Unveiling the ecology, taxonomy and metabolic capabilities of MBA03, a potential key player in anaerobic digestion

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Abstract

Biogas, a mix of CO₂, CH₄ and small proportions of other gases, is a biofuel obtained by anaerobic digestion (AD). Biogas production is often considered a black box process, as the role and dynamics of some of the microorganisms involved remain undisclosed. Previous metataxonomic studies in the frame of the MICRO4BIOGAS project (www.micro4biogas.eu) revealed that MBA03, an uncharacterised and uncultured bacterial taxon, was very prevalent and abundant in industrial full-scale AD plants. Surprisingly, no culturable specimen or genome of this taxon has ever been reported, so its role in AD has remained unclear. In the present work, thirty samples derived from anaerobic digesters were sequenced, allowing the reconstruction of 108 metagenome-assembled genomes (MAGs) potentially belonging to MBA03. According to phylogenetic analyses and genomic similarity indices, MBA03 constitutes a new bacterial order, proposed as Darwinibacterales ord. nov., which includes Darwinibacter acetoxidans gen. nov., sp. nov. of the family Darwinibacteriaceae fam. nov., along with Wallacebacter cryptica gen. nov., sp. nov. of the Wallacebacteriaceae fam. nov. Ecotaxonomic studies determined that AD processes are the main ecological niche of Darwinibacterales. Moreover, metabolic predictions identified Darwinibacteraceae members as putative syntrophic acetate oxidising bacteria (SAOB), as they encode for the reversed Wood-Ljungdahl pathway coupled to the glycine cleavage system. This suggests that Darwinibacteraceae members work in collaboration with hydrogenotrophic archaea to produce methane in industrial biogas plants. Overall, our findings present Darwinibacterales as a potential key player in anaerobic digestion and pave the way towards the complete characterisation of this newly described bacterial taxa.

Keywords: Metagenomics, anaerobic digestion, biogas, SAOB, MBA03, Darwinibacterales
The evidence supporting the need for a transition from fossil to renewable energy is overwhelming and unanimous (Yaqoob et al., 2021). In recent years, biogas has arisen as one of the leading alternatives to non-renewable fossil fuels such as natural gas. Biogas is produced by anaerobic digestion (AD) of organic material, through a complex microbiological process in which several groups of microorganisms with diverse metabolic abilities work in close collaboration to obtain methane as the main product (Schnürer, 2016). In addition to the production of renewable methane and fertiliser, biogas production contributes to the utilisation of today's global waste streams of various origins as a source of organic matter. In particular, waste from agriculture, the food industry, wastewater and municipal solid waste can be treated. (Tsang et al., 2019).

Biogas primarily consists of methane (CH₄), carbon dioxide (CO₂) and other gases like H₂S, N₂, NH₃, and H₂ in lower concentrations (Yaqoob et al., 2021). Despite the complexity of the AD process, the microbial metabolic reactions that occur in this process and that lead to the synthesis of methane can be summarized in four key steps: hydrolysis, acidogenesis, acetogenesis and methanogenesis (Schnürer, 2016). Briefly, during hydrolysis, complex macromolecules are broken down into various oligo-, di- and monomers, the final concentrations of which will depend on the biomass source (Nwokolo et al., 2020). These molecules are uptaken and used by acidogenic bacteria to produce H₂, CO₂, alcohols and volatile fatty acids (VFAs) during acidogenesis, while in acetogenesis, acetogenic bacteria synthesize acetic acid through the Wood-Ljungdahl (W-L) pathway or autotrophically from H₂ and CO₂. Subsequently, methanogens metabolize acetate, hydrogen and/or methylated compounds to produce methane, creating an equilibrium and a co-dependence between bacteria and archaea (Nwokolo et al., 2020; Vanwonterghem et al., 2016). Two main pathways can occur in the production of methane from acetate: the first one, directly by acetoclastic methanogens, which split acetate to methane and CO₂, with the methyl group yielding methane and the carboxyl group yielding CO₂. The other pathway results from syntrophic acetate-oxidizing bacteria (SAOB) oxidizing acetate to CO₂ and H₂, which is subsequently used by hydrogenotrophic archaea to obtain methane. Thus, this pathway must be coupled with hydrogenotrophic methanogenesis (SAO-HM). Although less abundant in anaerobic digester plants, there is a third pathway, which is the methylotrophic pathway (Abendroth et al., 2015). This pathway is regarded as an ancestral pathway of methanogenesis (Wang et al., 2001), where the respective archaea are converting methyllic C1 groups (Whitman et al., 2006).

The microbial community underlying AD shows very complex entangled ecologies, with the taxonomic composition additionally depending on many factors such as substrate, temperature or pH (De Vrieze et al., 2015; Lebuhn et al., 2015).

Despite the high economic potential, knowledge for the production of biogas with AD has remained stagnant, as the high variability of both the different operational parameters and the native microbial communities involved represent a challenge for its optimization and standardization. Improving the feasibility of the process and achieving a consistent and controlled way to obtain biogas...
would constitute a huge change in the energy industry. But reaching such an ambitious goal requires gaining insights in the microbial machinery behind AD and methanogenesis, and this goes through the study of a variety of anaerobic digestion systems with different chemical, taxonomical and operational parameters.

In this context, the EU-funded project MICRO4BIOGAS (https://micro4biogas.eu) was born in 2021. The ultimate objective of the project is to improve the yield, speed, quality and robustness of biogas production by optimising the AD microbiome. In the frame of the project, 80 different samples were collected from various AD plants in the Netherlands, Germany and Austria.

After metataxonomic analyses, a miscellaneous taxonomic group (i.e., MBA03) was found to be very prevalent and abundant. Moreover, MBA03 was among the taxa that correlated most strongly with operational (i.e., temperature) and chemical parameters (i.e., acetic acid) (Otto et al., 2023), suggesting that it may be a microorganism of great relevance for the AD process. MBA03 (AB114313.1) was described for the first time by Tang et al. as an uncultured clone obtained from a thermophilic anaerobic municipal solid-waste digester (Tang et al., 2004). The proposal of MBA03 exclusively relied on the 16S rRNA gene sequence. Since then, a series of metataxonomic studies have reported the presence of MBA03 in anaerobic digester systems (Dyksma et al., 2020; FitzGerald, 2018; Laguillaumie et al., 2022), representing up to 70% of the relative abundance in some samples (FitzGerald, 2018). This taxon has been associated with a higher production of methane and has been suggested as an indicator of the stability of the AD process (Fang et al., 2022; Laguillaumie et al., 2022).

Surprisingly, MBA03 has not been properly taxonomically delimited. In fact, no culturable specimen or genome of this taxon has ever been reported. However, according to SILVA nr database, MBA03 represents a heterogeneous group of more than 150 16S rRNA sequences. Therefore, a deeper characterization of this taxonomic group was urgently needed for unravelling its role in AD processes.

In this work, we used metagenomic data from MICRO4BIOGAS and other AD studies to propose MBA03 as a member Darwinibacteriales ord. nov., a not-yet-cultured bacterial taxa belonging to the class Clostridia. Darwinibacteriales includes two different genera: Darwinibacter acetoxidans gen. nov., sp. nov., of the Darwinibacteriaceae family, and Wallacebacter cryptica gen. nov., sp. nov., of the Wallacebacteriaceae fam. nov. Based on our results, both Darwinibacter and Wallacebacter predominantly inhabit AD environments. Furthermore, we have inferred that Darwinibacter may play a key role in anaerobic digestion processes related to biogas production, as it includes the reversed Wood-Ljungdahl pathway coupled to the glycine cleavage system, which is required for the synthesis of H₂ and CO₂ from acetate and the subsequent production of methane by hydrogenotrophic methanogenesis.
2. Materials and methods

2.1. Sample description

As part of the MICRO4BIOGAS project, a total of 80 samples were collected from anaerobic digestion systems at 45 large-scale reactors with different operational conditions and feedstocks. These samples, together with the corresponding metadata, were used to investigate the chemical and taxonomic profiles (Otto et al., 2023). The first 40 samples were collected by members of the Technische Universität Dresden (TUD, Dresden, Germany) from 25 different biogas plants in Germany and Austria, while the second set of 40 samples was collected by the team members from Bioclear Earth B.V. (Groningen, Netherlands) from 20 different biogas plants in the Netherlands. A subset of 30 of these samples was selected to be analysed through metagenomic sequencing. This subset included samples enriched in MBA03, hydrolytic, acidogenic and acetogenic bacteria and/or methanogenic archaea. A full description of the samples can be found in Supplementary Table 1.

2.2. DNA extraction and sequencing

The sludge samples were washed before DNA extraction to reduce the number of inhibiting substances. Specifically, 5-10 mL of each sample was mixed with 40 mL of sterile phosphate-buffered saline (PBS). Subsequently, the samples were centrifuged at 20,000 g for 10 min and the supernatants were discarded. This process was repeated twice, and the final pellets were used for DNA extraction with the DNeasy® PowerSoil® Pro kit (QIAGEN, Germany) based on the manufacturer’s instructions, with an extra incubation step at 65 °C for 10 min after adding the C1 solution to increase the efficiency of cell lysis. Qubit x1 dsDNA HS Assay kit (Qubit 4.0 Fluorometer, Thermo Fisher) was used for DNA quantification. The metagenomic DNA was randomly fragmented by sonication and DNA fragments were end polished, A-tailed, and ligated with the full-length adapters for Illumina sequencing: 5'-AGA TCG GAA GAG CGT CGT GTA GGG AAA GAG TGT AGA TCT CGG TGG TCG CCG TAT CAT T -3' (5' Adapter), and 5'- GAT CGG AAG AGC ACA CGT CTG AAC TCC AGT CAC GGA TGA CTA TCT CGT ATG CCG TCT TCT GCT GT -3' (3' adapter). PCR amplification with P5 and indexed P7 oligos was carried out, and the PCR products were purified with the AMPure XP system. Then, the libraries were checked for size distribution by an Agilent 2100 Bioanalyzer (Agilent Technologies, CA, USA), and quantified by real-time PCR. Libraries were sequenced at Novogene (UK) using the NovaSeq 6000 Illumina platform (150 bp x 2). The average sequencing depth was 45M reads/sample (min.: 35M reads; max.: 55M reads).

2.3. Metagenomic analysis

Adapters were trimmed from the raw reads using Cutadapt (v. 3.4; Martin, 2011), and a quality filtering step was performed with BBDDuk (BBTools v.38.84; https://sourceforge.net/projects/bbttools/). Reads shorter than 75 bp or with a
mean quality lower than Q20 (in the PHRED scale) were discarded. Bases with a quality below Q20 were also trimmed from both read ends. Human reads were then removed with bowtie2 (v. 2.4.4; Langmead & Salzberg, 2012) by aligning the reads against the human genome (GRCh38). The quality of the clean sequences was checked with FastQC (v0.11.5; http://www.bioinformatics.babraham.ac.uk/projects/fastqc). MEGAHIT was used with default parameters for assembling the filtered metagenomic reads (v1.2.9; (D. Li et al., 2015). Assembly statistics were calculated with QUAST (v. 5.0.2; (Gurevich et al., 2013). Two different binning strategies were applied: MetaBAT (Kang et al., 2019) and MaxBin (Wu et al., 2016). The MAGs obtained were refined with Das Tool (Sieber et al., 2018) and classified according to their quality: high quality (HQ, integrity ≥ 90% and contamination ≤ 5%), good quality (GQ, integrity ≥ 80% and contamination ≤ 10%) and medium-low quality (LQ, integrity ≤ 80% and/or contamination ≥ 10%), following the criteria of Bowers et al., 2017 and Feng et al., 2021. After that, Centrifuge (Kim et al., 2016) was used to study the taxonomic profile of the samples, while the resulting MAGs were taxonomically annotated with MiGA (Rodriguez-R et al., 2018).

2.4. Recovery of MBA03 genomes via phylogenetic analyses

First of all, 16S rRNA sequences were extracted from the high-quality MAGs using Prokka (v 1.14.6) (Seemann, 2014). These sequences were aligned together with the 157 available MBA03 16S rRNA sequences from SILVA nr database (SSU 138.1 ref NR) with the SINA Aligner (v 1.2.11) (Pruesse et al., 2012), and afterwards a phylogenetic tree was computed with FastTree (v 2.1.1) (Price et al., 2009) and visualized with the Interactive Tree Of Life (iTOL) (Letunic & Bork, 2021). Only the MAGs whose sequences formed a monophyletic group with MBA03 were selected for further analysis. Due to the limitations associated to the binning process, not all MAGs contained 16S rRNA sequences, while others contained more than one sequence, so not all the MAGs corresponding to MBA03 could be identified by 16S rRNA.

2.5. MBA03 identification from phylogenomic data

After identifying the first MAGs hypothetically belonging to MBA03, a two-step approach was followed to recover the remaining MAGs belonging to the MBA03 taxonomic group which did not contain rRNA operons.

As a first step, all the MAGs whose 16S rRNA gene formed a monophyletic group with MBA03 according to the phylogenetic tree were annotated using the whole genome classifier tool Genome Taxonomy Database toolkit (GTDB-tk; v 2.1.1) (Chaumeil et al., 2022). Although GTDB does not contain any genome identified as “MBA03”, the resulting annotation was used as a first hypothesis to elucidate which of the genomes actually corresponded to MBA03.

As a second step, all the HQ and GQ MAGs recovered in the project (n = 1053) were also annotated GTDB-tk. In addition, a phylogenomic tree was constructed using the UBCG pipeline (v 3.0) (Na et al., 2018). Once all the MAGs corresponding to MBA03 were identified, genomes belonging to the MBA03 order...
in the GTDB were downloaded and included in the dataset. Afterwards, genomes from the Biogasmicrobiome repository (https://biogasmicrobiome.env.dtu.dk/), and from (Dyksma et al., 2020) were annotated and incorporated into the dataset.

2.6. Taxonomic delimitation of the MBA03 taxonomic clade

Prokka (v 1.14.6) (Seemann, 2014) was used again to obtain all the 16S rRNA sequences from the MAGs and genomes identified as potential MBA03 representatives according to the previous phylogenomic analyses. To better study the phylogenetic patterns, 16S rRNA sequences corresponding to the most closely related taxonomic classes were set as outgroups. Sequences from class Clostridia and Limnochordia were downloaded from the Living Tree Project (LTP) and dereplicated using cd-hit (4.8.1) (W. Li & Godzik, 2006) at a 0.97 similarity threshold. Sequences were aligned with SINA Aligner (v 1.2.11) (Pruesse et al., 2012) and incorporated into the previous dataset. Finally, a phylogenetic tree was computed with FastTree (Price et al., 2009), and visualized with iTOL (Letunic & Bork, 2021). The reference sequence of the bacterial clone identified for the first time as MBA03 was also downloaded from the NCBI (accession number: AB114313.1) and included in the tree.

Following the same procedure as for the phylogenetic tree, genomes from Clostridia and Limnochordia were downloaded from RefSeq NCBI Assembly Database, filtering by “Assembly from Type Material” and “Complete Genomes” (Supplementary Table 2). These genomes were analysed altogether with the genomes identified as MBA03 and a phylogeny of clade was created with UBCG (Na et al., 2018) and visualized with iTOL (Letunic & Bork, 2021).

2.7. Analysis of the ecological niche of MBA03

In order to explore the ecological niches of MBA03, an analysis with IMNGS (Lagkouvardos et al., 2016) was carried out using a 97% identity cut-off. The same sequences used for the phylogenomic analysis were dereplicated using cd-hit (v 4.8.1) (W. Li & Godzik, 2006) at a 0.9 identity threshold. IMNGS results were filtered with the objective of gathering all the samples by their sample type to see which are the potentially relevant ecological niches of MBA03. As tags describing ecological niches were very diverse and, in many cases, ambiguous, a series of in-house Python scripts were needed to carry out a reclassification of niche categories. Prior to this filtering, samples containing the 16S rRNA sequence of interest in an abundance <0.1% were removed from the analyses. To reclassify the remaining samples, two subsequent strategies were followed: (1) description tags were filtered by the words “anaerobic”, “digester”, “biogas”, and “reactor”, assigning all of them to the same tag: “anaerobic digestion”; (2) the information of the remaining samples was scraped from NCBI. The output was manually curated to determine which samples belonged to “anaerobic digestion”, and other general tags were also reassigned to more specific classes (i.e., “metagenome” to “soil metagenome”) depending on the project description.
In order to determine the abundance of MBA03 among the samples, coverM (https://github.com/wwood/CoverM) was used, comparing the raw reads of the 30 sequenced reactors against the genomes assigned to MBA03.

2.8. Functional annotation

MAGs belonging to MBA03 were filtered according to two criteria: completeness > 90% and contamination < 5%. These MAGs were annotated with Bakta (Schwengers et al., 2021) and they were analysed with Kofamsan (Aramaki et al., 2020) and KEGG-Decoder (Graham et al., 2018).

As mentioned above, acetate as an intermediate is of particular importance during anaerobic reactions, especially for SAOB bacteria and for hydrogenotrophic methanogenesis. In order to elucidate if MBA03 was a SAOB, the protein sequences of all the enzymes involved in the reversed Wood-Ljungdahl (WL) pathway and the reversed WL pathway coupled to the glycine cleavage system (GCS) (Supplementary Table 3) were downloaded from UniProt (Coudert et al., 2023) (Swissprot), narrowing to all bacteria (taxonomy_id:2), and a database was created with Diamond (Buchfink et al., 2015). A Diamond BLASTp of the annotated protein sequences was performed against this database to detect presence and absence of each gene in all MAGs with 50% coverage and identity thresholds. For the whole metabolic analysis, any enzyme found in more than 70% of the MAGs of each family was considered to be present in the genome.

3. Results and discussion

3.1. Genomic reconstruction

MBA03 has proven abundant in a variety of anaerobic digesters (Dyksma et al., 2020; Liu et al., 2023). Although there are more than 150 16S rRNA sequences classified as MBA03 in the SILVA nr database (v. 138), no representative genome of any bacteria belonging to this taxon has ever been described. Given its importance in AD environments, this work focused on the recovery and characterisation of the genome of this taxonomic group using 30 metagenomic datasets derived from the MICRO4BIOGAS project and other genomic data available in public databases.

After metagenomic sequencing, assembly and binning of sludge samples, the metagenome-assembled genomes (MAGs) obtained were classified according to their quality, and 1053 MAGs (600 of high quality (HQ) and 453 of good quality (GQ)) were chosen for further analysis. After the annotation of the MAGs with MIGA and the metagenomes with Centrifuge, in neither case, MBA03 was reported despite being a highly abundant taxon according to the metataxonomic results (Otto et al., 2023). This was expected as genomic databases did not include MBA03 genomes, probably reflecting a lack of consensus on the nomenclature of this taxonomic group.
3.2. Taxonomic analysis of MAGs

In order to identify the genomes belonging to MBA03, a phylogenetic tree was constructed using the 16S rRNA sequences extracted from the HQ MAGs and 157 sequences classified as MBA03 according to the SILVA database (SSU 138.1 ref NR) (Supplementary Figure 1). The sequences from SILVA formed a monophyletic clade, and only 17 sequences from the metagenomic dataset fell within this subtree. The MAGs corresponding to these 17 sequences were annotated with GTDB-tk (v 2.1.1) and 13 out of the 17 MAGs shared the same annotation: they belonged to phylum Bacillota, class Limnochordia, order DTU010. From here, two different families were identified: DTU010 and DTU012 (Supplementary Table 4).

After this preliminary analysis, the HQ and GQ MAGs (n=1053) were annotated with GTDB-tk (Supplementary Table 5). A total of 45 MAGs were classified as members of the order DTU010, and they were subsequently treated as potential representatives of MBA03. The phylogenomic tree of HQ and GQ MAGs (Figure 1) also revealed a large proportion of MAGs belonging to Bacillota (n = 503), while 104 MAGs were classified as Bacteroidota, 77 as Actinobacteriota, 63 as Verrucomicrobiota and 32 as Chloroflexota, while a total of 80 MAGs corresponded to Archaea. These results were expected considering previous studies reporting the average AD microbiome (Sundberg et al., 2013; Abendroth et al., 2015; Treu et al., 2016; Kirkegaard et al., 2017; Otto et al., 2023). Moreover, it was confirmed that MBA03 formed a monophyletic clade with all 45 sequences annotated as DTU010. Two different subclades could be clearly distinguished, corresponding to families DTU010 and DTU012 according to the GTDB taxonomy.

**Figure 1.** (A) Dendrogram of all the HQ and GQ MAGs from MICRO4BIOGAS (n=1053), computed with UBCG and visualized with iTOL. The legend shows a classification by different colours corresponding to different phyla, according to the annotation obtained with GTDB-tk (v 2.1.1), and a highlight on MBA03 (within Bacillota). (B) Tree with 108 leaves, computed with FastTree and visualized with iTOL, corresponding to the MAGs belonging to order DTU010 according to GTDB-tk (v 2.1.1). Two clades corresponding to two families (DTU010 in orange and DTU012 in yellow) can be seen, for which the names
Darwinibacteriaceae and Wallaceae are proposed, respectively. In both figures, bootstrap values greater than 0.75 are represented by a small triangle in each branch.

For the enlargement of the dataset, 1804 MAGs from the Biogasmicrobiome repository (https://biogasmicrobiome.env.dtu.dk/) and 16 from Dyksma et al. were downloaded and annotated with GTDB-tk (Dyksma et al., 2016). After filtering, a total of 64 potential MBA03 MAGs were compiled. Together with the previous 45 MAGs (HQ and GQ) retrieved from MICRO4BIOGAS data, the final dataset was composed of 108 genomes. These MAGs were used for creating a phylogenomic tree (Figure 2), leading to an initial taxonomic delimitation of MBA03: (i) there were two families, named DTU010 (which accounted for 82 MAGs) and DTU012 (with 26 MAGs) according to GTDB database; (ii) DTU010 clade included 20 different taxa annotated in GTDB. The most common was DTU010 sp. 002391385, followed by sp. 900018335, and both formed monophyletic clades; and (iii) there were 10 different taxa belonging to DTU012, with the most common species being DTU012 sp.900019385.

Thereby, it was proposed the new order to be named Darwinibacteriales ord. Nov., and the two families Darwinibacteriaceae fam. nov., and Wallacebacteriaceae fam. nov., after Charles Darwin and Alfred Wallace, who are considered the fathers of the Theory of Evolution by natural selection.

### 3.3. Final taxonomic delimitation of MBA03

According to these results, the order Darwinibacteriales ord. nov. (DTU010 according to GTDB) was hypothesized to be equivalent to MBA03 (SILVA’s name), but further phylogenetic evidence was needed to get a proper delimitation of the clade. For that reason, phylogenetic analyses based on the 16S rRNA gene were carried out. A total of 52 16S rRNA sequences were identified by Prokka (Seemann, 2014) in the 108 MAGs (i.e., not all MAGs contained 16S rRNA sequences), and they were computed together with the 157 16S rRNA sequences downloaded from the SILVA (SSU 138.1 ref NR) database in a phylogenetic tree (n=209). Moreover, 772 sequences from Clostridia were added as outgroups.

The phylogenetic tree (Figure 2) confirmed the abovementioned hypothesis: the sequences annotated as MBA03 in SILVA formed a monophyletic clade with a clear subdivision of two clades, which included two distinct families (Darwinibacteraceae/DTU010 and Wallacebacteriaceae/DTU012). Clade I was formed by 52 sequences and included the MBA03 reference sequence, whereas Clade II accounted for 44 sequences.
Figure 2. Phylogenetic tree based on 16S rRNA sequence from our own database and from SILVA ref NR SSU 138.1 database. The sequences in bold letter belong to the MAGs from our database, while the other sequences are the ones downloaded from SILVA. Coloured clades correspond to the outgroup sequences, and they have been collapsed for a better visualization of the MBA03 clade.

It is worth highlighting that, although both *Hydrogenispora* and *Darwinibacteriales* (MBA03) are annotated as class *Limnochordia* in the SILVA database, *Hydrogenispora* is annotated as *Clostridia* in the LPSN (List of Prokaryotic names with Standing in Nomenclature, Parte et al., 2020), which provides the internationally accepted and regulated taxonomic classification. This annotation, supported by other authors (FitzGerald, 2018; Westerholm et al., 2016), was confirmed once the tree was computed, as both *Hydrogenispora* and *Darwinibacteriales* appeared to be two different orders from class *Clostridia*. In other words; *Darwinibacteriales* is proposed as a new order that belongs to class *Clostridia*.

Finally, a phylogenomic tree (Figure 3) was computed with the 108 genomes classified as MBA03 together with 206 genomes from *Clostridia* and one genome from *Limnochordia* as outgroups (Supplementary table 2). This confirmed the hypothesis that MBA03 belongs to a new order within class *Clostridia*, and it comprises two families.
Figure 3. Phylogenomic tree showing only the *Darwinibacteriales* (MBA03 in SILVA) clade, which includes two potential new families: *Darwinibacteriaceae* (DTU010) shaded in pink and *Wallacebacteriaceae* (DTU012) shaded in blue. Output groups from *Clostridia* and one genome from *Limnochordia* are collapsed in the tree for a better visualization. MAGs in green (n=9) belong to *Darwinibacteriales* (MBA03), as there is evidence at the genetic and genomic level. MAGs in red (n=9) may be MBA03, but their 16S rRNA phylogeny is not concordant. This could be explained by errors during the binning process. The remaining MAGs (n=90) are incomplete and/or 16S rRNA sequences could not recovered from the MAGs. Although they form a cluster with the rest of the MBA03 MAGs, they should remain as hypothetical members of this order.

According to these results, there is enough evidence to confirm that MBA03 belongs to a new taxonomical order, *Darwinibacteriales* ord. nov., which includes two potential new families, *Darwinibacteriaceae* fam. nov., (DTU010 according to GTDB) and *Wallacebacteriaceae* fam. nov., (former DTU012). It is important to note that samples from MICRO4BIOGAS provided the higher number of MAGs belonging to MBA03, meaning that the samples and datasets from this project...
marked a significant breakthrough towards the genetic isolation of MBA03 genomes. The formal description of these taxa can be found in Supplementary Material.

Once the two families from the order *Darwinibacteriales* (*Darwinibacteriaceae* and *Wallaceae*) were properly delimited, the read coverage of both families in the samples was calculated. 10 MAGs from DTU010 and 4 MAGs from DTU012 were used as reference for read mapping. The results showed that DTU010, with a mean abundance of 4.1% and a maximum abundance of near 18%, was considerably more abundant than DTU012, with a mean abundance of 0.6% (Supplementary Table 6).

### 3.5. Ecological distribution *Darwinibacteriales*

The ecological distribution of *Darwinibacteriales* was evaluated using IMNGS. This tool quantifies the prevalence and abundance of any 16S rRNA sequence of interest introduced as input, giving information about the number of coincidences with the samples included in its database. IMNGS also informs about the type of sample (e.g., soil, water, anaerobic digester, environmental) showing a hit with the input sequence. A total of 8856 SRA accessions containing 16S rRNA sequences from *Darwinibacteriales* were detected according to sequence similarity. Due to the high amount of data obtained, only the samples with an abundance of *Darwinibacteriales* higher than 0.1% were considered at first (Figure 4A). An exhaustive analysis of the sample metadata revealed that more than 99% of them belonged to anaerobic digestion studies, meaning that *Darwinibacteriales* is a very specialized bacterial group that grows and develops in this specific environment. This result reinforces the hypothesis that MBA03 *Darwinibacteriales* may play a central role in the AD process.

![Figure 4](image)

**Figure 4.** (A) Ecological niches of *Darwinibacteriaceae* based on coincidences of 16S rRNA sequences with samples of the IMNGS database with an abundance above 0.1%, (B) Ecological niches of *Darwinibacteriaceae* according to the samples with an abundance below 0.1% and excluding all the samples related to anaerobic digestion for a better insight into the wide range of environments from where it may be introduced to anaerobic reactors.
To study the ecological niches besides anaerobic digestion, a similar analysis was performed considering only the samples with an abundance of *Darwinibacteriales*-related sequences below 0.1%. The resulting 8650 hits were then filtered in two steps: (i) first, those related to anaerobic digestion were discarded, and (ii) results with less than 5 hits in total were removed from the dataset, leaving 1636 samples that were categorised into different groups (Figure 4B).

Soil was the second most common environment in which MBA03 could be found, followed by rhizosphere and sediment samples. However, the abundance of *Darwinibacteriales* in these habitats was substantially lower than in the case of anaerobic digesters. Therefore, it can be hypothesized that these bacteria are found in soil, plants and grounds, in a quiescent or inactive form. When agricultural residues, soils, muds or sludges are taken as substrate for anaerobic digestion, bacteria from the *Darwinibacteriales* order thrive in the anaerobic digester environment. Similarly, the presence of MBA03 in gut and milk microbiomes can be explained by the dietary habits of the animals. As *Darwinibacteriales* exists in the soil, herbivorous and omnivorous animals may ingest this group of bacteria during their feeding. Another hypothesis would be that *Darwinibacteriales* is part of the native gut microbiota of the animals, but at a very low abundance, and would thus be a reservoir which would guarantee the prevalence in AD plants fed with those substrates. Therefore, when manure is used as a substrate, *Darwinibacteriales* would be introduced into the bioreactors.

### 3.6. Unveiling the functional potential of Darwinibacteriales

In anaerobic digesters, the metabolic diversity of *Darwinibacteriales* remains challenging due to the absence of known cultivable representatives. A total of 45 and 16 MAGs from *Darwinibacteriaceae* (former DTU010) and 16 from *Wallaceae* (former DTU012), respectively, were analysed to gain insights into the potential metabolic pathways of this bacterial order (Supplementary Table 7).

*Darwinibacteriales* are chemoorganoheterotrophs that derive energy from anaerobic respiration through oxidative phosphorylation. They utilize inorganic molecules of ferric iron (Fe3+) as potential electron acceptors (K02012 and K02011). These bacteria also possess sulphide:quinone oxidoreductase (EC:1.8.5.4), an enzyme involved in the oxidation of sulphides, including hydrogen sulphide, enabling them to be chemolithotrophic. Their ability to grow on H2S is interesting, as H2S is usually not oxidized under anaerobic conditions. However, there are some exceptions. For example, it has recently been suggested to add reducing agents to control the H2S level. In this regard, Jung et al. (2020) highlighted the possibility to add magnetite. Jung et al. suggested that the phenomenon of direct interspecies electron transfer (DIET) can not only increase methanation activity, but can also reduce H2S, which results in an intracellular accumulation of S0. In the present study, no hints for DIET have been found so far. However, even in the case that DIET is not involved, it is conceivable that electrons are transferred via alternative routes via the
mediated electron transfer (MIET). Due to the importance of sulfur-metabolism in regard to industrial biogas formation, it would be of high interest, to study the involvement of Darwinibacteriales in the sulfur-metabolism in further detail. Apart from using anoxic electron donors, there is also the possibility to treat H$_2$S due to micro-aeration in anaerobic digesters (van der Zee et al., 2007). Because of this, plant operators inject sometimes small amounts of oxygen into anaerobic digesters. In turn, this enables microbial oxidation of H$_2$S to sulfate, which improves the overall biogas quality. The link to sulfur metabolism is particularly interesting because some members of Darwinibacteriales appear to be syntrophic microorganisms (as discussed later in 3.7). In fact, in recent years and alongside Jung et al., several other articles have been examining the influence between syntrophy and sulfur metabolism. To give here few examples: Igarashi and Kuwabara (2016) applied a syntrophic community to dacite pumice to induce oxygen supported H$_2$S oxidation. Shakeri Yekta et al. (2017) highlighted the complex interplay between iron, sulfur metabolism, volatile fatty acid turnover and syntrophic relation.

Darwinibacteriaceae and Wallacebacteriaceae have complete sets of genes for glycolysis (Embden-Meyerhof pathway; M00001), gluconeogenesis (M00003), and pyruvate oxidation (M00307) pathways. However, they lack certain essential genes required for the tricarboxylic acid (TCA) cycle, specifically isocitrate dehydrogenase. Both bacterial families can convert glycogen to glucose-6P. Most MAGs from both families contain genes encoding beta-N-acetylhexosaminidase for bacterial cell wall peptidoglycan degradation and recycling, pullulanase for starch degradation, and beta-glucosidase for cellobiose metabolism. Darwinibacteriaceae members also possess additional genes encoding chitinase for chitin degradation, D-galacturonate isomerase for pectin utilization, and alpha-amyrase for starch hydrolysis. Wallacebacteriaceae members carry the gene encoding oligogalacturonide lyase, allowing them to degrade pectin. Both Wallacebacteriaceae and Darwinibacteriaceae exhibit the ability to degrade catechol and methylcatechol through catechol 2,3-dioxygenase enzymes (EC 1.13.11.2). However, the presence of these compounds can be toxic to microorganisms in anaerobic digesters, inhibiting the digestion process. These findings suggest that each bacterial family can exploit different organic substrates as carbon and energy sources, maybe facilitating their coexistence. This also indicates that the importance of MB03 does not only restrict to one single phase of anaerobic digestion. Apart from acetogenesis and syntrophic acetate oxidation, MB03 could also be involved in the hydrolysis phase as well.

Members of Darwinibacteriales assimilate nitrogen by converting NH$_3$ to organic nitrogen through the enzymatic activities of glutamine synthetase (EC 6.3.1.2) and glutamate (NADPH) synthase (EC 1.4.1.13). However, genes associated with other nitrogen metabolism processes, such as nitrate reduction, denitrification, nitrogen fixation, nitrification, and anaerobic ammoniumoxidation, were not observed. Most representatives of Darwinibacteriales have auxotrophic
requirements for various amino acids, while *Wallacebacteriaceae* representatives are auxotrophic for a smaller set of amino acids (Supplementary Table 7).

Both bacterial families share several ABC transporters, including ABCB-BACATP binding cassettes and NitT/TauT family transporters. They also possess multiple sugar transport systems. *Darwinibacteriaceae* exhibits ABC transporters for iron (III), phosphonate, alpha-glucoside, aldouronate, sn-glycerol 3-phosphate, cobalamin, and haemolysin transport systems. Both *Darwinibacteriaceae* and *Wallacebacteriaceae* show transport systems for maltose/maltodextrin, arabinogalactan/maltooligosaccharide, raffinose/stachyose/melibiose, glucose/mannose, trehalose/maltose, cellobiose, N,N'-diacetylchitobiose, L-arabinose/D-xylene, rhamnose, ribose, fructooligosaccharides, sorbitol/mannitol, arabinooligosaccharide, polygalacturonan/rhamnogalacturonan, pectin-derived oligosaccharide, cobalt, spermidine/putrescine, nickel, and biotin transport systems.

Members of *Darwinibacteriales* possess resistance genes against vancomycin (vanW) and tetracycline (tetM and tetO). They also encode multidrug efflux pumps (mdlA/smdA; mdlB/smdB; efrA/efrE) to withstand various antimicrobial compounds.

Both families are mobile, as indicated by the presence of flagella synthesis genes. However, *Darwinibacteriaceae* lacks genes for the MS/C ring type III secretion system. Members of both taxa possess the enzymatic machinery for chemotactic movement and detection and transport of D-ribose and D-galactose via ABC transporters.

### 3.7 Members of *Darwinibacteriaceae* as potential SAOB

Members of the MBA03 lineage were previously hypothesised to be SAOB in AD based on correlation studies (Perman et al., 2022; Zheng et al., 2019; Zeng et al., 2021) together with other unclassified *Clostridia* (Dyksma et al., 2020); however, the metabolism of this bacterial lineage, classified in this study as *Darwinibacteriaceae*, remains poorly understood and characterised due to the lack of cultured representatives. SAO bacteria possess either the known reversed Wood-Ljungdahl (WL) pathway or an alternative route coupled to a glycine cleavage system (GCS) (Zeng et al., 2021). Recently, the co-occurrence of members of MBA03 with other potential SAO bacteria, such as DTU014 and *Syntrophaceticus*, and with hydrogenotrophic methanogens such as *Methanothermobacter, Methanoculleus* and *Methanosarcina*, has been observed (Otto et al., 2023). This suggests that several taxonomic groups identified as potential SAOBs, including *Darwinibacteriaceae*, grow simultaneously with methanogens to provide a continuous supply of H₂ and CO₂ for hydrogenotrophic methanogenesis. Usually, hydrogenotrophic methanogens are associated with high performance plants, whereas the acetoclastic methanogens *Methanothrix* (formally *Methanosaeta*) are found specifically in low performance systems such as sewage sludge digesters (Sunderberg et al. 2013; Abendroth et al., 2015). Therefore, due to the co-occurrence between members of *Darwinibacteriaceae* and hydrogenotrophic methanogens, we hypothesise that the presence of
Darwinibacteriaceae could be associated with a higher biogas production in industrial AD plants, although further analyses are required.

To confirm whether members of the Darwinibacteriaceae and Wallacebacteriaceae are SAOBs, their MAGs were mined to identify genes of the reversed Wood-Ljungdahl pathway (Figure 5). In all members of the Darwinibacteriaceae family, gene Acka was consistently detected, whereas Pta, Acs, the CODH complex, and MetF were conspicuously absent. Most of the Darwinibacteriaceae MAGs contained FolD, Fhs, and Fdh enzymes, which play vital roles in the acquisition of CO₂ and H₂ from formate. The absence of gene Pta and Acs could be justified by the presence of glycine reductase (Grd) in the glycine cleavage system, which facilitates the conversion of glycine for entry into the GCS (as reported by C. Li et al., 2022). Thus, the entire reverse WL-GCS pathway was encoded by members of Darwinibacteriaceae, with only two genes, namely Pta and Acs missing, which are in fact not necessary for the completion of the pathway, as in the GCS, CH₃-CO-Pi is directly converted to glycine through the function of glycine reductase (Grd) (Li et al., 2022). The remaining enzymes were identified in more than 70% of the Darwinibacteriaceae MAGs, with the exception of Dld, which was present in only 60% of them.

In the case of the family Wallacebacteriaceae, only genes AckA and Fhs were identified, being the other genes of the pathway were absent. The absence of critical enzymes of the reversed Wood-Ljungdahl pathway, including Sda, Grd, FolD, and Fdh suggests that members suggests that Wallacebacteriaceae are not syntrophic acetate oxidising bacteria.

Based on the evidence found, it could be hypothesized that the family Darwinibacteriaceae is a potential SAOB, as it encodes for the necessary genes to perform the reversed Wood-Ljungdahl pathway coupled to the glycine cleavage system. Within the intricate anaerobic digestion metabolic web, this pathway would allow these bacteria to perform the oxidation of acetate to H₂ and CO₂, reaction that would constitute a key step in anaerobic digestion, as hydrogenotrophic archaea would take this H₂ as a substrate to ultimately produce methane. However, the family Wallacebacteriaceae lacks several key enzymatic of this pathway. Considering the relatively low abundance of this taxon in the analyzed bioreactors (as detailed in Supplementary Table 6), our hypothesis suggests that Wallacebacteriaceae may not play a significant role in the functioning of anaerobic digesters, unlike Darwinibacteriaceae. This could be attributed to the absence of a symbiotic relationship between Wallacebacteriaceae and hydrogenotrophic archaea.
Figure 5. Mining of reverse WL pathway (black lines) coupled with glycine cleavage system (purple lines) (WL-GCS) genes in MAGs of the families Darwinibacteriaceae (n=45) and Wallacebacteriaceae (n=16). Red to green colors indicate the percentage of MAGs in which the genes were identified. A gene was considered to be present in each family if it was found in more than 70% of the MAGs. Bacteria from the family Darwinibacteriaceae encode the genetic repertoire necessary to metabolise acetate to produce CO2 and H2, which are then used by hydrogenotrophic archaea to produce methane, whereas Wallacebacteriaceae showed a truncated pathway. Abbreviations: AckA; acetate kinase, Pta; Phosphate acetyltransferase, Acs; Acetyl-coA synthetase, Por; pyruvate-ferredoxin reductase, Sda; L-serine dehydratase, GlyA; glycine hydroxymethyltransferase, Grd; glycine reductase, GcvP; glycine dehydrogenase, GcvH/T; glycine cleavage system H/T protein, Dld; dihydrolipoyl dehydrogenase, PflD; pyruvate-formate lyase, AcsE; iron sulfur protein methyltransferase, metF; methylenetetrahydrofolate reductase, Fdh; methyltetrahydrofolate dehydrogenase/cyclohydrolase, Fhs; formate-tetrahydrofolate ligase, Fdh; formate dehydrogenase.

4. Conclusions

In this work, a cryptic bacterial group previously referred as MBA03 in SILVA database, was described for the first time by recovering and analysing representative genomes belonging to this taxon. Phylogenetic analyses confirmed that MBA03 is a representative of a new taxonomic order with the proposed name Darwinibacteriales ord. nov. Two different families can be distinguished within this order: Darwinibacteriaceae fam. nov. (also known as DTU010 according to the GTDB taxonomy) and Wallaceae fam. nov (also known as DTU012).

Order Darwinibacteriales, particularly family Darwinibacteriaceae, is among the most abundant genera in anaerobic digesters, regardless of operational and chemical parameters such as feedstock and type of reactor. Darwinibacteriales is
a very specific bacterial group, being anaerobic digestion its main ecological niche. Bacteria from this order are also present in soils, sediments, plants and animal gut microbiomes at very low abundances. These materials are often used as substrates to feed anaerobic digesters, acting as a reservoir and an inoculum that allows Darwinibacterales to thrive when the AD process begins.

The presence of the reversed Wood-Ljungdahl pathway coupled to the glycine cleavage system in the family Darwinibacteriaceae and the co-occurrence of Darwinibacterales with several hydrogenotrophic archaea (Otto et al., 2023) reveals that they are potential syntrophic acetate oxidising bacteria, and thus that they may act as competitors for acetate with acetoclastic archaea, favouring hydrogenotrophic methanogenesis. Furthermore, these bacteria may be important during hydrolysis, like other members of the Clostridia class, since Darwinibacterales harbour a wide repertoire of hydrolytic enzymes. Nevertheless, it is important to highlight that this description has been entirely based on genome characterization, and so its culture and further laboratory analyses are needed to confirm what has been hypothesized.

The high abundance of Darwinibacterales in AD environments strongly suggests an exaptation mechanism to this industrial process. In other words, Darwinibacterales are naturally occurring hydrolytic and potentially SAOB bacteria that are pre-adapted to AD and are thus massively enriched in biogas producing and other AD facilities, where they may play an industrially relevant role that our results have only begun to reveal.
References


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