +  | +  | D  | +  | D  | +  | D  | +  | D  | +  |
|                                   | Lactate                              | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  |
|                                   | Succinate                            | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  |
| Butyric acid fermentation        | Butyrate                             | +  | D  | +  | D  | +  | +  | +  | +  | D  | +  | D  | +  | D  |
| Homoaacetogenesis                | Acetate                              | +  | D  | +  | D  | +  | D  | +  | D  | +  | D  | +  | D  | +  |
| Lactic acid fermentation         | Homolactic acid fermentation         | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  |
|                                   | Heterolactic acid fermentation       | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  | ND | +  |
|                                   | Lactate                              | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  | −  |
|                                   | Acetate                              | +  | D  | +  | D  | +  | D  | +  | D  | +  | D  | +  | D  | +  |
|                                   | Ethanol                              | +  | ND | +  | +  | +  | +  | +  | ND | +  | ND | +  | ND | +  |

Genomic loci encoding enzymatic functions participating to the corresponding fermentation type for each bacterial strain analyzed are listed in Additional file 2.

+ , synthesis of the corresponding fermentation end-product is predicted; − , pathway incomplete or misses key enzymes, the synthesis of the corresponding fermentation end-product is doubtful; EP , experimental proof; D, the corresponding fermentation product has been experimentally detected; GP, genes predicted applying metabolic reconstruction within the GenDB 2.0 system [39]; NA, not analyzed; NC, not confirmed; ND, fermentation product has been experimentally not detected

a Unpublished data  
b Data published in [20]  
c Data published in [54]  
d Data published in [55]  
e Data published in [26]  
f Data published in [27]  
g Pathways for propionic acid synthesis via succinate decarboxylation or amino acid degasation were not included
screened for encoded enzymes involved in homolactic and heterolactic acid fermentation. With the exception of the strain *Sporanaerobacter* sp. PP17-6a, all other bacterial genomes were predicted to perform homolactic acid fermentation. They harbor all genes encoding necessary enzymes including the gene for lactate dehydrogenase Ldh (EC: 1.1.1.27) converting pyruvate to lactic acid. Furthermore, some genetic determinants of the heterolactic acid fermentation pathway were identified. However, none of the strains encodes a full set of the genes needed. Hence, the question which strains are responsible for lactic acid production remains unsolved.

**Prediction of methanogenesis pathways based on sequence information for the subset of archaean genomes**

The formation of CH$_4$, the last step in the AD of biomass, is performed by methanogenic Archaea (Fig. 3). Based on their genetic repertoire, methanogens are able to perform either the hydrogenotrophic, acetoclastic, or methylotrophic pathway utilizing CO$_2$ and H$_2$, acetate, or methylamine and methanol, respectively, for CH$_4$ production [67]. To predict the pathway by which the analyzed Archaea produce CH$_4$, genes involved in the different methanogenesis pathways mentioned above
| Strain name                        | Features predicted | Methanobacterium formicicum MF<sup>T</sup> | Methanobacterium formicicum Mb9 | Methanobacterium sp. Mb1 | Methanobacterium conglobense Buetzberg | Methanothermobacter wolfeii SIV6 | Methanoculleus bourgensis MS2<sup>T</sup> | Methanoculleus chiku-goensis L21-II-0 |
|-----------------------------------|--------------------|------------------------------------------|---------------------------------|-------------------------|----------------------------------------|-----------------------------------|--------------------------------------------|------------------------------------------|
| Methanogenesis-related hydrogenase genes encoded in the genome | eha, ehb, fth, mvh, hdr | eha, ehb, fth, mvh, hdr | eha, ehb, fth, mvh, hdr | eha, ehb, fth, mvh, hdr | eha, ehb, fth, mvh, hdr | eha, ehb, fth, mvh, hdr | ech, fth, mvh, hdr | ech, fth, mvh, hdr |
| Substrates used for methanogenesis | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F | H<sub>2</sub>/CO<sub>2</sub>, F |
| Predicted metabolites required for growth | Acetate, cysteine<sup>a</sup>, vitamin B<sup>b</sup> | Acetate | Acetate | Acetate, lactate | Acetate | Acetate, lactate<sup>b</sup> | Acetate, lactate<sup>b</sup> |

F, formate; H<sub>2</sub>, hydrogen; CO<sub>2</sub>, carbon dioxide

<sup>a</sup> Utilization of cysteine and vitamin B by the strain MF<sup>T</sup> was described previously [50]

<sup>b</sup> No growth or methane production was detected on lactate for Methanoculleus species described previously [49, 82]
were examined interpreting functional KEGG assignments calculated within GenDB (Table 4).

All Archaea analyzed encode a full set of genes involved in CH₄ production from CO₂ and H₂. This result was as expected, as members of the families Methanobacteriaceae and Methanomicrobiaceae are known to solely perform hydrogenotrophic methanogenesis [68]. Additionally, genes for the formate dehydrogenase complex FdhA-B and a formate transporter FdhC for growth on formate as an alternative methanogenic substrate were identified in all seven analyzed genomes. For acetyl-CoA production from acetate, all seven genomes encode the acetyl-CoA synthetase Acs. Interestingly, methanogens from the genus Methanothermobacterium, namely the strains MS2T and L21-II-0, also harbor a lactate dehydrogenase gene involved in conversion of lactate to pyruvate or vice versa. However, no growth or CH₄ production from lactate has been described for the Methanothermobacterium species so far.

For activation of H₂ during methanogenesis, all seven Archaea analyzed encode the cytoplasmic coenzyme F₄₂₀-reducing [NiFe]-hydrogenases FrhA-D, the cytoplasmic [NiFe]-hydrogenase MvhADG, and the heterodisulfide reductase HdrABC in their genomes. The latter two enzyme complexes interact with the cytoplasmic [NiFe]-hydrogenase MvhADG, which was also identified in all investigated methanogens, for the coupled H₂-driven reduction of ferredoxin and heterodisulfide CoM-S-S-CoB [69]. Furthermore, methanogens of the family Methanobacteriaceae encode the membrane-bound energy-converting [NiFe]-hydrogenases EhaA-T and EhbA-Q [70], whereas the Methanomicrobiaceae strains encode the energy-converting [NiFe]-hydrogenase EcaA-F in their genomes. Members of the order Methanomicrobiales were described to exhibit a high affinity for H₂ (ca. 0.1 µM resp. 15 Pa H₂ pressure [71]), possibly providing an advantage over certain Methanobacteriales under conditions of low H₂ partial pressure.

Prevalence of bacterial and archaeal isolates in different microbial biogas communities analyzed by metagenome fragment mappings

To determine the prevalence or rather the abundance of the bacterial and archaeal isolates analyzed in this study in communities of production-scale BGPs, metagenome fragment mappings were done using deeply sequenced metagenomes from three mesophilic (BGP1-3) and one thermophilic (BGP4) agricultural BGPs which were published recently [41]. Configurations and process parameters corresponding to these BGPs are documented in the publication cited above. To identify metagenome sequence reads of the BGPs that match the genome sequences of the biogas isolates, these were mapped to the genomes applying Kallisto. Reads assigned to certain genomes were summed up and normalized according to dataset and genome sizes analogous to TPM (transcripts per million, [72]) values in RNASeq studies, to allow for quantitative comparisons.

Metagenome fragment mapping results were distinguished into the following groups: (I) abundant fully covered genomes, (II) less abundant but fully covered genomes, (III) rare but fully covered genomes, and (IV) rare, partially covered genomes (examples for each group are shown in Additional file 1).

Only three genomes, namely those of Methanothermobacterium bourgensis MS2T, D. tunisiensis L3, and Clostridium sp. N3C, fall into group I. M. bourgensis is abundant in all mesophilic BGPs studied and slightly less abundant in the thermophilic BGP, whereas D. tunisiensis and Clostridium sp. N3C are prominent in the thermophilic BGP (Fig. 4, Additional file 3).

Several of the analyzed strains were clearly detectable in the mesophilic BGPs but appeared to be only moderately abundant (group II). The strains H. luporum SD1T, M. chikugoensis L21-II-0, Sporanaerobacter sp. PP17-6a, and M. wolfei SI6 fall into this category. They are supposed to perform functions that are also taken by other community members. In other words, the corresponding microbial guilds are composed of several species featuring similar functionalities. Specific adaptation of species within a guild may refer to slight fluctuations in environmental conditions with one or the other species being more competitive under a particular condition.

The strains C. bornimense M2/40T, F. caenicola ING-E5B, H. hemicellulosilytica T3/55T, and C. thermocellum BC1 seem to be rare in most of the analyzed BGPs (group III), whereas the isolates Proteiniborus sp. DW1, Peptoniphilaceae bacterium str. ING-D1G, P. mucosa ING-E5A, Methanobacterium sp. Mb1, P. saccharofermentans M3/6T, B. thermoamylovorans 1A1, Propionispora sp. 2/2-37, M. formicicum MF7, M. formicicum Mb9, M. conglolense Buetzberg, and C. cellulosi DG5 seem to be, if at all, of minor importance in most BGPs (group IV).

Furthermore, the non-cultivable fractions of the biogas microbiomes residing in BGPs 1 to 4 were studied by Stolze et al. [41], applying metagenome assembly combined with a binning method. This approach enabled the identification of novel and uncharacterized species represented by MAGs, namely 206_Thermotogae, 175_Fusobacteria, 138_Spirochaetes, 244_Cloacimonetes, and 120_Cloacimonetes. To determine the prevalence of these MAGs in the biogas microbiomes analyzed, fragment recruitments were performed. The obtained results showed that the species represented by the bin 175_Fusobacteria is abundant in the mesophilic BGP3, whereas both Cloacimonetes MAGs were abundant in BGP2 and
Furthermore, all three MAGs represent fully covered genomes and therefore fall into the groups I and II in the case of 175_Fusobacteria and both Cloacimonetes MAG, respectively. The bin 138_Spirochaetes is detectable in the mesophilic BGP3 but appeared to be only moderately abundant (group III). The MAG 206_Thermotoga MAG was not further considered for fragment recruitments.

Among the publicly available reference species, only the genomes of M. bourgensis MAB1 [74] originating from a laboratory-scale biogas reactor and Amphicillus xylanus NBRC 15112 [75], isolated from compost of manure with grass and rice straw, were almost completely covered with metagenome sequences featuring high matching accuracy. The bacterial species A. xylanus NBRC 15112 was found to be highly abundant within the BGP1 microbiome, whereas the hydrogenotrophic methanogen M. bourgensis MAB1 was dominant in the mesophilic digesters 2 and 3 (Fig. 4). The genomes of both strains fall into group I regarding their fragment recruitments.
recruitment profiles. Among the microorganisms of group II, the species *Clostridium thermocellum* involved in hydrolysis of cellulose and hemicellulose [76] and *Streptococcus suis* BM407, a human pathogen [77], were found to be nearly fully covered but less abundant.

Based on these findings, metagenome fragment mappings clearly showed that the culturomics approach led to isolation and characterization of dominant and therefore important members of the biogas microbiome. However, since it is assumed that many biogas community members cannot be cultured by currently available cultivation techniques, further prevalent key microorganisms remain to be discovered.

**Conclusions**

Application of high-throughput and -omics technologies such as metagenomics, metatranscriptomics, metaproteomics, and genomics for the analysis of biogas microbial communities is becoming increasingly important. However, currently, the interpretation of generated data is limited due to the restricted availability of the corresponding and appropriate reference genome sequences connected with functional and metabolic information in public databases.

In this study, whole genome sequence information for 22 bacterial and archaeal strains was analyzed with respect to their metabolic functions in AD communities. For 15 bacterial strains, their participation in hydrolysis and/or acidogenesis/acetogenesis of plant biomass decomposition was predicted and partially verified by in vivo characterization of pure cultures. *Clostridium cellulolyticum* DG5, *H. hemicellulosilytica* T3/55T, *H. luporum* SD1D3, and *C. thermocellum* BC1 represent cellulose degraders, while the nine remaining bacteria presumably play a role in acidogenesis and/or acetogenesis. The seven analyzed methanogenic *Archaea* were predicted to produce CH₄ via the hydrogenotrophic pathway, representing the final phase of the AD chain.

Among the microorganisms analyzed in this study, only two species, namely *Methanoculleus bourgensis* and *D. tunisiensis*, were identified to play a dominant role within biogas microbial communities. *Defluviitoga tunisiensis* was proposed as a marker organism for the thermophilic biogas processes. This species is very versatile in the utilization of different sugars that can be converted to metabolites serving as substrates for methanogenesis. *Methanoculleus bourgensis* has frequently been found to dominate methanogenic sub-communities residing in production-scale BGPs and is assumed to be well adapted to high-osmolarity conditions and ammonia/ammonium concentrations prevailing when manure is used as a substrate for biogas production. Furthermore, the fragment recruitment analysis of MAGs published by Stolze et al. [41] could also show that in addition to the classical cultivation and isolation strategy, the metagenome assembly and binning approach may also enable the identification and characterization of previously unknown but abundant species featuring important functional potential in the context of the anaerobic digestion process.

It appeared that among the publicly available genomes only those of the species *A. xylanus*, *C. clariflavum*, and *C. thermocellum* were found to be well represented within biogas microbiomes, but do not reach the level of abundance as observed for *M. bourgensis* and *D. tunisiensis*. Surprisingly, among 5061 complete genome sequences archived in the public database NCBI, only those mentioned above seem to be of pronounced importance for agricultural biogas systems. Accordingly, the applied culturomics approach led to the isolation of further key AD species, thus providing genome sequence information for novel biogas community members. In the future, the non-cultivable fraction of AD communities should also be accessed by single-cell genomics to uncover genome sequence information of further, so far unknown biogas community members.

**Additional files**

Additional file 1. Fragment recruitment of metagenome sequences derived from four biogas-producing microorganisms to the genome sequences of the exemplarily chosen strains *Amphibacillus xylosus* NBRC 15112T, *C. cellulolyticum* N3C, *Fermentimonas caenicola* ING2-ESB3, *Methanobacterium formicicum* MF1 and *Methanoculleus bourgensis* MAB1. The x-axis: microbial genome analyzed, y-axis: percent identities of mapped metagenome reads.

Additional file 2. Genomic loci encoding enzymatic functions participating in the propionic acid, ethanol, formic acid, butyric acid and lactic acid fermentation for each strain analyzed.

Additional file 3. List of the 72 most abundant bacterial and archaeal strains within the biogas microbial communities analyzed, their GPM (genomes per million) values and further coverage statistics.

**Abbreviations**

AD: anaerobic digestion; BGP: biogas plant; CCC: circulary closed chromo-some; CAZymes: carbohydrate-active enzymes; CSTR: continuous stirred tank reactor; DSMZ: Leibniz Institute German Collection of Microorganisms and Cell Cultures; GH: glycosyl hydrolase; GPM: genomes per million; HT: high-through-put; KEGG: Kyoto Encyclopedia of Genes and Genomes; qPCR: quantitative 'real-time' polymerase chain reaction; TPM: transcripts per million; TRFLP: terminal restriction fragment length polymorphism; UASS: upflow anaerobic solid-state reactor; VFA: volatile fatty acids.

**Authors’ contributions**

IM performed the phylogenetic classification, genome assembly, and annotation of microbial isolates, participated in the prediction of bacterial fermentation pathways based on genome sequence information, coordinated drafting, and drafted the corresponding parts of the manuscript. AB carried out the fragment recruitment analyses for the 5061 publicly available genomes plus the 22 strains from this study, contributed to the "Results and discussion" section, and revised the manuscript. YS participated in the prediction of bacterial fermentation pathways based on genome sequence information and revised the manuscript. SH contributed to isolation and characterization of acidogenic bacterial strains and additional methanogenic Archaea and drafted the corresponding parts of the manuscript. KGC isolated and characterized acidogenic
bacterial strains and revised the manuscript. DEK isolated and characterized cellulolytic, hydrolytic, and acidogenic bacterial strains, participated in the analyses of bacterial genes encoding carbohydrate-active enzymes and revised the manuscript. YSK and JK contributed to the isolation of methanogenic Archaea and revised the manuscript. JH contributed to the phylogenetic classification of the analyzed bacterial and archaeal isolates and drafted the corresponding part of the manuscript. DW participated in the genome assembly and annotation of microbial isolates, submitted the 22 genome sequences to the EBI database, and revised the manuscript. AW participated in bioinformatic data analysis and revised the manuscript. SO participated in the isolation and characterization of methanogenic archaeal strains and contributed to the results and discussion part of the manuscript on archaeal isolates. RS participated in the isolation and characterization of methanogenic archaeal strains and contributed to the revision of the manuscript. VVZ and WL contributed to the discussion section and revised the manuscript. PS participated in the analysis of the hydrogenase genes in methanogenic archaeal isolates and revised the manuscript. ACM participated in bioinformatic data analysis and revised the manuscript. AScz participated in bioinformatic data analysis and discussion of bioinformatics results. MK participated in the design of this study, contributed to the "Results and discussion" section, and revised the manuscript. AP and AS conceived the study, participated in manuscript coordination, drafted the fragment recruitment section, supervised all biological analyses, and revised the manuscript. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

Availability of supporting data
The metagenome datasets supporting the conclusions of this article are available at the NCBI database in the short read archive (SRA): http://www.ncbi.nlm.nih.gov/sra/?term=SRA357211. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357213. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357208. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357209. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357214. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357221. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357222. http://www.ncbi.nlm.nih.gov/sra/?term=SRA357223.

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