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Production of biohythane from cow dung using novel microbial synthetic consortia designed by heat-treated and acclimatized combined wastes†

Rashmi Ira,^a Vikas Sharma,^a Shrawan Kumar,^a Mira Koul,^a Lalita Sharma,^{ID b} Aditi Halder^{ID b} and Tulika Prakash^{ID *a}

The world's increasing dependency on fossil fuels has become a significant energy and environmental concern as they contribute 83% of the global energy supply and produce large amounts of carbon dioxide. Biohythane, a blend of biomethane (5–10%) and biohydrogen (50–60%), is emerging as a promising and environmentally friendly alternative fuel derived from organic wastes and offers a sustainable solution. The existing methods of biohythane production suffer from major limitations of being cost- and labor-intensive due to adopting bulk substrate pretreatment to enhance biohythane yield thereby limiting their industrial applications. In this study, we have developed a synthetic microbial consortium (E(C2)Tx) for anaerobic digestion by combining various organic wastes and subjecting them to heat pre-treatment and acclimatization to enrich biohydrogen producers and methanogens, respectively. Raw cow dung was anaerobically digested as the substrate with E(C2)Tx and this resulted in the production of biohythane with 3% biohydrogen and 36% biomethane. The consortia designing strategy avoided any bulk substrate pretreatment and only included the pretreatment of the inoculum which is used in four times less volume than the substrate. A 16S rRNA gene based metagenomic analysis revealed that the CD samples treated with E(C2)Tx were enriched in cellulolytic and hydrogen-producing *Firmicutes*, along with methylotrophic and hydrogenotrophic methanogens. The developed technology offers promising commercial benefits by requiring less energy for biohythane production. In addition, it offers environmental advantages by providing an efficient CD waste management alternative and reducing climatic impact by lowering greenhouse gas emissions associated with fossil fuel burning. Using a waste complementarity approach for consortia designing aligns with the principles of circular economy and presents a sustainable, scalable energy solution. The developed method can support the growing energy market by increasing biohythane yield and lowering its production cost.

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Environmental significance

Biohythane is a blend of CH₄ and H₂ and is an excellent replacement for non-renewable energy sources. The current biohythane production method involves the pre-treatment of a bulk substrate, which is a cost, energy, and labor-intensive process. Therefore, there is a need for the development of an improved synthetic microbial consortium for this issue, especially in single-stage anaerobic digestion. In this work, we have developed a synthetic microbial consortium using various organic wastes and adopted a heat pre-treatment strategy to enrich H₂-producing microbes combined with the acclimatization process to augment methanogens. Following this, we tested the performance of two types of designed consortia (E(C2)Tx and E(C2)UTx) for biohythane production *via* anaerobic digestion of raw cow dung as a substrate. In addition, we explored the role of different microbial taxa in the anaerobic digestion process, revealing that our synthetic microbial consortium can overcome extreme environmental conditions and work more robustly. The organic composition of the substrate and inoculum allowed the dilution and counteracting of the various toxic compounds or acids from the substrate and inoculum blend, making suitable conditions for microbes' growth, degrading the substrate, and producing biogas/methane. Additionally, our consortia designing approach and findings would stimulate further research in understanding the waste complementarity concept at the substrate level and with various wastes as potential inoculum.

Introduction

Due to the current global environmental and energy scenarios, the world's increasing dependency on fossil fuels has become a major concern. The global energy demand rose 2.9% in 2018 and is expected to reach 740 million terajoules by 2040, with an expansion of 30%.¹ Currently, 83% of this energy is obtained

^aSchool of Biosciences and Bioengineering, Indian Institute of Technology, Mandi, Kamand, Mandi, Himachal Pradesh 175005, India. E-mail: tulika@iitmandi.ac.in; Tel: +91-1905-267140

^bSchool of Chemical Sciences, Indian Institute of Technology, Mandi, Kamand, Mandi, Himachal Pradesh-175005, India

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from fossil fuels (mainly coal, oil, and natural gas) which substantially raises greenhouse gas (GHG) emissions (especially carbon dioxide (CO₂)) and is leading to changing climatic conditions.² Renewable energy sources such as biogas, wind, solar energy, *etc.*, are emerging as better alternate options in terms of being more sustainable and generating a lower carbon footprint.³ In addition, implementing circular economy principles (namely reusing, reducing, and recycling wastes) helps lower the carbon footprints by reducing resource extraction and generation of waste.^{4,5} Following appropriate carbon pricing mechanisms, renewable energy policies are expected to reduce greenhouse gas emissions and lead to a lower carbon price level.⁵ As a result, renewable energy sources, including biogas, are forming a relatively significant part of primary energy sources in several countries worldwide.⁴

The transition to renewable energy sources is not very swift and profitable and suffers from its own challenges including lower production and reduced supply of inputs.⁴ For any technology to be sustainable it must be profitable and able to attract investors' interest. From an investor's perspective, investing in funds with guaranteed returns may be a preferred choice,⁶ although a technology shock is shown to be associated with a growth in private investments. In addition to this, government policies to promote a circular economy *via* adopting renewable energy as a mechanism are also likely to play a significant role in investments in this direction.⁷ Another bottleneck in the utilisation of renewable energy sources is cost competitiveness. For example, microalgae emerged as a promising potential feedstock for biodiesel production while treating wastewater. However, the commercial and economic aspects of the microalgae biodiesel business depend on microalgae production cost, which itself suffers from several limitations.^{8,9}

Recently, the blend of biomethane (CH₄) with biohydrogen (H₂) (concentration between 10% and 30% v/v), known as biohythane¹⁰ or H₂-enriched compressed natural gas (HCNG), has emerged as another promising clean fuel. CH₄ and H₂ are widely used by chemical industries due to their high calorific values of 143 kJ g⁻¹ and 55 kJ g⁻¹, respectively.¹¹ Biohythane offers a strong potential to enhance energy recovery by ~8–43% along with a significant improvement in economic viability.¹² As biohythane can reduce carbon monoxide (CO) emissions by up to 50%, it has been used directly as a commercial vehicle fuel in various countries including India, California, China (Beijing Hythane Bus Project), Canada (Montreal Hythane Bus Project),^{13,14} and the well-established automobile companies such as Volvo, Fiat, *etc.*^{15,16}

The valorization of biohythane from biomass feedstocks or its microflora is emerging as a promising renewable energy generation technology. Biohythane fuel is derived from biomass feedstocks *via* the anaerobic digestion (AD) of organic waste, for example, food, microalgae, sewage sludge, and animal manure, including cow dung (CD).¹⁷ H₂ production in a single-stage dark fermentation AD process yields biohythane with 5–10% H₂, 50–60% CH₄, and 30–40% carbon dioxide (CO₂).^{10,18} It is estimated that animal manure produces 25 million tons of pollutants annually throughout the world.¹⁹ If untreated, these pollutants lead to the emission of noxious gases and repugnant odor,

thereby necessitating the development of efficient waste management strategies. CD is an organic waste, which is rich in crude proteins, minerals, fibers, lignocellulosic components, and a diverse microbial community.²⁰ Some of the CD waste management methods include the production of biogas (*via* anaerobic fermentation) and biochar (through pyrolysis) and recovery of plant nutrients^{21–23} among others. The utilization of CD as an independent substrate or in co-digestion with other organic wastes for biogas production through the AD process has been very widely explored.^{19,24–26} Cow manure of 1 kg can produce 35–40 L of biogas when mixed with water for a hydraulic retention time (HRT) of 55–60 days at an ambient temperature of 24–26 °C.²⁷ However, there are only a limited number of studies that have focussed on using CD as a substrate for biohythane production.

The type of feedstock substrate, inoculum, and preprocessing strategies that are used during the AD process significantly affect the yield and quality of biohythane. Various studies have explored different approaches to optimize H₂ and CH₄ production to meet the requirements of standard biohythane composition.^{1,12,28} Many recent studies employing various different pretreatment strategies on different types of substrates (*e.g.*, food waste, municipal waste, sludge, algal biomass, animal waste, and agricultural waste) and inoculums (pure or mixed cultures) to enhance H₂ yield during biohythane production have been reviewed.^{1,12,13,29} It is clearly apparent that, first, there are very few studies that have utilised CD as a substrate for biohythane production. Second, in some of the studies simple and easily consumable carbon sources such as glucose, glycerine, cellulose, fruit juices, *etc.*, which themselves are obtained by preprocessing, have been used as substrates.¹³

Third, in order to facilitate the hydrolysis of lignocellulosic components of a substrate such as CD and make them available for microbial digestion to improve biohythane (or biogas) production several pre-treatment strategies of the substrate have been adopted. These pre-treatment strategies utilize various physical (heat, sonication, *etc.*) and chemical methods (treatment using acid, alkali, *etc.*) or a combination of both.^{30–33} Biological pre-treatment is another alternative method of inoculating industrial cellulolytic enzymes or microbes into the substrate to degrade lignocellulosic components under mild and controlled environmental conditions. However, a major limitation of these substrate pre-treatment procedures is the increased consumption of energy and chemicals, which can lead to an extreme pH or temperature change and the formation of toxic by-products.³⁴ Besides, these substrate pre-treatment procedures are labor- and/or energy-intensive which makes them economically not preferable for managing large quantities of the substrate in real-life applications.³⁵

Fourth, a large majority of studies have adopted bulk substrate pre-treatment instead of pre-treating inoculum, which is used in much lower quantities as compared to the substrate. Some of the adopted pre-treatment methods of inoculum include heat treatment,³⁰ sequential re-inoculation for enrichment,³⁶ blending of different manures,³⁷ acclimatizations,³⁸ *etc.*, however, they have been employed independently of each other. Furthermore, some studies have used pure cultures and



synthetic media to maintain them for enhancing H_2 production. Pure cultures, although effective, require highly sterile operating conditions and extremely pure substrates, making them non-economic and impractical for industrial use. Mixed cultures, while economically viable, face challenges from non-hydrogen-producing organisms like methanogens, lactic acid bacteria, and homoacetogenic bacteria.¹³ Finally, some studies have performed two-stage AD or continuous AD under different organic load rates and HRT. Maintaining such a setup at the industrial scale requires additional costs limiting their economic viability.

Taken together, the existing methods of biohythane production suffer from several limitations including being labor-, energy-, and cost-intensive which make their real-life application very challenging. This indicates a need for the development of technologies for biohythane production which are environment friendly and free from bulk substrate treatment, use readily available feedstock, contribute towards waste management and thereby the circular economy, and most importantly are cost-effective. Towards this, in the present study, we have developed a technology for the preparation of novel synthetic microbial consortia by combining various types of wastes, namely, aged sludge (AS), paper mill sludge (PM), sewage wastewater, and CD, which is capable of producing biohythane from the CD substrate. Our approach mainly relies on pre-treating the synthetic inoculum, which is used in four times less quantity as compared to the bulk substrate in a single-stage dark fermentation reactor. This microbe-based AD process entails a complex and interconnected anaerobic

microbial consortium that performs hydrolysis, acidogenesis, anaerobic/syntrophic oxidation, and methanogenesis in synergy. In such a process, the heat treatment method is known to inhibit the H_2 consumers and the methanogenic activity, whereas the spore-forming H_2 -producing bacteria survive under such stressful conditions. Similarly, a pH range of 5.0 to 6.5 is known to favor the growth of H_2 -producing microbes.^{29,39,40} We adopted the heat pre-treatment and acclimatization strategies during our synthetic microbial consortia designing and compared the biohythane production among the treated and untreated inoculum with respect to the control. Furthermore, in order to understand the role of the microbial community dynamics during the AD processes, we have performed a 16S ribosomal RNA metagenomic sequencing analysis of the samples inoculated with heat-pre-treated (E(C2)Tx), untreated (E(C2)UTx), and control samples at the start of the AD process and after 40 days in a single-stage anaerobic digester at the lab-scale.

Materials and methods

In this study, we have used fresh CD as a feed to evaluate the potential of novel synthetic microbial consortia developed by us to produce biohythane. The complete workflow and study design are given in Fig. 1.

Novel synthetic microbial consortia as the inoculum

The novel synthetic microbial consortia used in this study consist of four different waste products as the natural mixed

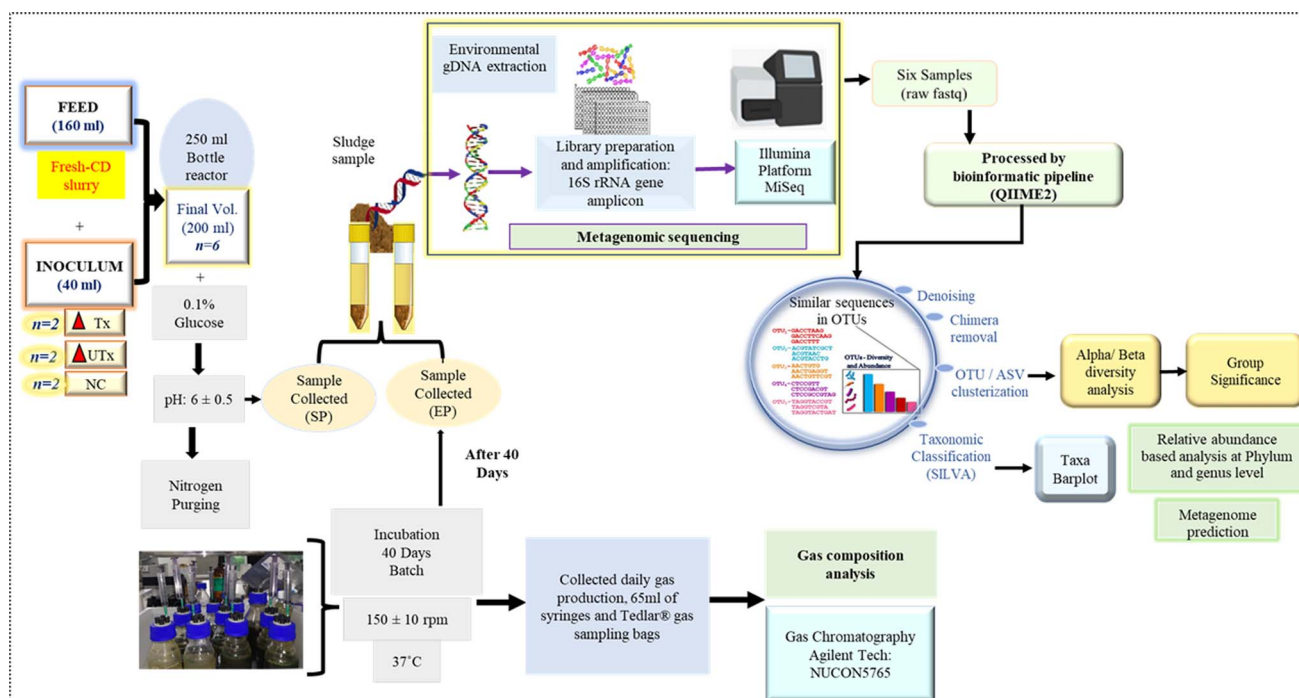


Fig. 1 Schematic representation of the detailed workflow of the experimental setup and analysis. CD: cow dung, Tx: heat-pre-treated (at 80 °C for 2 hours with continuous magnetic stirring), UTx: heat untreated, NC: negative control (cow dung slurry), SP: starting point of the experimental set-up before nitrogen purging (at 0 hours), EP: endpoint of the experiment (after 40 days), gDNA: genomic deoxyribonucleic acid, OTU/ASV: operational taxonomic unit/amplicon sequence variants.



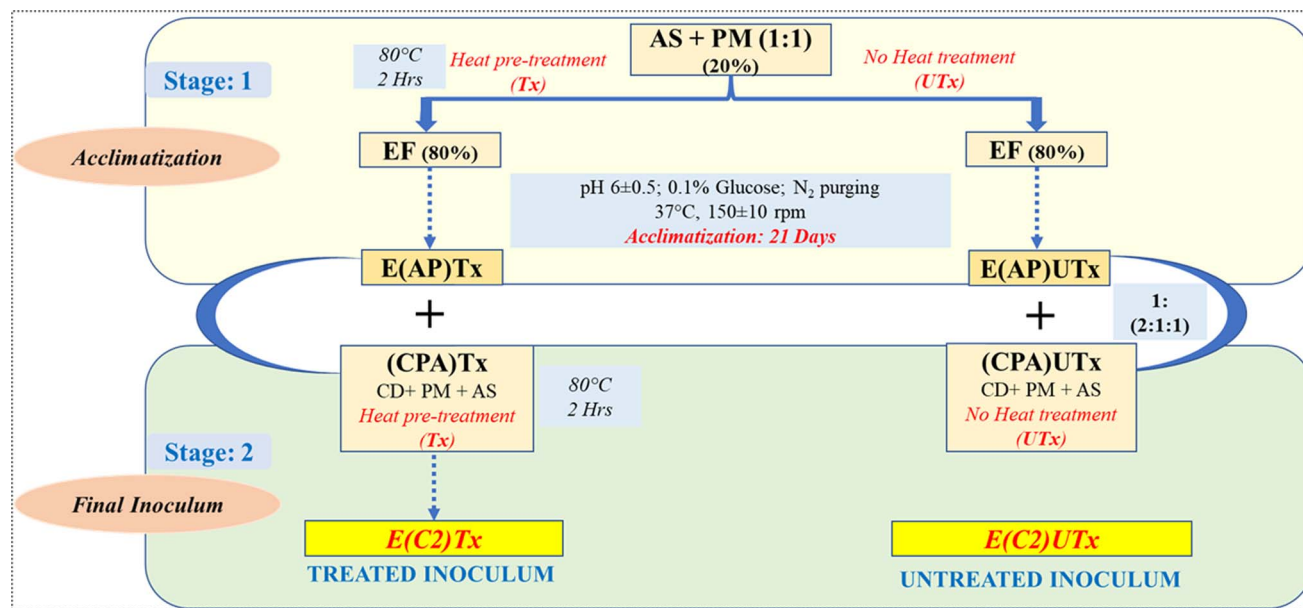


Fig. 2 Schematic representation of the preparation and composition of the synthetic microbial consortia. AS: aged sludge, PM: paper mill sludge, EF: electro-flocculated wastewater, CD: cow dung, Tx: heat-pre-treated (at 80 °C for 2 hours with continuous magnetic stirring), UTx: heat untreated, E(C2)Tx: treated inoculum and E(C2)UTx: untreated inoculum.

cultures: aged sludge (AS) obtained from the local sewage treatment plant (STP) at IIT Mandi (Kamand, Mandi, Himachal Pradesh, India), paper mill sludge (PM) obtained from Shreyans Paper Mill Ltd, Banah (Punjab, India), flocks of electro-flocculated (EF) wastewater were freshly obtained after performing the electro-flocculation technique on freshly collected sewage wastewater at IIT Mandi, as mentioned by Lalita *et al.* 2021,⁴¹ and CD obtained from local cowshed at Salgi, near IIT Mandi. The detailed method for the preparation of the inoculum is described in Fig. 2.

Two types of synthetic microbial consortia were used in this study, namely, heat-pre-treated E(C2)Tx and untreated E(C2)UTx, and CD was used as a control. The compositions of both inocula were the same, however, the processing of the constituents to prepare the final inoculum differed on the basis of the heat pre-treatment step in the case of E(C2)Tx in order to retain the H₂-producing microbial community. Heat-pre-treatment was performed by boiling the constituents at 80 °C for 2 hours with continuous magnetic stirring.

The preparation of the synthetic inoculum was carried out in two stages. In stage 1, acclimatization of the primary constituents (AS + PM) was performed for 21 days using EF as a carbon source for the microbial community at a pH of 6 and mesophilic temperature of 37 °C. 0.1% glucose (HiMedia) was added during this step to facilitate microbial growth and purging of N₂ gas for 5 minutes to maintain an anaerobic environment. In stage 2, the acclimatized sludge prepared by using heat-pre-treated primary constituents was mixed with the heat-pre-treated CPA (CD + AS + PM) component in order to retain the effect of heat treatment throughout the process of inoculum designing for preparing the final treated inoculum (E(C2)Tx). The initial parametric conditions such as total solids (TS), volatile solids

(VS), chemical oxygen demand (COD), and pH have been listed for the inoculum E(C2)Tx, in the recently published study by Thakur *et al.* (2023).²⁴ However, for preparing the final untreated inoculum (E(C2)UTx), we mixed the acclimatized sludge prepared by using untreated primary constituents with untreated CPA components.

Feed preparation

Fresh CD was collected from a local cowshed at Salgi, near North Campus, IIT Mandi, (Kamand, Mandi, Himachal Pradesh, India). The cow was feeding on local vegetation, plants, and grass. Manual cleaning of the sample was performed by removing long grass. CD slurry was prepared by adding tap water in a proportion of 1:1 (v/v). The initial parameters including VS and TS (g VS_{added} and g TS_{added}, respectively) for CD as a substrate were calculated as previously described.²⁴

Experimental conditions

A mini-scale dark-fermentative single-stage bottle bioreactor was prepared in two replicates by using transparent 250 ml Duran® glass bottles that were sealed tightly with rubber corks having four septa, which were fitted with a needle of 65 ml syringes (Dispo Van®) for the collection of gas (Fig. 1). 160 ml of CD slurry as feed and 40 ml of synthetic microbial inoculum were added to prepare a final working volume of 200 ml in these 250 ml mini bottle reactors. This resulted in a substrate and inoculum composition of 4:1 (v/v) *i.e.*, 20% of synthetic microbial inoculum and 80% of the feed substrate. The final mixture was supplemented with 0.1% of glucose (HiMedia) and the pH was adjusted to 6 ± 0.5. To maintain the anaerobic conditions, the small-scale bottle bioreactor was purged with



nitrogen gas. The bottle bioreactors were placed in an incubator in the dark at a temperature of $37 \pm 1^\circ\text{C}$ at 150 ± 10 rpm for a period of 40 days (Fig. 1). Analysis of the gaseous mixture was carried out with the help of a gas chromatograph (Agilent, 7890B, Santa Clara, California, United States) which was equipped with two TCDs (Thermal Conductivity Detectors) and one FID (Flame Ionization Detector). All hydrocarbons were detected at the FID and hydrogen was detected at the TCD also known as auxiliary detectors where N_2 was used as a reference gas. The injector and detector were maintained at 60°C .

DNA extraction and metagenomic sequencing

Metagenomic DNA extraction. For the extraction of metagenomic DNA an appropriate amount of sampled sludge from each bottle was taken in 15 ml falcons at the beginning (SP) and end (EP) of the experiment (Fig. 1) and was immediately stored at 4°C until further use. The stored samples were thawed and washed thrice in phosphate buffer saline before DNA extraction. Metagenomic DNA extractions and purification were performed by using the genomic DNA from a soil extraction kit (Nucleo-Spin®96 Soil, MN, Takara) according to the manufacturer's protocol. The integrity of the extracted metagenomic DNA was determined on 0.7% agarose gel by electrophoresis, and a gel documentation system (Bio-rad, California, United States) was used for visualization. Qubit 3.0 fluorometer (Invitrogen Corporation, Thermo Fisher Scientific, United Kingdom) was used for evaluating the concentration of DNA with the Qubit dsDNA BR Assay Kit.

Next-generation sequencing library construction for 16S ribosomal RNA amplicon-based metagenomic analysis. To capture the microbial communities by metagenomic sequencing of the samples we proceeded with a single replicate of each type of bioreactor. The metagenomic 16S ribosomal RNA gene library was constructed by following the "16S Metagenomic Sequencing Library Preparation" guide by Illumina (Illumina, San Diego, CA, United States).

The forward (5'TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCTACGGG-NGGCWGCAG3') and reverse (5'GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGGACTACH-VGGGTATCTAATCC3') primers with overhang adapters were used for the amplification of the V3–V4 hypervariable regions of the microbial 16S rRNA gene by a polymerase chain reaction (PCR) (ProFlex PCR, Thermo Fisher Scientific, United States). PCR clean-up was performed using AMPure-beads (Beckman Coulter, California, United States) to purify the amplicons to remove free primers and primer dimers. Subsequently, bar-coding for pooling multiple samples was performed using an index PCR with Illumina sequencing dual-index primers provided in Nextera XT Index Kit v2 (Illumina, San Diego, CA, United States) followed by PCR clean-up to generate the final library (4 nM). Spiking in with 25% Phi-X was used as a sequencing control. Equimolar amounts of the respective 16S rRNA gene amplicon pools were sequenced by using a MiSeq platform (Illumina, San Diego, CA, United States) available in the in-house NGS facility of IIT Mandi, by using MiSeq v3

reagents with paired-end 300-bp read chemistry (2×300 bp). After trimming the adapters and primer sequences, raw fastq sequence files were acquired and subjected to further bio-informatics analysis.

Post-sequencing bioinformatics analysis

Sequenced reads in the FASTQ format were subjected to a comprehensive bioinformatics processing and analysis using the QIIME2, version 2021.8. (Quantitative Insights into Microbial Ecology) software suite.⁴² Raw reads were imported into QIIME2 with the "Casava 1.8 paired end demultiplexed fastq" method (Fig. 1). The raw file was filtered and trimmed first to remove all the sequences with an average Phred score or quality value of below 20 or size less than 20 bp. The DADA2 (Divisive Amplicon Denoising Algorithm 2) plugin was used to denoise and filter the reads, which includes trimming and truncating low-quality regions, read dereplication, chimera filtering, and singleton removal.⁴³ After denoising, open-reference clustering was performed with the VSEARCH plugin at 99% identity to generate operational taxonomic units (OTUs),⁴⁴ where reads were organized in features, and a feature table was generated with a feature-table plugin (<https://github.com/qiime2/q2-feature-table>). In the feature table, exactly one sequence represents each feature. The Amplicon Sequence Variants (ASVs) that have a frequency below 0.1% of the mean depth were excluded from further analysis. The feature-classifier plugin was used to assign taxonomy to OTUs against a pre-trained Sk-learn classifier⁴⁵ based on the SILVA 138 database⁴⁶ with a 99% similarity clustering threshold to generate taxonomy tables. Taxa filtering for eukaryota, mitochondria, and chloroplast was performed, and a final representative sequence file was generated for filtered taxonomy. The taxa bar plots depicting taxonomic hierarchy were generated using the QIIME2 taxa plugin.

Alpha rarefaction for the phylogenetic diversity was performed by alpha_rarefaction.py script using the minimum and maximum depth parameters as 10 and 2227, respectively. The analysis of both alpha and beta diversity was performed by the q2-diversity plugin in QIIME2. The alpha diversity metrics to the evenness, observed OTUs, and Shannon were measured based on ASV levels. The alpha diversity metric dissimilarities were tested by the non-parametric Kruskal-Wallis test. Alpha-diversity-group significance was determined from community richness in addition to the Shannon index.⁴⁷ Beta diversity metrics were measured by the Bray–Curtis distance, and the structure of the microbial community in all samples was explored with PERMANOVA⁴⁸ by using the beta-group-significance command.

Results and discussion

Designing of synthetic microbial consortia for producing biohythane by anaerobic digestion of cow dung

Establishing an enriched and efficient inoculum of microbial members to produce H_2 is crucial for the fermentation process during AD. For microbial enrichment, mixed cultures can be directly used from the natural environments. In various natural



environments, a variety of mixed cultures are found that have the extensive ability for fermentative H_2 generation. These include hot spring sediment, anaerobic sludge, compost, municipal sewage sludge, and soil. Such mixed cultures have an additional advantage due to the simplicity of their operation and control under non-sterile conditions which is not feasible while using pure cultures.⁴⁰ For the development of efficient synthetic microbial consortia the principle of “division of labor” has been used. This allows the microbial community to complete the process more quickly than monocultures by assigning various tasks to different members of the community. This enables a coordinated environment to support each process, *i.e.*, each consortium member produces a molecule that is then used by the following link in the supply chain.⁴⁹

We have designed our synthetic microbial consortia by combining four different natural mixed cultures, namely aged sludge (AS), paper mill sludge (PM), cow dung (CD), and electro-flocculated wastewater (EF) (Fig. 2). PM is rich in compounds of fiber wood (lignocellulosic), such as carbohydrate polymers, mainly cellulose and hemicellulose, lignin, lipids, and other extractives. Some members of *Firmicutes* (*Bacillus* sp., *Clostridium* sp., and *Paenibacillus* sp.), and *Actinobacteria* (*Streptomyces*, *Microbacterium*, and *Cellulomonas xylanilytica*) present in PM sludge showed cellulase activity, which may be important for the breakdown of feedstock.⁵⁰ AS obtained from STPs is rich in microbial diversity and contains a high concentration of organic matter with various nutrients including N_2 , P, K, Ca, Fe, Mg, Cu, Zn, Mn, and Si.⁵¹ AS contains aerobic and anaerobic microbes like bacteria, archaea, fungi, and protists. The predominant bacterial phyla in AS include *Proteobacteria*, *Bacteroidota*, *Firmicutes*, and *Actinobacteria*. The most dominating genera include *Clostridium*, *Treponema*, *Propionibacterium*, *Syntrophus*, *Desulfobulbus*, *Brevundimonas*, *Paludibacter*, *Cloacibacterium*, and *Methylobacterium*.⁵² The CD is enriched in hydrolyzing and acidifying bacteria and methanogenic archaea, such as *Bifidobacterium*, *Bacillus*, and *Lactobacillus*.⁵³ EF is obtained by the electro-flocculation of sewage raw wastewater from the STP of IIT Mandi and is rich in Fe content in the form of Fe^{2+} and Fe^{3+} ions,⁴¹ which play important roles in the AD process.⁵⁴ The production of H_2 and CH_4 requires different kinds of metal ions as micronutrients, namely, Fe^{2+} , Zn^{2+} , Ni^{2+} , Na^+ , Mg^{2+} , and Co^{2+} as cofactors, facilitators of transport processes, and structural skeletons of numerous enzymes mainly for the growth and metabolism of microbes.⁴⁰ Based on the form of Fe available, namely Fe^{2+} or Fe^{3+} , the microbial community is shifted towards CH_4 or H_2 production, respectively.⁵⁴

Another method for enriching a community is based on employing a selection mechanism to favor bacteria that produce H_2 while eradicating those that consume it. The majority of H_2 -producing bacteria are spore-forming, whereas methanogens and H_2 -consuming bacteria are not, except for a few hydrogenotrophic methanogens.⁵⁵ The production of endospores can be encouraged by providing a stressed environment to the microbes which will be favorable for the endospore formation of bacteria, such as the H_2 -producers while removing the others. In the present study, we used two types of

synthetic microbial inoculum, namely, heat-pre-treated and untreated, and tested the biohythane efficiency with CD feedstock as compared to the control group. *Bacillus* and *Clostridium* are the two most popular sporulating bacteria that are known to be mainly responsible for producing H_2 . Similarly, a slightly acidic pH, to avoid methanogenesis and solventogenesis (5.0 to 6.5), is one of the favoring parameters to support the growth of H_2 -producers and enhance the H_2 -production rate,^{29,40} whereas an optimum pH for methanogenic bacteria ranges between 6.0 and 7.5 and anaerobic bacteria work well below pH 6.⁵⁶ So, in order to favor both H_2 and CH_4 production, we performed our experiments at a pH of 6–6.5.

For the design of the synthetic microbial consortia, at first, we used AS + PM to prepare two types of mixed sludge with (EAPT_x) or without (EAPUT_x) heat pre-treatment (Fig. 2). These mixed sludges were independently acclimatized with EF for 21 days to allow the microbial community to adapt to the AD process. After 21 days of acclimatization under anaerobic conditions, the microbial community is likely to be shifted towards methanogenesis and H_2 scavenging conditions which may impact H_2 yield. In general, the methanogenic archaeal community is a slow grower as compared to the anaerobic bacteria. Therefore, it is important to overcome this risk of complete community shifting towards H_2 consumption and methanogenesis. Towards this, in the next step, we have mixed CPA consisting of AS, PM, and CD in our acclimatized sludge (EAPT_x or EAPUT_x). The heat-pre-treated CPA (CPA(T_x)) is combined with EAPT_x and the untreated CPA ((CPA)UT_x) is mixed with EAPUT_x to result in the final synthetic microbial consortia, namely, E(C2)_{Tx} and E(C2)UT_x (Fig. 2). The effect of heat pre-treatment is again to enhance the H_2 producers, which are mixed with a methanogenic microbial community enriched in the acclimatized sludge (EAPT_x). It is important to note that we have used CD as the substrate for AD in our analysis which is classified as a lignocellulosic waste due to the presence of a large proportion of lignocellulose (50% in dry matter).³³ Therefore, it is important to enhance the degradation of recalcitrant lignocellulose in CD so that the overall yield of biohythane can be increased. The use of PM and CD in the last step (CPA) is also to provide a community rich in lignocellulose-degrading microbes to facilitate the process of hydrolysis and fermentation.

Gas production and composition

A mini-scale dark-fermentative single-stage bottle bioreactor system was used for biohythane production from CD as feed and synthetic microbial inoculum for a period of 40 days. The total gas production was measured under two experimental conditions, namely, the digestion of the CD substrate with E(C2)_{Tx} and E(C2)UT_x with respect to the control (only CD as feed without any inoculum). Fig. 3 represents the cumulative gas volumes for different gases produced in 40 days for this single-stage fermentation process. CD samples inoculated with E(C2)_{Tx} showed the highest volume of total gas (1247 ml) produced followed by CD inoculated with E(C2)UT_x (1052.5 ml) and CD control (963.5 ml).



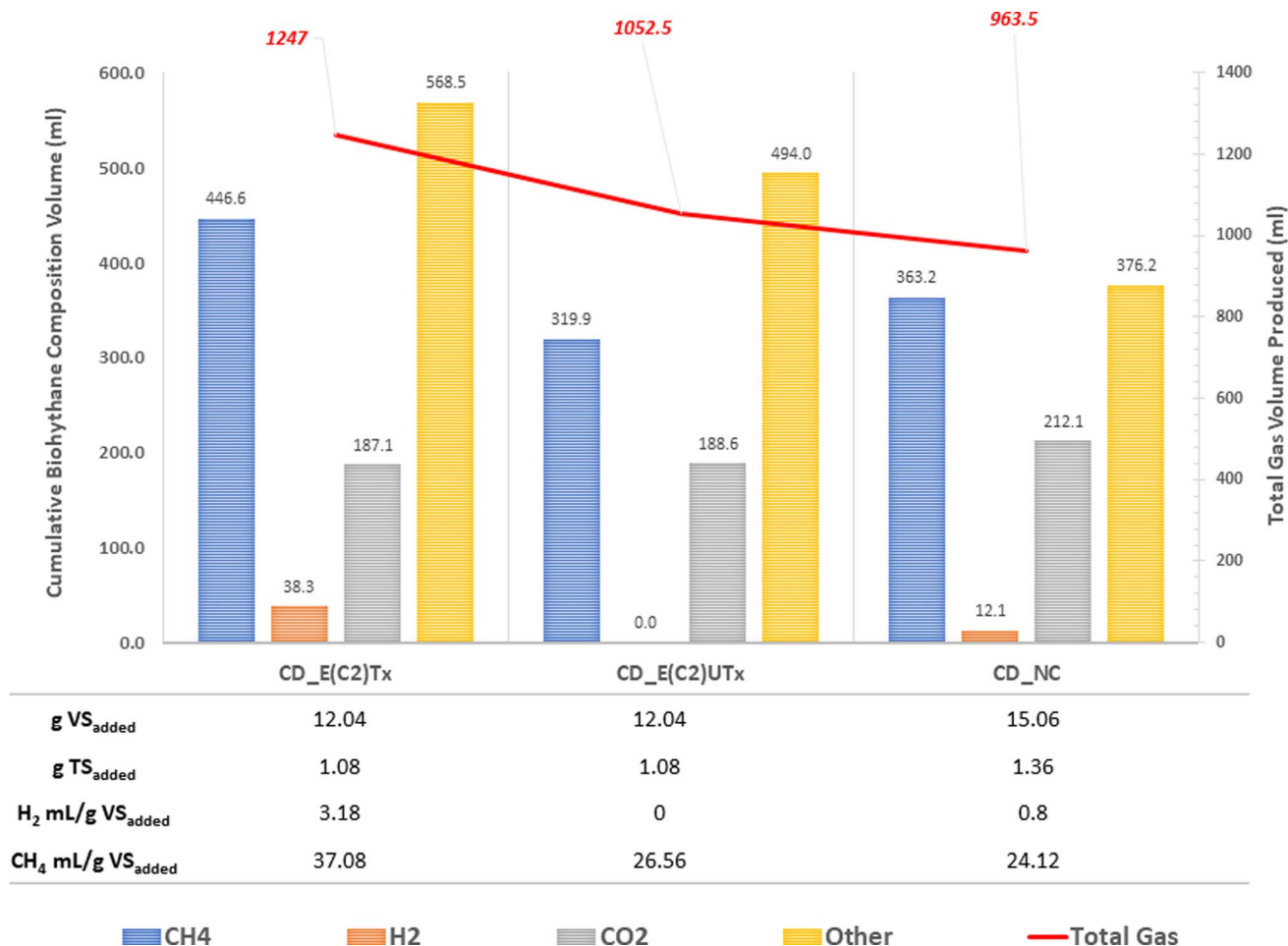


Fig. 3 Bar plot representing cumulative and total gas volumes produced during the anaerobic digestion of the CD substrate with different inocula. The gram volatile solids (g VS) and gram total solids (g TS) for the CD substrates are provided for each inoculum. Biohydrogen and biomethane yields are provided in $\text{mL g}^{-1} \text{VS}_{\text{added}}$. "Other gases" include the other gases or gaseous components, mainly nitrogen. CD: cow dung, E(C2)Tx: heat-pre-treated inoculum, E(C2)UTx: heat untreated inoculum, H₂: biohydrogen, CH₄: biomethane, CO₂: carbon dioxide.

We also measured the biogas production daily to estimate the duration of a cycle for the production of H₂ and CH₄ (Fig. 4). In the case of E(C2)Tx, a drastic increase in gas production was observed on the 7th day of the experiment (Fig. 4(a)). In the case of E(C2)UTx samples, a rise in gas production was observed only after the 11th day, whereas for control samples this was observed after the 15th day (Fig. 4(a)). For the E(C2)Tx, the highest gas volume was obtained on the 8th day (112.5 ml) (Fig. 4(a)), whereas for E(C2)UTx and control samples, the highest gas volume was produced only on the 31st day (137.5 ml) and 26th day (125 ml), respectively. These observations show that there is a significant reduction in the cycle length of biogas production in the case of CD samples mixed with E(C2)Tx. A similar decrease in biogas production cycle length was observed previously while using E(C2)Tx inoculum for the anaerobic co-digestion of food waste (FW), CD, and sewage wastewater (FCR_E(C2)Tx) for 20 days.²⁴

In the single-stage biohythane production, the rate-limiting step is H₂ production, which ultimately also governs the composition of the biohythane. Cumulative H₂ volume is 38.3 ml (3.17 ml H₂ per g VS_{added}) for CD inoculated with E(C2)Tx

and 12 ml (0.79 ml H₂ per g VS_{added}) for the control, whereas the samples inoculated with E(C2)UTx do not show any H₂ production (Fig. 3). There is a substantial increase in the H₂ production in the E(C2)Tx sample on the 1st and 5th days and the maximum volume was obtained on the 7th day (12.6 ml) (Fig. 4(b)). For the control sample, H₂ production started on the 2nd day and reached the maximum on the 3rd day (6.4 ml), although the total volume of H₂ produced was less than that in the case of E(C2)Tx samples. However, no H₂ was obtained in the E(C2)UTx sample till the end (Fig. 4(b)).

The production of H₂ is followed by the production of CH₄. The cumulative volume of CH₄ is found to be the highest for CD inoculated with E(C2)Tx (446.6 ml, 37.08 $\text{mL g}^{-1} \text{VS}_{\text{added}}$) (Fig. 3). This is followed by the CH₄ production in E(C2)UTx (319.9 ml, 26.56 $\text{mL g}^{-1} \text{VS}_{\text{added}}$) and control (363.2 ml, 24.12 $\text{mL g}^{-1} \text{VS}_{\text{added}}$) samples (Fig. 3). A gradual increase in CH₄ production can be observed from the 7th day onwards for E(C2)Tx, with a maximum volume obtained on the 8th day (55.4 ml) (Fig. 4(c)). CH₄ production continued to increase at the rate of approximately 16 ml per day up to day 32, after which only ~5 ml per day of CH₄ was obtained till the end of the experiment. While in



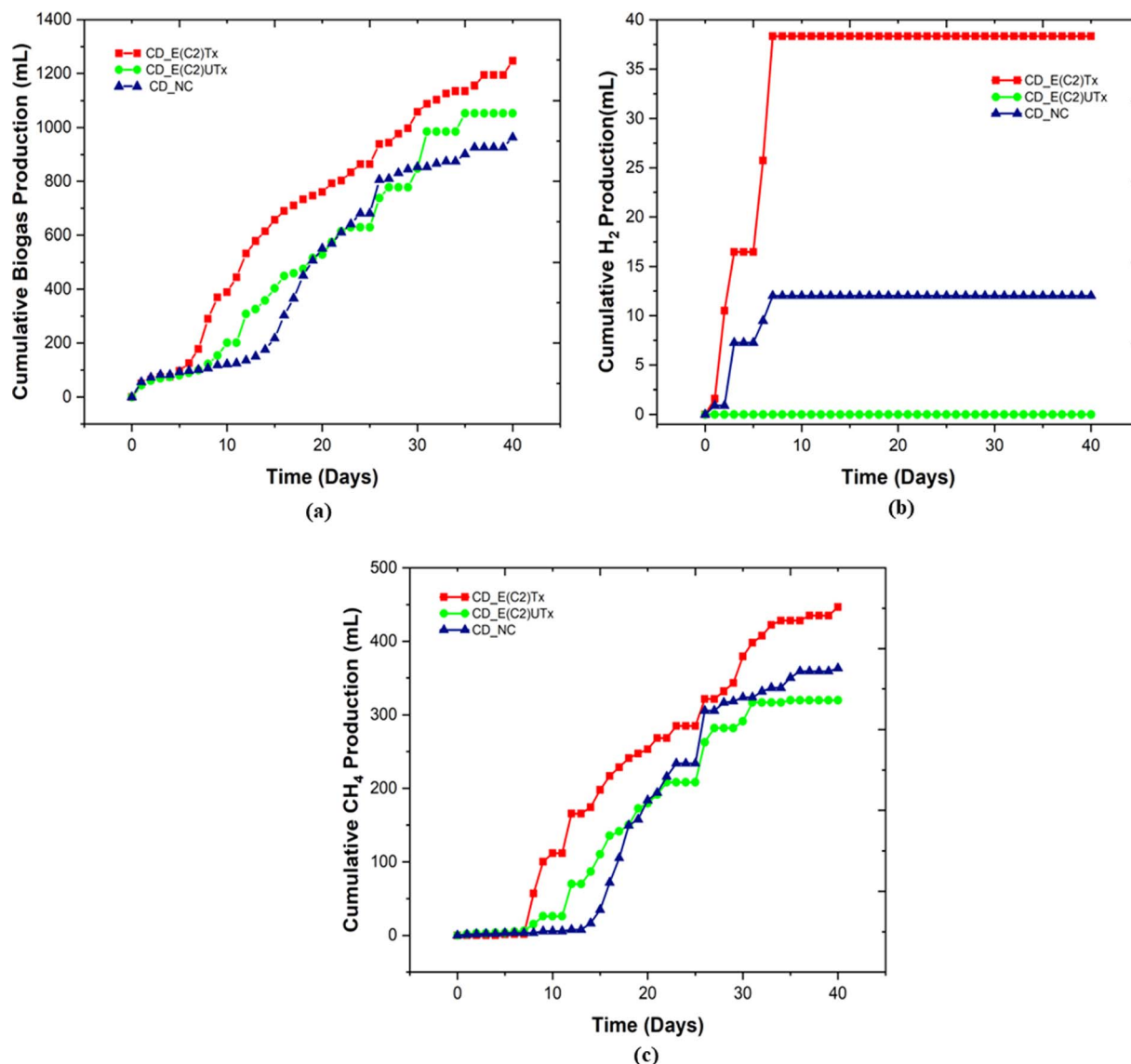


Fig. 4 Plots representing the cumulative volumes of (a) biogas, (b) biohydrogen and (c) biomethane produced during the complete operation period of 40 days. E(C2)Tx: heat-pre-treated inoculum, E(C2)UTx: untreated inoculum, CD_NC: negative control (CD alone).

the case of E(C2)UTx, a slight increase in CH_4 production can be observed only from the 11th day onwards with a maximum CH_4 volume of 54.7 ml obtained on day 25. In the case of the control samples, an increase in CH_4 production was observed only by day 15, and the maximum volume was obtained on day 26 (71.7 ml).

These observations show that with E(C2)Tx, a higher proportion of H_2 (3%) was obtained, whereas in control samples a lesser percentage of this gas was recovered (1.3%). In E(C2)UTx, no H_2 production was observed. Additionally, with E(C2)Tx, the cycle of H_2 production begins on the 1st day itself and that of CH_4 production starts on the 7th day onwards, whereas in the control and E(C2)UTx samples a delayed CH_4 production with less or no H_2 production, respectively, was observed. This implies that the anaerobic digestion of the CD substrate with

the E(C2)Tx inoculum is capable of generating biohythane, although with only 3% of H_2 gas.

Biohythane production, targeting more than 5% H_2 , has been explored in several studies previously.^{57–63} For example, Nguyen *et al.*, (2022)⁶⁴ used a mixture of swine manure and pineapple waste for the production of biohythane in a single-stage two-chamber digester with varying HRT (96 h to 6 h) and observed 60.5% of peak H_2 content in biohythane at an HRT of 6 h. Ta *et al.*, (2020)⁶⁵ performed biohythane production in a single-stage anaerobic digester using synthetic nutrient media as feed and separately entrapped H_2 - and CH_4 -producing bacteria and obtained a peak of H_2 (64.6 ml/L-d, 15%) and CH_4 (395 ml/L-d). In another study, biohythane with a maximum of 3.21 mol H_2 per mol hexose and 3.63 mol CH_4 per mol hexose from acetic acid and glucose (synthetic wastewater) in a continuous stirred tank reactor (CSTR) was obtained.⁶⁶ The AD



of CD for biohythane production has also been explored, however, utilizing other co-digestion substrates.^{67,68} For example, Sufyan *et al.*, (2023)⁶⁸ performed a two-step AD using a mixture of domestic wastewater and CD with varying pH ranges and maintaining thermophilic conditions (55 °C) for H₂ production and mesophilic conditions (35 °C) for CH₄ production. They observed the highest yield of H₂ (108.04 ml H₂ per g VS) and CH₄ (768.54 ml CH₄ per g VS) at pH 5.5 and pH 7.5, respectively, and the highest cumulative biohythane (811.12 ml g⁻¹ VS) was obtained at a pH of 7.5.⁶⁸ As is evident, most of the studies are based on either the two-stage AD process or with a variety of different substrates (such as wastewater, palm oil mill effluent, food waste, distillery spent-wash, swine manure, fruit waste, algae, *etc.*) under energy or labor-intensive conditions, thereby leading to limited real life applications.

Taxonomy analysis of CD treated with different inocula

In order to estimate the microbial diversity dynamics before the start (SP) and end (EP) of AD, we performed the 16S rRNA-based metagenomic analysis. The amplicon sequencing yielded a total of 131 121 raw reads and 128 051 quality-filtered reads for all 6 samples (Table 1). The quality-filtered reads resulted in a total of 779 Operational Taxonomic Units (OTUs). After removing singletons and contaminants 756 OTUs were finally retained which were distributed among 30 phyla and 365 genera. A varying sampling depth across the samples is known to introduce bias in the estimation of rare taxa. A rarefaction analysis to estimate the minimum number of reads required to capture the complete diversity of each sample (ESI Fig. 1†) indicated that the sampling across all samples was found to be sufficient to capture the complete diversity of each dataset. The CD samples with E(C2)UTx exhibited the highest microbial diversity as they harbor the maximum number of unique OTUs followed by the samples with E(C2)Tx and controls (Table 1 and ESI Table 1†). This could be due to the loss of heat-intolerant microbial community during the heat pretreatment step of E(C2)Tx inoculum preparation. During heat pretreatment, the high temperature could lead to the disruption of chemical bonds of the microbial cells leading to a loss of heat labile community.⁶⁹ This indicates that the type of inoculum pretreatment causes different microbial communities which can affect the overall H₂ and CH₄ production during the AD process. The beta-diversity analysis demonstrated no significant clustering of samples.

Phylum-level microbial community composition. For estimating the microbial dynamics across samples at the phylum

level, all taxa with a relative abundance of 0.5% or higher are considered. A detailed analysis of the predicted OTUs showed that the number of bacterial OTUs was much higher than that of the archaeal OTUs (ESI Table 1†). At the phylum level, most sequences were classified within three dominant bacterial phyla, namely, *Firmicutes*, *Bacteroidota*, and *Proteobacteria*. However, the relative abundance of these taxa was found to vary among samples based on the process of treatment and sampling points. *Bacteroidota*, *Firmicutes*, and *Proteobacteria* are found to be abundant in several AD systems.⁷⁰ Furthermore, these phyla contain several species that are known to participate in one or more phases of the general AD process.

The top ten most abundant bacterial or archaeal phyla from each sample are shown in Fig. 5. The microbial community of the E(C2)Tx samples was mainly enriched with phylum *Firmicutes*, whereas in all other samples, the relative abundance of this phylum was comparatively less. The higher abundance of *Firmicutes* in E(C2)Tx samples throughout the AD process may be responsible for a higher hydrolysis rate and enhanced H₂ and biohythane production.⁷⁰ The members of this phylum participate in the hydrolysis, acidogenesis, and acetogenesis steps of AD and are known to degrade a wide range of substrates (protein, lipids, and carbohydrates).^{54,71} The abundances of *Proteobacteria* and *Bacteroidota* were the highest in the SP of E(C2)UTx samples. *Proteobacteria* include important acetogenesis bacteria and are well known for degrading or utilizing glucose, proteins, and various kinds of VFAs in manure and other lignocellulosic wastes.⁷¹ The phylum *Bacteroidota* can hydrolyze carbohydrates or proteins and transform organic substances to produce VFAs and is considered to participate in the hydrolysis and acidogenesis steps of AD.⁷⁰ Interestingly, the EP of control samples showed the highest abundance of phylum *Chloroflexi*, which is found to be negligible in all other samples. The members of phylum *Chloroflexi* are often treated as primary fermenters and are ubiquitous in various AD systems including cattle manure, although their metabolism is not well defined though they are assumed to utilize glucose and break down tough-to-degrade organics.⁷¹

Among the archaeal taxa, *Euryarchaeota* is found to be enriched in samples with E(C2)Tx as compared to E(C2)UTx and controls, whereas *Halobacterota* is present in least abundance. A majority of the members of phylum *Euryarchaeota* utilize the hydrogenotrophic and methylotrophic pathways for methanogenesis.⁵⁴ *Halobacterota* can act on a wide range of substrates

Table 1 Table representing the summary of the raw and quality filtered sequence data

| Sr. no. | Sample ID | Total reads | Read length | Reads passing quality filtering | % reads passing quality filtering | Number of observed OTUs/feature count | Number of unique OTUs |
|---------|----------------|-------------|-------------|---------------------------------|-----------------------------------|---------------------------------------|-----------------------|
| 1 | CD_E(C2)Tx_SP | 24 416 | 301 | 23 801 | 97.50% | 3140 | 98 |
| 2 | CD_E(C2)Tx_EP | 25 539 | 301 | 24 905 | 97.50% | 3403 | 96 |
| 3 | CD_E(C2)UTx_SP | 21 991 | 301 | 21 505 | 97.80% | 4275 | 124 |
| 4 | CD_E(C2)UTx_EP | 23 072 | 301 | 22 401 | 97.10% | 3474 | 111 |
| 5 | CD_NC_SP | 22 272 | 301 | 21 847 | 98.10% | 2912 | 74 |
| 6 | CD_NC_EP | 13 831 | 301 | 13 592 | 98.30% | 2227 | 58 |



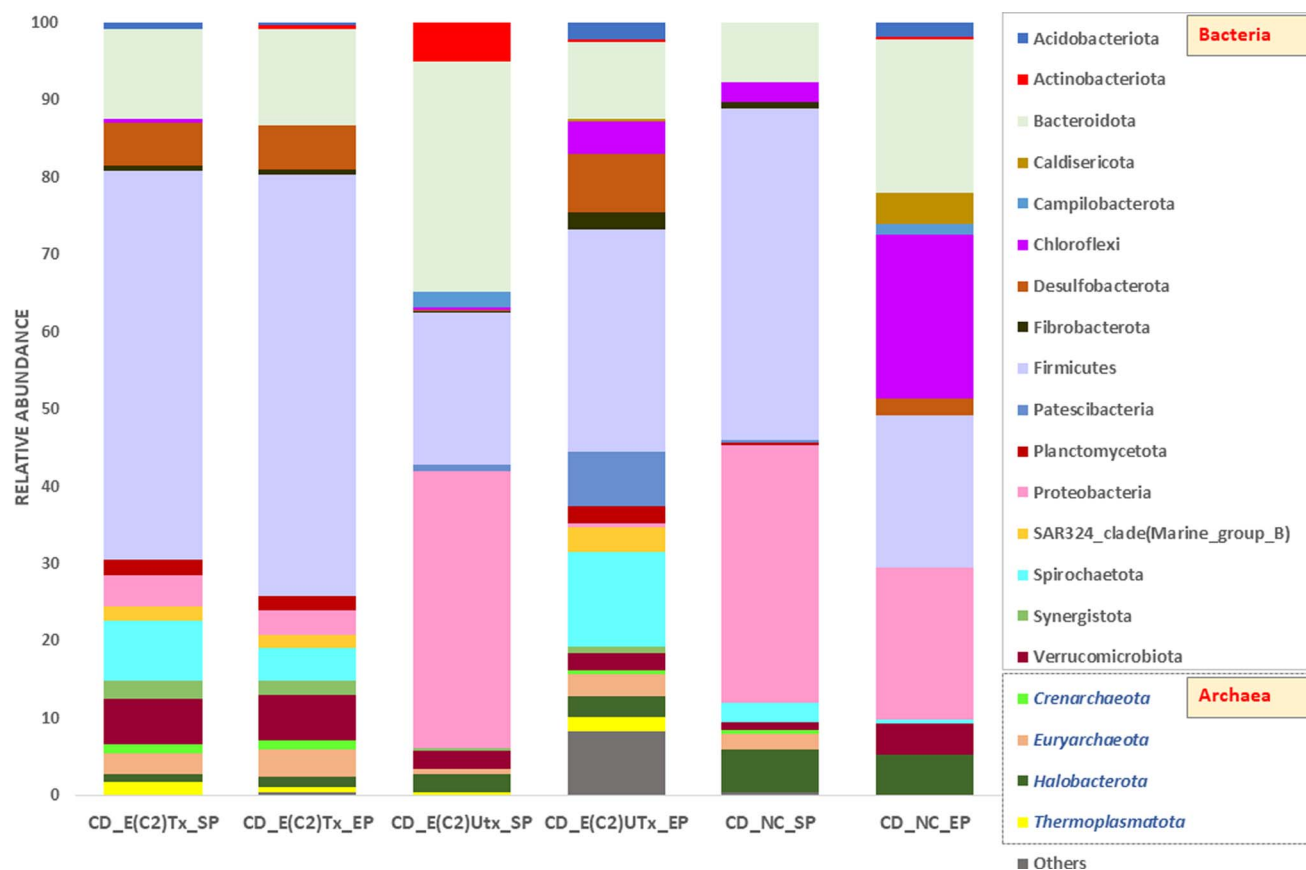


Fig. 5 Stack bar plots representing the relative abundance of the top ten bacterial or archaeal phyla. Samples were collected for each category at the starting (SP) and end (EP) points of the anaerobic digestion of CD for a period of 40 days. The remaining microbial community is marked in the category of "Others". E(C2)Tx: heat-pre-treated inoculum, E(C2)UTx: untreated inoculum, NC: negative control.

and produces CH_4 using any of the methanogenesis pathways (acetoclastic, hydrogenotrophic, and methylotrophic).^{54,72}

Genus-level microbial community composition. For estimating the microbial dynamics across samples at the genus level, all the bacterial taxa with a relative abundance of 1% or higher (ESI Table 2(A)†) and archaeal taxa with a relative abundance of 0.5% or more are considered (ESI Table 2(B)†). The microbial community of the E(C2)Tx samples did not change much upon digestion for 40 days. This is evident from similar relative abundance of taxa at the genus level in the SP and EP of these samples. Nevertheless, in these samples, the relative abundance of genera belonging to Clostridiales is higher as compared to the other taxa with that of *Clostridium_sensu_stricto_1* being the highest in both SP and EP samples (Fig. 6). Earlier, the abundance of >90% Firmicutes belonging to class Clostridia has been observed in all the digesters with different total solid percentages of cow manure.⁷¹ *Clostridium* sp. is an obligate anaerobic bacterium that can utilize a wide range of carbohydrates (such as xylose, arabinose, galactose, glucose, cellobiose, sucrose, and fructose) from various renewable substrates for the production of H_2 with a notable yield of 2.1–2.2 mol H_2 per mol sugars.²⁹ The variations at the genus level between the SP and EP of E(C2)UTx and control samples are extremely high. The relative abundance of the genus *Pseudomonas* is the highest in the SP followed by the

EP of control samples and the SP of E(C2)UTx, whereas it is absent in E(C2)Tx samples (Fig. 6). *Pseudomonas* is of great importance for an efficient AD process and acts as a key player for hydrolysis, fermentation, and degradation of cellulose-rich biomass and all the important methanogenic pathways.^{22,73} The EP samples of E(C2)UTx are mainly dominated by uncultured members of *Spirochaetaceae* and in the EP of controls *Pseudomonas* is enriched and a significant rise in *Leptolinea* is observed. *Spirochaetes* and *Leptolinea* are acidogenic bacteria and associated with the metabolism of carbohydrates into acetate, whereas *Spirochaetes* are also involved in amino acid metabolism and production of H_2 and CO_2 .⁷⁴ The fact that the overall microbial community of the E(C2)Tx samples is not varying much throughout the experiment from SP to EP implies that this community is more stable over time and rich in cellulose-degrading microbes throughout the cycle. On the other hand, the microbial communities of SP and EP of the E(C2)UTx and control samples show drastic changes in their diversity and abundance of taxa. One of the prime reasons for the stable microbial community in the heat-pre-treated (E(C2)Tx) samples is the elimination of numerous microbes that cannot withstand harsh conditions due to the effect of heating.³⁹



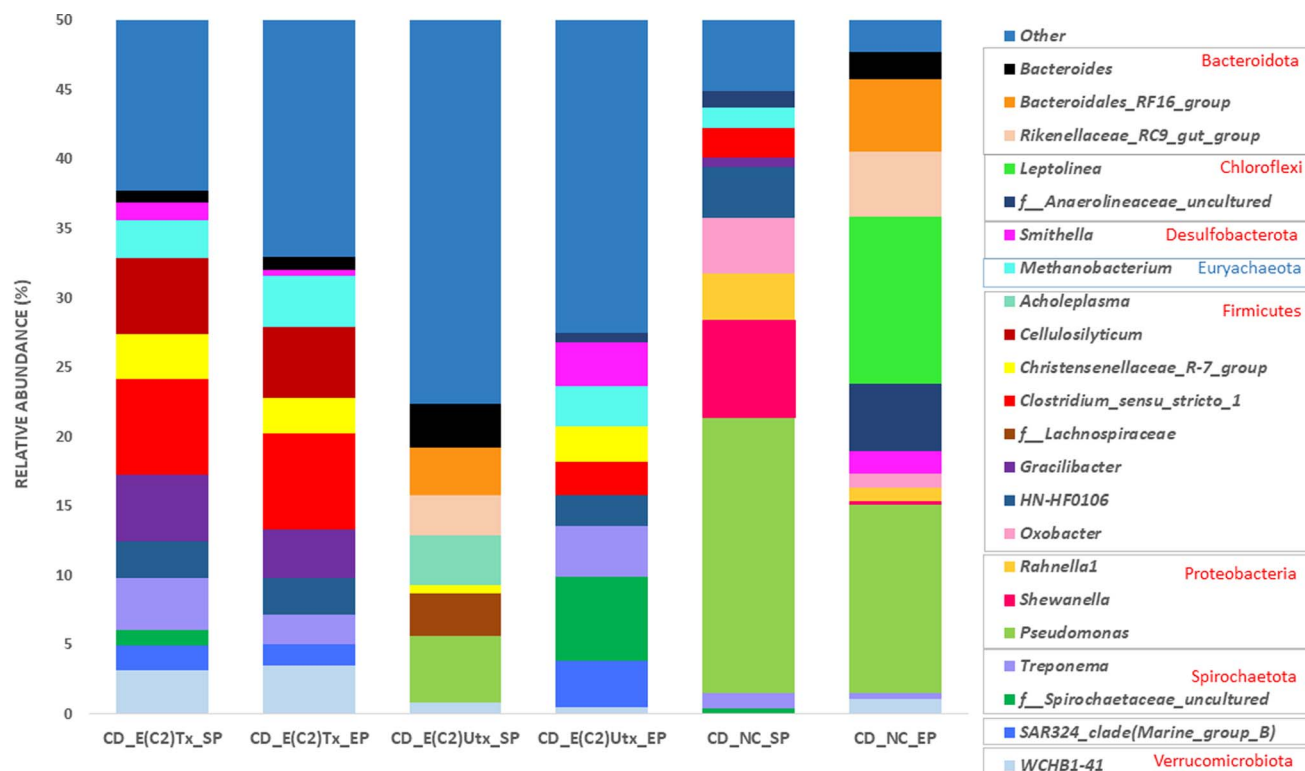


Fig. 6 Stack bar plots representing the relative abundance of the top five bacterial or archaeal genera. The samples were collected for each category at the starting (SP) and end (EP) points of the anaerobic digestion of CD for a period of 40 days. The remaining microbial communities are marked under the category of "Others". E(C2)Tx: heat-pre-treated inoculum, E(C2)Utx: untreated inoculum, NC: negative control.

Roles of microbial community during anaerobic digestion of cow dung to produce biohythane

The process of microbial AD consists of four phases, namely, hydrolysis, acidogenesis, acetogenesis, and methanogenesis, where bacteria mediate the first three stages and archaea carry out the final one (Fig. 7).

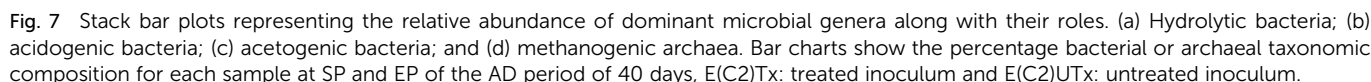
Hydrolysis. During hydrolysis the fermentative bacteria facilitate the hydrolysis of recalcitrant and complex organic matter (e.g., proteins, carbohydrates, and lipids) into soluble and biodegradable simpler substrates (e.g., monosaccharides, higher fatty acids, amino acids, and alcohols).⁷⁴ The hydrolytic fermentative bacteria (such as *Firmicutes*, *Bacteroidota*, *Proteobacteria*, *Chloroflexi*, and *Actinobacteria*) initiate this process by producing extracellular enzymes (such as cellulases, proteases, and lipases), which adhere to their cell wall and increase the digestibility of organic matter. The microbes involved in the hydrolysis process are mainly governed by the composition of the feedstock being used. CD as a feedstock has recalcitrant lignocellulosic content (50% in dry matter) and an inherent set of different fermentative microbes (such as species of *Bacillus*, *Corynebacterium*, *Lactobacillus*, *Pseudomonas*, *Citrobacter*, *Ruminococcus*, etc.).²² Therefore, the microbial members involved in the hydrolysis process are expected to be mainly rich in the cellulolytic bacterial community (including *Clostridium*, *Corynebacterium*, *Ruminococcaceae*, *Pseudomonas*, *Ruminiclostridium*, *Ruminococcus*, *Ruminofilibacter*, etc.).^{54,73–75} Overall, the abundance of cellulolytic microbes is the highest in the E(C2)Tx

samples as compared to the others (Fig. 7(a, b) and ESI Table 2(A)†).

There are other prominent differences in the dominant microbes involved in the hydrolysis process in the E(C2)Tx and E(C2)Utx samples as compared to the control (Fig. 7). At the genus level, the cellulolytic activity to degrade the lignocellulosic feedstock is mainly carried out by the genus *Clostridium* and several other members of *Firmicutes* in E(C2)Tx samples. In the E(C2)Utx samples, the cellulolytic and hydrolytic activities are carried out by the members of the genus *Pseudomonas* and several other members of *Bacteroidota*. In the control samples, this is mainly carried out by *Pseudomonas* and *Shewanella*. A few studies have previously reported the microbial community of the AD process inoculated with heat-shock pre-treated inoculum on various substrates.^{39,69,76,77} Corroborating with our results, a dominance of H₂ producers belonging to *Clostridium* sp. mainly *Clostridium_sensu_stricto_1* and other *Firmicutes* (*Prevotella*) and *Bacteroidota* (*Romboutsia*), etc., have been observed in those studies. In addition, they also observed *Enterococcus*, *Pseudomonas*, and a few less explored taxa including *Paraclostridium*, *Exiguobacterium*, *Aneurinibacillus*, *Ureibacillus*, *Xanthomonas*, and *Paenibacillus* which could be due to the use of other substrates.^{39,69,76,77}

Acidogenesis. The acid-forming bacteria (most commonly *Firmicutes*, *Clostridium*, *Bacteroidetes*, and *Enterobacter*) continue the fermentation process to metabolize the products of hydrolysis towards short-chain volatile fatty acids (VFAs, such





The H₂-production process can be hindered by several microbes including hydrogenotrophic methanogens, homoacetogens, nitrate-reducing, sulfate-reducing, iron-reducing, propionic acid-producing, and lactic acid bacteria. Propionogens or homoacetogens can produce propionic acid, which inhibits the dark fermentation process and reduces H₂ yield.⁷⁸ *Akkermansia* and *Alistipes* are the two propionate producers and *Candidatus_Falkowbact*, *Fusibacter*, *Caldisericum*, and *Sulfurospirillum* are sulfate and thiosulfate reducing bacteria which are mainly present in E(C2)UTx and control samples which can impact H₂ production adversely. Consequently, a reduction in H₂ production has also been observed in these samples as

compared to E(C2)Tx (ESI Table 2(A)†). Another important bacterial genera, namely *Acinetobacter* (*Proteobacteria*) showed a slightly higher abundance in E(C2)Tx than E(C2)UTx and was completely absent in control samples (ESI Table 2(A)†). Most of the members of this genera are pathogenic but play very important roles in hydrolysis, acidogenesis, and H₂S removal. In addition, *Acinetobacter* is also an effective phosphate solubilizer which makes it an important taxa for phosphorus nutrient enrichment in E(C2)Tx.

Acetogenesis. Few of the fermentative products (including acetic acids, H₂, and CO₂) produced during the acidogenic stage may be directly employed by methanogens for biogas production,⁷⁹ whereas the VFAs (such as butyrate, propionate, and valerate) are further consumed by H₂-utilizing acetogens or syntrophic acetate-oxidizing bacteria (SAOB) and converted into acetic acid, H₂, and CO₂ in the acetogenesis step.⁷⁴ During this step, the microbial community of E(C2)Tx samples is shifted towards the members of class *Clostridia* (including *Clostridium_sensu_stricto_1*, *Clostridium_sensu_stricto_6*, *clostridia-vadinBB60_group*, members of genus *Acetivibrio*, *Anaerovorax*) which are SAOB,⁷⁴ *Azospira*, *Bacteroides_UCG-001*, *Terrisporobacter*, *SG8-4*, *SAR324_clade* (*Marine-group-B*), and *DMER64* and other syntrophic bacteria including *Syntrophorhabdus*, *Syntrophus*, and *Trichlorobacter* (Fig. 7(c) and ESI Table 2(A)†).

Methanogenesis. Finally, acetic acid is consumed by acetoclastic methanogens to generate CH₄. Alternatively, H₂-consuming methanogens generate CH₄ from H₂ and CO₂ through the hydrogenotrophic pathway. In some cases, the methylotrophic methanogenesis pathway is followed in which CH₄ is produced from methyl compounds (e.g., methylamines, methanol, and methyl sulfides).^{54,74} The methanogenic community of E(C2)Tx samples is dominated by *Methanobacterium*, and *Bathyarchaeia* (Fig. 5, 7(d) and ESI Table 2(B)†). *Methanobacterium* belongs to the major methanogenic archaeal phylum *Euryarchaeota* and is a hydrogenotrophic or methylotrophic methanogen that lives in metabolic symbiosis with fumarate-reducing bacteria and H₂-producing cellulolytic bacteria.^{54,80} *Methanobacterium* is not affected by VFA concentration and has been observed in a syntrophic association with acetate oxidizers.^{54,81} The associations between H₂-producing bacteria (such as *Ruminococcaceae*) and hydrogenotrophic methanogens (*Methanomicrobiaceae*) are well known.⁸¹ In the presence of both *Ruminococcus* and *Methanobacterium*, acetate was found as the main fermentation product, H₂ did not accumulate, and a significant amount of CH₄ was formed.⁸¹ Towards this, the taxa *Ruminococcus*, uncultured members of the *Ruminococcaceae* family, and *Methanobacterium* are found in E(C2)Tx samples and their relative abundance was found to be increasing from SP to EP of the AD process.

In E(C2)UTx samples, the methanogenic community is mainly dominated by *SCGC_AAA011-D5*, *Methanobacterium*, and *Methanosaeta* among others, whereas the control samples exhibit a higher abundance of *Methanosaeta*, *Methanosarcina*, and *Methanospirillum*. *Methanosarcina* and *Methanosaeta* are expert taxa in the acetate cleavage process that produces methane, however, *Methanosarcina* grows best in environments with high acetate concentrations and encompasses both

acetoclastic and hydrogenotrophic pathways for methane production.^{82,83} *Methanospirillum* sp. is a member of the *Methanomicrobiales* of *Halobacterota* and is crucial for methanogenesis.⁸⁴

Methylotrophic microbes are known to play a lesser role in methanogenesis as compared to hydrogenotrophic and acetoclastic ones.⁷⁴ The known substrates for the methylotrophic pathway are methyl compounds, alcohols, and acetate. In addition, some of the methylotrophic microbes are strictly dependent on H₂ for their activity. At the same time, the available alcohol and acetate produced during acidogenesis might be consumed *via* the methylotrophic pathway for CH₄ production. These indicate the possibility of an active participation of the methylotrophic pathway also for methanogenesis in E(C2)Tx. It is known that acetoclastic processes contribute more towards CH₄ production.⁸⁵ Towards this, some acetoclastic microbes are observed in E(C2)Tx samples, such as *Methanosarcina*. However, a higher abundance of methylotrophic and hydrogenotrophic microbes are found for CH₄ production in E(C2)Tx samples (ESI Fig. 2†). This implies that due to the effect of heat treatment the methylotrophic and hydrogenotrophic pathways might be favored for methane production in E(C2)Tx. Corroborating with our results, in another study, acetoclastic and methylotrophic methanogens have been observed while performing thermal hydrolysis pre-treatment of protein rich tofu as the substrate.⁸⁶

Proposed mechanism of biohythane production *via* anaerobic digestion of CD using E(C2)Tx inoculum

The E(C2)Tx samples are found to be enriched with the H₂-producing and cellulolytic microbial communities throughout the experiment duration as compared to E(C2)UTx and controls. Due to this, an enhanced substrate degradation is expected to take place and subsequently, more H₂ will be produced till the end of the experimental cycle. A major hindrance in the retrieval of H₂ in AD is caused by the presence of H₂-consuming methanogens, as they use up all the available H₂ to form CH₄. It is important to note that the indigenous microbial community of the substrate CD is rich in methanogens. However, the fact that methanogens are very slow-growing microbes⁷⁴ and the microbial community in the E(C2)Tx samples is enriched in cellulolytic bacteria suggests that there may be a faster degradation of a substrate and subsequent H₂ production but a relatively slower consumption of H₂ for CH₄ formation in the initial phase of the experiment. Therefore, due to the high cellulolytic activity, the abundance of H₂ may be so much that even after the production of CH₄ by consuming the available H₂, sufficient H₂ is left which can be detected and recovered in E(C2)Tx samples. Subsequently, methanogens will start to adapt to the compounds available in the environment which include H₂, acetic acid, alcoholic compounds, methyl compounds, and others, and will start growing. This will lead to the enrichment of hydrogenotrophic, acetoclastic, and methylotrophic microbes during the AD process for CH₄ production. Interestingly, in the E(C2)Tx samples, a dominance of methylotrophic



and hydrogenotrophic communities was observed throughout the experiment (ESI Table 2(B)†).

The overall stable community composition throughout the AD process of 40 days in the E(C2)Tx samples also offers other advantages. For example, the microbial community in these samples has a negligible abundance of nitrifying and denitrifying microbes and sulfate- and thiosulfate-reducing microbes throughout the experiment. This implies that the process of ammonification and sulfur reduction is not predominant in this case. These processes are known to increase the alkalinity of the system, leading to a hindrance to the growth of H_2 -producing microbes and an overall disruption of the AD process, as well as the production of biogas.⁷⁴ Another factor that is mainly responsible for the cessation of the AD process is the accumulation of VFAs and CO_2 , which leads to an acidic environment and is known to hinder the growth of methanogenic and H_2 -producing microbial communities.⁵⁴ The methylotrophic pathway already plays an important role in the metabolization of alcohol and acetic acid. These observations imply that there exists an optimal balance between the shifting of the environment to excessively acidic or alkaline conditions, which can adversely affect the microbial community dynamics and prevent the community from shifting to the extreme.

The main driving mechanism behind efficient biohythane production from CD in this study is the optimized designing and pre-treatment of the synthetic microbial consortia – E(C2)Tx. For the designing of E(C2)Tx consortia, heat pre-treatment was used to selectively enrich the community with spore-forming H_2 -producers. Heat treatment can increase the activity of hydrolytic enzymes in the inoculum and can accelerate the degradation of complex organic compounds into simpler molecules, which can be more easily utilized by microbes for H_2 and CH_4 production.⁸⁷ Acclimatization of the inoculum's components under AD conditions allowed the microbial community to adapt to the new components such as CD, AS, PM, EF, and environmental conditions. Acclimatization promotes the development of synergistic interactions between various microbial species, improving the stability and overall metabolic performance of microbial consortia. During the acclimatization period, the microbial community becomes more tolerant towards AD inhibitors such as VFAs leading to more consistent biogas production.^{88–90} There are some studies that have adopted heat pretreatment and acclimatization of inoculum to enrich biohythane with H_2 . For instance, Rafieenia *et al.* (2018)⁹¹ used heat pre-treatment of inoculum as one of the methods to enrich H_2 -producing bacteria and performed the two-stage AD of food waste. However, food waste (FW) as a substrate is known to be rich in organic content (lipids, proteins, carbohydrates, starch, cellulose) which is very different from CD composition. In another study, Shi *et al.*, (2023)⁸⁶ demonstrated the use of heat pretreatment in combination with acclimatization, but they performed thermal hydrolysis pretreatment on a bulk substrate and acclimatization of inoculum. Besides, the substrate used by them was tofu, which is a highly protein-rich food product.

Suitability of the cow dung substrate for biohythane production

Among different substrates, carbohydrate-rich substrates are best suited for biohythane production, followed by those rich in proteins, while fatty acid-rich substrates are least suited.²⁹ For example, the H_2 production from FW is reported to yield 205 L H_2 per kg VS, whereas cellulosic wastes such as hyacinth yielded 51.7 L H_2 per kg VS.⁹² However, water hyacinth plants need harvesting, cutting/chopping, and pretreatment before AD and for FW continuous pH maintenance is required during the AD process to get H_2 and CH_4 because FW contains various inhibitory components which can cause instability of the system.²⁴

The substrates with a carbon/nitrogen (C/N) ratio of more than 20 are ideal for stable H_2 production, whereas a C/N ratio between 20 and 30 is observed as suitable for efficient CH_4 yield.^{92,93} The C/N ratio of CD is in the range of 16–25,⁹⁴ which means it is the best-suited feedstock for the production of both H_2 and CH_4 or even biohythane. Furthermore, in the biohydrogen production system, H_2 -producing bacteria need vital nutrients such as carbon (C), nitrogen (N), and phosphorus (P). The cellulose component of CD can act as a very good C source. In contrast, N can be supplied by proteins, nitrates, nitrites, and ammonium salts (also function as a buffer for organic acids) from the undigested organic matter of CD.

Phosphorus concentration plays a crucial role in the DF process by maintaining the buffer capacity of the system during any reaction.⁹² Although P is an extremely important nutrient for various life forms, including plants and animals, its ready availability is a great challenge.^{23,95} Recovery of phosphorus from CD is often limited by the extremely small size of phosphorus-containing struvite crystals.⁹⁶ Phosphate solubilizing microbes can hydrolyze organic and inorganic phosphorus into soluble forms.⁹⁷ Thus, phosphorus is likely provided in the form of phosphates in the AD process,⁹⁸ and its concentration is positively associated with H_2 production. In addition to this, CD from indigenous Indian cows is also enriched with certain inorganic metals, including zinc (Zn), copper (Cu), cobalt (Co), manganese (Mn), and traces of sulfur (S), iron (Fe), and magnesium (Mg), which are all required for microbial growth.²²

Economic perspective for biohythane production

A techno-economic analysis for the production of biohythane from wheat straw estimated the production cost as 23.04 \$ per kg biohythane.⁹⁹ The nutrient cost involved in this process was found to be the major cost driver. It was additionally highlighted that a reduction in the nutrient cost by 80% could lead to a decrease of 44% in the total production costs of the process. The production cost of biohythane was estimated previously as 7.20–48.96 \$ per kg biohythane using a two-stage dark fermentation and AD of potato steam starch.¹⁰⁰ The nutrients (yeast extract) required for both dark fermentation and AD were found to contribute the highest towards the total production cost. Subsequently, it was highlighted that using fewer or less costly nutrients for the AD process for single- or two-stage- DF and AD process would be critical for lowering the production



costs. Furthermore, it was also noted that the two-stage systems for biohythane production demonstrated a net positive energy balance than the single-stage H_2/CH_4 reactors while also considering the energy required for pretreatment and reactor heating.^{101,102}

For exploring the industrial applications of any technology, the capital investment estimates need to be explored. For setting up a two-stage AD process for biohythane production, an approximate amount of 12 687.7 € has been estimated for treating 13.4 tons of waste biomass (*i.e.*, 946.84 € per ton).¹⁰⁰ In Europe, a pilot experiment was set up in a two-stage thermophilic AD on food waste, which was found to generate an annual income of 540 874 € per year from 27 tons of daily waste feed (*i.e.*, 54.88 € of income per ton).¹⁰³ For any investments in this direction, a payback period varying from 2 to 6 years has been reported, which also depends upon the nature of feedstocks.¹⁰⁴ Governmental schemes and policies under the waste-to-energy theme have been known to play major roles in the proliferation and implementation of novel technologies for biohythane production in the market.

The economic viability and competitiveness of renewable energy sources, including biohythane, are significantly influenced by the cost of traditional fossil fuels, particularly crude oil.¹⁰⁵ Any fluctuations in the oil or fossil fuel prices on a global scale impact the economic sustainability of renewable energy sources by influencing the currency exchange rates, which will further affect the cost of energy import and export.¹⁰⁶ For an effective competition of biohythane as a future energy source with fossil fuels, despite the fluctuations in their costs, biohythane production technology needs to be economically viable. This emphasizes the need for more innovation to develop cost-effective strategies for the production of biohythane to ensure its financial sustainability.

Limitations of the study and future prospects

There are a few limitations of this study that may be addressed to enhance the commercial potential of the technology developed. The lab-scale experimental set-up with small working volumes, small study sample size, and the purity of reactants used limits the direct transferability of the findings of the study to the industrial settings. Another important challenge is a lower than required (>5–10%) amount of H_2 gas (3%) in biohythane composition recovered during the AD process. Furthermore, a parametric analysis was outside the main frame of this study, which might be one of the major factors for not attaining the appropriate range of biohythane composition.

Keeping these limitations in mind, a comprehensive standardization of the bioprocess and substrate parameters (such as COD, TS, VS, *etc.*) can be taken into consideration in future studies. Second, lab-based or small-scale experiments may not accurately represent the challenges and complexity of commercial-scale operations. Therefore, future research should focus on scaling up the experiments to increase working volumes, which may match the industrial demands. Third, co-digestion of the CD substrate with other wastes can be attempted to enhance the proportion of H_2 to meet the standard

biohythane composition. Previously, we have attempted co-digesting food waste, CD, and sewage wastewater mixed in an appropriate proportion by using the E(C2)Tx inoculum for enhancing the H_2 production.²⁴ However, the AD process requires continuous pH maintenance during the entire process, which is likely to increase production costs if applied at the commercial scale.

The cost-effective utilization of fermentative or AD residue is crucial for improving the financial performance of biohythane production. For future studies, the use of anaerobic digestate for generating various cost-effective and sustainable products, such as effective biofertilizers, biochar, bio-cement, bio-bricks, *etc.*, may be targeted. Converting anaerobic digestate into biochar or bio-cement not only addresses waste management issues but can sequester carbon and provide an alternative solution for soil health and fertility with new market opportunities.^{107,108} The future research may also focus on enhancing the efficacy of the system by providing key elements (C, H, N, and P) in addition to essential micronutrients (Fe, Ni, Co, Mo, Cu) in the form of nanoparticles for the growth of anaerobic bacteria.¹⁰⁹

Conclusion

The current study investigated the potential of synthetically designed microbial consortia for biohythane production using single-stage anaerobic digestion of CD waste. The E(C2)Tx inoculum generated biohythane with 3% H_2 and 36% CH_4 without requiring any pretreatment of the bulk substrate, thereby, proving to be more economically viable. In contrast the existing approaches heavily rely on cost- and labor-intensive bulk substrate pretreatment strategies limiting their commercial potential. The presented approach involved a pretreatment of the inoculum which is used in four times less volume than the bulk substrate and is made up of different wastes as well. The strategy for the pretreatment of inoculum included heat treatment and acclimatization, which resulted in a stable microbial community enriched in hydrolytic and cellulose degrading microbes as well as hydrogenotrophic and methylo-trophic methanogens, which have their own advantages in increasing H_2 and CH_4 yields by also reducing the cycle length of obtaining these gases. The presented technology shows promising cost-, resource- and energy-effectiveness, which can offer biohythane as a competitive alternative renewable energy source over the traditional fossil fuels, which also have their own environmental impacts. In addition to this, the developed technology for generating biohythane from CD presents a very efficient CD waste management strategy and a good example of waste to energy conversion supporting circular economy and sustainable development initiatives. Future research in this direction should focus on developing economic models for quantifying financial benefits and scaling up the process with comprehensive process parameters and open the avenues for its commercial adoption worldwide.



Data availability

Raw sequencing reads have been submitted to the NCBI Short Reads Archive (SRA) under BioProject accession number PRJNA828359, with respective SRA IDs (*viz.*, SRR18827896, SRR18906513, SRR18827912, SRR18828078, SRR18828097 and SRR18828098). These data can be accessed using the following link: <https://www.ncbi.nlm.nih.gov/sra/PRJNA828359>.

Author contributions

Rashmi Ira and Vikas Sharma designed the experimental setup, and performed sample collection, DNA extraction, and meta-genomic analysis. Shrawan Kumar and Tulika Prakash performed NGS library preparation and sequencing. Mira Koul assisted in metagenomic data analysis. Lalita Sharma carried out EF synthesis and Aditi Halder provided inputs and leads for EF production. Tulika Prakash conceptualized the problem, arranged resources, managed the project, and provided guidance for all experiments and computational analysis. Rashmi Ira and Tulika Prakash wrote the original draft and final manuscript. All authors have read and approved the final version of the manuscript.

Conflicts of interest

The authors have declared no conflict of interests.

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