Can the addition of biochar improve the performance of biogas digesters operated at 45°C?

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Abstract

Biogas from anaerobic digestion (AD) may contribute towards a green energy scene in the future. Despite many benefits, AD plant operators are still challenged by limitations of the technology. This study aimed at evaluating the effect of biochar addition to AD reactors operated at 45°C to optimize productivity, efficiency and stability of the process. Biochar is characterized by a large surface area and may therefore facilitate the creation of biofilms on the one hand and remove environmental contaminants on the other hand. Moreover, biochar may serve as pH control agent due to its alkaline properties, counteracting reactor acidification. Our results showed that the addition of biochar reduced efficiently the amount of H₂S that initially occurred in the headspace during reactor start-up. Nevertheless, no impact on biogas- and methane yields was observed. This might be explained with an already well-working AD system without any severe disturbances, where the balancing nature of biochar did hardly count. Future studies, however, need to evaluate the effect on stressed systems. The microbiome analysis provided a detailed view of the microbial dynamics during the start-up phase, revealing *Methanosarcina* as dominant methanogen in the end. The addition of biochar did not alter the microbial community composition.

Keywords: Anaerobic digestion, AD, Process optimisation, Methane, methanogens, 16S rRNA
1. Introduction

Anaerobic digestion (AD) is an established management and energy extraction technology for organic wastes and purpose-grown energy crops of various types under anoxic conditions by consortia of microorganisms to produce biogas [1, 2]. Although AD offers many benefits, the process presents some limitations as commercial AD plants typically operate below optimal performance levels [3, 4]. The limitations of AD can be broadly categorised into three groups: (1) the restriction on substrate types fed to the digester without pre-treatment, (2) process and product quality assurance, and (3) management of process effluents and emissions [5-10]. The preference of AD for wet organic waste requires that lignocellulose-rich agricultural biomass is pre-treated if high biogas yield is expected without sacrificing digester volume. This pre-treatment adds to the energy demand and can introduce inhibitors, leading to AD instability. The produced biogas requires further treatment before it can be used in all other applications aside from combustion. High specific volume, low pressure and low relative density characterise biogas obtained from an AD process. These undesirable characteristics with a high proportion of non-combustible carbon dioxide makes biogas unsuitable for energy storage. According to Budzianowski et al. [11], reduction in CO₂ concentration in biogas will reduce the required resources and scale of biogas upgrading systems.

Various attempts to improve the AD process stability and product quality have been made. Approaches such as biological, chemical, physical, mechanical, thermal and ultrasound techniques have been implemented [12-16]. More recently, bio-inspired materials have been investigated for their potential to facilitate microbial community clustering, and manage some of the indicators for process stability in AD. Among the varieties of bio-inspired materials is
biochar produced from the thermal decomposition of biomass in the absence of oxygen (pyrolysis). Biochar is a cost-effective material and an environmentally friendly approach for the production of carbon-based adsorbent with properties similar to activated carbon [17-19]. Carbon-based materials such as graphite and activated carbon have been considered to stabilise the AD process and increase the biogas yield [8, 20], but the production process of commercial-grade activated carbon is limited due to environmental-, health- and safety concerns [21]. This challenge is not a limitation for bio-inspired materials during biochar production and the subsequent in-situ application for AD as biochar has long been applied to soil for carbon sequestration. Biochar has the potential to effectively remove both organic and inorganic environmental contaminants as well as CO₂ from biogas due to its high surface area, stable structure, porosity, high ion exchange capacity and the presence of various surface functional groups (e.g. −OH, −COOH, C−O, O=C=O) [19]. An exhaustive review of the characteristics of biochar for AD and recent applications to AD has been presented by Masebinu et al. [10]. In the AD process, biochar has been applied mostly for pH control due to its alkaline nature, biofilm formation due to the porous structure and large surface area and for toxic compound inhibition [18, 22, 23]. Chen et al. [24] reported the ability of biochar to adsorb NH₃ in AD while its role in volatile fatty acid (VFA) management and CH₄ production was investigated by Sunyoto et al. [25]. In AD, biochar selectivity towards certain groups of parameters, however, cannot be controlled. Mumme et al. [26] reported that biochar likewise adsorbed inhibitors, nutrients and useful metabolites. Therefore, uncontrolled addition of biochar to AD can result in a negative interaction with the process. In a recent investigation, Shao et al.
[27] concluded that biochar effectiveness is limited to stressed AD processes and its effect on well-buffered systems is insignificant.

Among the observed parameters for the performance of biochar in AD is the operating temperature. The effectiveness of biochar tends to be higher in AD systems operated thermophilically (50-60°C). Elevated temperature offers higher metabolic rates, better pathogen inactivation, increased biogas- and CH₄ yields, as well as reduced foaming [1, 28, 29]. The high temperature can also accelerate leaching and dissolution of alkali metal cations from biochar, which improves the digestibility of substrate and endothermic adsorption of biochar [30-32]. Though the merits of thermophilic digestion are well outlined in the literature, yet the process requires intensive monitoring and microbes are very sensitive to fluctuation in operating temperature. AD at mesophilic temperature (30-40°C) is generally considered more stable, less sensitive to inhibitory components and lowers the inhibitory effects of long chain fatty acid [33] and ammonium. Moreover, it is characterized by a lower endogenous thermal energy demand [34, 35] and shows a significantly higher dewatering potential of the digestate [36], which is crucial since digestate handling is a major cost factor in an industrial-scale biogas facility. For these reasons, many biogas plants operate at mesophilic temperature. However, Hupfauf et al. [4], Tian et al. [37] and Hupfauf et al. [38] highlighted that the general acceptable temperature for mesophilic digestion at 37°C is not optimal due to a low substrate conversion rate and insufficient hygienisation. Instead, a preferable process temperature of 45°C was proposed by the authors that would combine the advantages of both, thermo- and mesophilic operation, accompanied by a promotion of members of the phylum Firmicutes with their superior capabilities to produce methanogenic precursors. The results of Hupfauf et al. [38] further
indicated that higher organic loading rates might be tolerated at this temperature without an inhibiting accumulation of VFAs particularly acetate. Hence, a process temperature of 45°C was chosen for this study, in order to contribute towards finding the optimum conditions for future AD systems.

Running a reactor at 45°C still has two major drawbacks: a prolonged initial lag phase and a comparably low CH$_4$ concentration in the biogas compared with colder temperatures. Biochar has been reported to reduce the lag phase and increase the CH$_4$ concentration at the same time. Moreover, it is assumed to promote clustering of microbes and to improve the process stability. Therefore, this research investigates biochar addition to lab-scale AD systems operated at 45°C, which is novel and has never been done before. Moreover, a strong focus lies on microbial community characterisation. Relevant research is scarce and urgently required since a properly working microbiota is crucial for a balanced AD process. So far, little data are available on microbial communities at 45°C process temperature with cattle slurry as main substrate, and none at all concerning biochar addition.

We hypothesise that (i) biochar addition increases the methane production in an AD system treating agricultural waste at 45°C, (ii) the addition of biochar facilitates the degradation of lignocellulose-rich biomass, and (iii) the addition of biochar shortens the initial lag phase typically found in AD at 45°C. Moreover, we hypothesise that biochar addition does not evoke a fundamental change of the microbiota in the bioreactor and that the whole consortium still works in a balanced way.

2. Material and Methods
2.1. Inoculum and Substrate

Fresh cattle slurry, collected at a local farm (Abenthum, Innsbruck, Austria) was used as start-up substrate and for all upcoming feeding events. Immediately after collection (November 6, 2018), the slurry was sieved (~ 2 mm) and stored at 4°C until use. Corn straw that was cut by hand into small pieces served as co-substrate. One gram of coarsely cut straw fibres was filled together with glass beads (to avoid floating) into mesh bags and the bags were then sealed with plastic clips (Laurel, Aichwald, Germany).

2.2. Biochar Production and Characterisation

The biochar was produced from corn stover biomass, which was obtained from MTN Bree market, Johannesburg, South Africa. The material was chopped to a particle size of 1 mm before pyrolysis. A tube furnace under slow pyrolysis condition of 600°C at a heating rate of 11°C min\(^{-1}\) with a residence time of 1 h was used. Nitrogen gas flowing at 230 mL min\(^{-1}\) was used to create anoxic conditions during heating. Proximate, ultimate and pH analyses were conducted in duplicate [39, 40]. The Brunauer-Emmet-Teller (BET) method for surface area, total pore volume and pore size determination by measuring of N\(_2\) adsorption using an ASAP 2460 instrument (Micromeritics, Georgia, USA) was implemented [41, 42]. The surface morphology of the biochar was determined using a VEGA 3 XMU scanning electron microscope (SEM) (Tescan, Czech Republic) equipped with an X-Max energy dispersive x-ray (EDX) analyser (Oxford Instruments, UK). For SEM, samples were mounted unto a stud using a double-sided carbon tape and analysed with an accelerating voltage of 15 kV.
2.3. Experimental Setup

Nine lab-scale biogas reactors [4] with a total volume of 2.35 L were used to assess the effect of biochar addition to an AD system. At the start of the experiment, the reactors were filled with 700 mL of sieved cattle slurry that was diluted 1:4 (v/v) with deionized water. Every three reactors were amended with 0 (reactors 1-3), 0.25 (reactors 4-6) and 0.5 g biochar g⁻¹ volatile solids (VS) (reactors 7-9), the reactors were shaken, and the straw-filled mesh bags were added. After sparging the headspace with pure nitrogen gas (purity 2.5, Messer, Germany) to establish anoxic conditions, the reactors were sealed and placed in an incubator set to 45°C. The reactors were then operated in batch mode over 22 d. Physico-chemical and microbiological characteristics were determined regularly. After the daily gas measurements, the reactors were manually shaken for 10 s to avoid sedimentation and to guarantee a homogeneous reactor content. The reactors with 0 g of biochar addition served as controls.

Through the data collection process, it was noticed that reactor 2 (one of the control reactors) could not hold pressure, possibly due to leakages caused by an improper alignment of the sealing gasket of the reactor. Hence, reactor 2 was completely excluded from the study and the results for the control group were thus calculated from duplicates rather than from triplicates.

2.4. Biogas Yield and Composition

The differential pressure (ΔP) between the headspace of each reactor against ambient pressure was measured daily with a digital manometer (GDH 200-13, GHM Greisinger, Germany). This differential pressure was used to calculate the biogas volume (\(Yield_b\)) according to Eq. (1) corrected to standard conditions (0°C, 1013 hPa). To account for the water vapour phase and
correct the biogas to dry conditions (which is beneficial when comparing gas yields with other studies), the temperature specific dry biogas factor (DBF) [43] was included:

\[
Yield_b [NmL] = \frac{\Delta P [hPa] \times \text{headsapce [mL]} \times DBF(T)}{1013 hPa}
\] (1)

Biogas quality (CH\textsubscript{4}, CO\textsubscript{2} and H\textsubscript{2}S) was measured daily except for the first day using an infrared-based biogas analyser (Geotech 5000, Geotechnical Instrument, UK). For each measurement, 100 mL of biogas was extracted from the headspace of each reactor immediately after the pressure measurement. Thereafter, the pressure in the reactor headspace was zeroed to avoid any over-pressure or vacuum in the digester at the beginning of the new measurement period. In the case of negative pressure, the headspace was flushed with pure N\textsubscript{2} gas (purity 2.5, Messer, Germany).

2.5. Physico-Chemical Analyses

Dry matter (DM) content, VS and the pH were determined twice a week. Fresh sludge samples from each reactor were dried overnight at 105°C (UF110, Memmert, Germany) and the weight loss was measured. For VS, the weight loss after incinerating the dried samples in a muffle furnace (CWF 1100, Carbolite, Gero, Germany) at 550°C for 5 h was calculated. The pH was assessed in freshly extracted reactor sludge with a glass electrode (Primatrode 6.0228.020, Metrohm, Switzerland) connected to a portal pH meter (826 pH mobile, Metrohm, Switzerland). Reactor alkalinity was determined at the beginning and the end of the study with 2-point titration to pH 4.4 and pH 5.0 using 0.1 N sulphuric acid (Merck, Germany). Based on the amount of acid added, the FOS/TAC ratio was then calculated according to Bensmann et al. [44] with Eq. (2):
To quantify the degradation of lignocellulose-rich straw fibres, the mesh bags were weighed at the beginning and the end of the experiment and the weight loss was determined. Prior to weighing the mesh bags at the end of the study, they were washed with deionized water and dried overnight at 105°C as described for DM. VFA and NH₄⁺ were quantified out of frozen (-20°C) sludge samples on a weekly basis. After thawing, the samples were centrifuged for 10 min at 20,000 g (Eppendorf 5804R, Germany) and the supernatant was diluted with distilled water: 1:2 (v/v) for VFA and 1:10 (v/v) for NH₄⁺. Samples were then sterile filtered through a cellulose filter with a pore size of 0.2 µm (Chromafil Xtra, Macherey-Nagel, Germany). VFAs were analysed on a high-performance liquid chromatography (HPLC) system (Prominence, Shimadzu, Japan) equipped with a Rezex RFQ-Fast Acid column (Phenomenex, California, USA). The HPLC analysis was done as described in Hupfauf et al. [4]. Ammonium contents were determined on the same HPLC system using the flow injection method [45] and a modified OPA/NAC reagent mix as described by Meseguer-Lloret et al. [46].

2.6. DNA Extraction and 16S Smplicon Sequencing

Microbial community analysis was conducted at the start and the end of the experiment. DNA was extracted with a NucleoSpin Soil extraction kit (Macherey-Nagel, Germany) out of frozen (-80°C) reactor sludge samples. The extraction protocol followed the instruction manual with some minor adaptations: 450-500 mg of sludge was poured into Bead Tubes and SL1 was used as lysis buffer. Sample lysis was accomplished with a FastPrep-24 device (MP Biomedicals,
California, USA) at room temperature, following the suggested settings (5 m s\(^{-1}\), 30 s). DNA elution was done in two steps with each 30 µL of PCR water (Rotipuran Low Organic Water, Roth, Germany). The final DNA extracts were stored at -20°C in low-binding reaction tubes (Eppendorf, Germany) until use. Gel electrophoresis (12 min, 100 V) on a 1% (w/v) agarose gel with Midori Green DNA dye (Nippon Genetics, Japan) was used to appraise the DNA extraction. Finally, DNA was quantified with a Nanodrop 2000C spectrophotometer (Thermo Fisher Scientific, Massachusetts, USA) and the samples were sequenced by a commercial company (Microsynth, Switzerland) using next-generation sequencing (NGS).

NGS was done on an Illumina MiSeq device targeting the bacterial/archaeal V4 region of the 16S rRNA (2 x 250 bp). The universal primers 515f (5’-GTGCCAGCMGCCGCGGTAA-3’) and 806r (5’-GGACTACHVGGGTWTCTAAT-3’) were used as suggested by Caporaso et al. [47] and NGS data analysis was done using the CoMA pipeline [48]. After assembling paired-end reads, primers and linker sequences were trimmed off. In the following quality filtering step, reads with an average Phred quality score < 30 (representing a base call accuracy > 99.9%) and/or a deviation > ± 5% compared with the mode sequence length were dropped from the analysis. Sequence alignment on a 97% level and the taxonomic assignment was done with the BLAST algorithm and the SILVA SSU (release 132) database. In cases where an operational taxonomic unit (OTU) could not be assigned to SILVA SSU, Greengenes (release 13_5) served as backup database. During post-processing, rare OTUs with < 5 occurrences over all samples were excluded and samples were randomly subsampled to a common depth of 19,048 reads after checking rarefaction curves. Both are commonly applied strategies in order to allow a proper comparison of the microbial communities in each reactor.
2.7. Statistical Analysis

Significance was determined at $p \leq 0.05$. Descriptive statistics, analyses of variance (ANOVA), Pearson correlations and post-hoc tests (with Bonferroni correction) were calculated with Excel 2016 (Microsoft, Washington, USA) and SPSS 24 (IBM, New York, USA). Principal component analysis (PCA) and redundancy analysis (RDA) were conducted with Canoco 5 [49]. Prior analyses, data were centred, standardised (only PCA) and log-transformed ($\ln(x+1)$). Shannon-Wiener diversity ($H'$), Pielou’s evenness ($J'$) and Chao1 richness were calculated based on archaeal and bacterial OTUs using the following Eq. (3), (4) and (5), respectively:

$$H' = - \sum_{i=1}^{R} p_i \cdot \log_2 p_i$$  \hspace{1cm} (3)

$$J' = \frac{H'}{H'_{\text{max}}}$$  \hspace{1cm} (4)

$$\text{chao1} = R + \frac{F_1(F_1-1)}{2(F_2+1)}$$  \hspace{1cm} (5)

where $p_i$ is the proportion of reads belonging to the $i^{\text{th}}$ OTU and $R$ is the samples OTU count. $H'_{\text{max}}$ is the maximum Shannon-Wiener index for a sample and can be calculated as follows:

$$H'_{\text{max}} = \log_2 R$$

$F_1$ is the number of singletons (OTUs with only one sequence), $F_2$ the number of doubletons (OTUs with only two sequences). Graphics were created either with the CoMA pipeline or with in-house Matlab- and Python scripts. In the taxonomic plots (Fig. 7, Fig. 8), OTUs with an abundance < 1% were excluded.

3. Results and Discussion

3.1. Substrate and Biochar Characteristics
Cattle slurry, maize straw and biochar were initially analysed in order to characterize all applied substrates in detail (Supplementary Table S1). The O/C ratio of the biochar indicates the extent of hydrophobicity of the biochar. The lower the O/C ratio, the less water can be absorbed by the biochar [50]. This is important as water absorption in an AD process reduces the interaction capacity of the microbial community with the substrate. Furthermore, the extent of hydrophobicity is a property of an adsorbent material indicating the capacity for CO₂ adsorption [51]. A lower ratio of H/C indicates higher stability of the biochar with highly carbonised and aromatic structures. It also signifies that possible organic content in the biochar has been devolatilised and therefore, the biochar will not necessarily degrade under anoxic conditions during the period of digestion [52]. Low O/C (0.28) and H/C (0.03) ratios in this study indicate the extent of aromatic condensation in the biochar [17, 50, 53]. The SEM images show the structural profile of the biomass before (a) and after (b) pyrolysis (Supplementary Fig. S1). The biochar surface area (0.49 m² g⁻¹) is very low compared to 3.19-29 m² g⁻¹ for non-activated biochar from corn stover [54, 55]. The pore size indicates that the pyrolysis process temperature facilitated the formation of meso-pores on the biochar surface. Micro-pores and meso-pores are preferred pore sizes on biochar surface to facilitate the creation of biofilms, promote the enrichment and the clustering of similar microbial communities [23, 50, 56].

3.2. Gas Measurement and Physico-Chemical Analysis

The biogas yield followed the similar trend of a long acclimatisation phase that characterises AD at 45°C as previously reported [4] (Fig. 1(a)). It was noticed that despite an increasing total biogas yield the share of CH₄ was low compared to CO₂ during the first 17 d of the experiment.
Biochar addition did not shorten this initial lag phase, neither at the low concentration nor at the higher one. This is contrary to what we had expected since we had assumed that the pH-balancing properties of biochar as well as its potential for creating biofilms might facilitate the microbial acclimatisation phase and reduce the duration until a steady state is reached. Obviously, the complex community structure including various syntrophic interactions, which is found at this temperature demands a certain time to establish, irrespective of the addition of a supporting agent. This however, needs to be proven also for higher biochar concentrations as well as for other substrate combinations. Particularly the latter may play a crucial role since highly specific microbial consortia are introduced to the process with the substrate. After day 17, the methane concentration started to increase rapidly, resulting in a CH$_4$/CO$_2$ ratio of 1.2 at the end of the experiment (Fig. 1(b)). As a result, the methane yield dramatically increased from day 17 onwards (Fig. 1(c)). Surprisingly, the addition of a low concentration of biochar negatively affected the biomethane yield by 10.2%, whereas the high concentration of biochar increased the methane yield by 3.3% compared to the control. However, these changes were not statistically significant and we assume that in this setup biochar addition does not affect the biogas production at all. This supports the results of Shao et al. [27], who concluded that the supporting properties of biochar are only decisive for stressed or inhibited AD systems. In undisturbed AD systems as in this study, however, only minor effects are to be expected.

Aside from the CH$_4$ concentration, the addition of biochar evoked a reduction of the H$_2$S concentration in the biogas by 23.3% and 37.5% for low and high concentration, respectively, compared to the control (Fig. 1(d)). The reduction of the H$_2$S was tested to be statistically significant (p = 0.002). We assume that biochar functions here as an effective sorbent agent, a
property that has already been reported for soil and other aqueous environments [57]. This is important since H$_2$S negatively affects human health and causes major problems in biogas facilities due to its corrosive properties damaging metal compounds of bioreactors, sensors, gas turbines and biogas upgraders [58, 59]. Our data show that biochar addition might be a potential approach for managing H$_2$S problems in AD. However, these findings need to be confirmed also for other substrates or substrate combinations.

Although, biochar addition lowered H$_2$S concentration in biogas, its effect in increasing the calorific value of the gas through increased methane concentration could not be supported because there was no significant difference in the CH$_4$ concentration between the treated reactors and the control reactor. Notwithstanding, several research results showed that the effect of biochar to AD is still not yet well understood due to divergent results. Shen et al. [60] reported that as soon as the biochar concentration increased in the AD, the accumulated biogas yield decreased and the methane concentration increased at the same time. Taking both factors together, the biomethane yield was similar to that of the control and no clear effect was found for the addition of biochar. Shao et al. [27] concluded that the addition of biochar to an already well-buffered system is insignificant but may show a positive effect for stressed AD systems. This is also supported by the results of Giwa et al. [61]. At low organic loading rates (1.02-1.62 g VS L$^{-1}$ d$^{-1}$), the performance of the control and the reactors with biochar were similar but as organic loading rate was increased beyond 1.62 g VS L$^{-1}$ d$^{-1}$, the control digester gradually acidified while the biochar-amended digester continued with stable biogas production. Taking all this together, we assume a general balancing effect of biochar at 45°C when added to AD systems.
disturbed by various reasons, including the accumulation of H₂S, NH₄⁺, or VFAs. For NH₄⁺- and
VFA accumulation, however, this still needs to be verified.

Among other parameters, the stability of an AD process can be verified by testing the pH,
ammonium, DM, VS, and VFA levels in the reactor sludge. The pH in all samples decreased as
the digestion process proceeded until day 17 (Fig. 2(a)); thereafter, the pH started to increase.
This can be explained with an ongoing accumulation of VFAs until this time point. Even though
the pH in the biochar-amended reactors was slightly higher, the trend was not significant. We
assume that biochar may indeed be used as a pH-balancing agent for AD reactors as suggested
by other authors [2, 18]; however, higher biochar concentrations as applied in the present study
might be required in order to gain a significant effect. Moreover, digesting other substrates than
cattle slurry and corn straw might lead to divergent results and need to be tested before making
general suggestions.

The ammonium concentrations in all samples were similar without any fluctuation
throughout the experiment (Fig. 2(b)). Furthermore, the ammonium concentration lay within the
expected range for cow manure (ca. 2 g L⁻¹) when considering the dilution in the presented
experiment. Contrary to our assumption, biochar addition did not affect the ammonium
concentration in any way (p = 0.588). This contradicts previous reports according to which
biochar is highly effective when it comes to the removal of inhibitors, particularly ammonium,
from biogas reactors [62-64]. Ammonium levels did not exceed an inhibitory concentration [65]
in this study and were hence not causing any process disturbances. However, further studies
investigating AD systems suffering from high ammonium loads are needed. In this respect, also
the substrate compositions needs to be considered as discussed for pH effects.
The DM content was significantly affected by the addition of biochar \((p = 0.005)\), but not by the factor “time” \((p = 0.406)\). Except for day 20, reactors amended with the high biochar concentration showed the highest DM content (Fig. S2(a)). However, no accumulation over time was observed for any of the investigated bioreactors, pointing towards a stably running system without overload or starvation. Contrary to DM, VS was neither affected by biochar addition \((p = 0.053)\) nor by the factor “time” \((p = 0.149)\) (Supplementary Fig. S2(b)).

Looking at VFAs shows that throughout the entire experiment, no butyrate was detected in any of the reactors (data not shown). The same was reported by Hupfauf \[4\] in a long-term experiment over 6-months at 45°C. In another long-term experiment, butyric acid was not detected at 35°C and only at a concentration of 8 mg L\(^{-1}\) at 45°C \[37\]. However, as temperature increased to 55°C, butyric acid levels increased to 131 mg L\(^{-1}\). It could not be ascertained if biochar facilitated the rapid reduction of butyric acid, however, it can be concluded that over a long term, digesting at 45 °C may eliminate butyric acid in the AD process. The acetate concentration gradually increased over time until day 17 after which it suddenly decreased below detection limit (Fig. 3(a)). The peak of acetate was recorded on day 17 corresponding to the sharp rise in the biogas production process. This is a strong indication that the AD process was first imbalanced until day 17, leading to a continuous accumulation of VFAs, particularly acetate. As soon as the microbiota fully adapted to the process conditions, the system recovered and all of the acetate was turned over into methane. Such an extensive initial lag phase was already reported before by others \[4, 37\], pointing towards a general characteristic of AD at 45°C. It was observed that biochar addition had no influence on the acetate concentration in the samples \((p = 0.987)\).
Propionate was not detected during the first week of the experiment, but it was found from day 10 onwards and reached a peak on day 17, again corresponding to the sudden rise in biomethane production (Fig. 3(b)). At the end of the experiment, a decline was noticed in the propionate concentration especially for the samples with high biochar concentration. The explanation for that might be the same as for acetate: as soon as a balanced and proper-working AD system has been established, accumulated VFAs were utilized and methane was produced out of it (either directly through acetoclastic AD or indirectly through syntrophic acetate oxidation and hydrogenotrophic AD). The addition of biochar did not show any significant effect on the propionate concentration in the reactors at $p < 0.05$ ($p = 0.621$).

The FOS/TAC gives information about the ratio of VFAs and the buffering capacity of the digester. Therefore, it is an important parameter for controlling the stability of a biogas reactor and indicates process disturbances caused for instance by organic overloading. All reactors showed considerably higher values at the beginning of the experiment than at the end (Fig. S3). The control had a 59.7% reduction, whereas the low and high biochar amended digesters showed a 64.3% and 65.8% reduction, respectively, compared to day 1. Biochar addition, however, did not show any significant effect ($p = 0.709$). This was unexpected since biochar is usually known to increase the buffering capacity of AD systems [2]. The insignificance of the buffer capacity of biochar in this study may be attributed to the already well-buffered AD system facilitated by the type of substrate used (cattle slurry). We assume that the amount of added biochar was not high enough to induce a measurable effect.

In order to quantify the degradation of lignocellulose-rich biomass, the weight loss of straw fibres was determined inside mesh bags. After 22 d, 60.2%, 61.8% and 58.5% of the straw
biomass were degraded within the control reactors, the reactors with a low biochar concentration
and the reactors with a high biochar concentration, respectively. At least in the short term, no
significant differences between the treatments were detected (p = 0.637). This is surprising and
contradicts previous reports of an accelerated degradation of fibrous biomass [66], even though
the authors suggested an optimum biochar concentration of 15 g L\(^{-1}\), which is approximately six
times and three times higher than seen for the low and the high concentration in this study,
respectively. This may at least partially explain the divergence of both studies; however, more
research is needed to investigate the degradation of lignocellulose-rich biomass at 45°C, since it
represents one of the major drawbacks of AD when treating agricultural residues.

3.3. Microbiological Analysis

NGS data showed that biochar addition did not affect the OTU richness (number of different
OTUs found in each reactor; p = 0.44) (Fig. 4(a)). The same was found for the Shannon-Wiener
diversity (p = 0.516; Fig. 4(b)), the Chao1 richness (p = 0.78; Fig. 4(c)) as well as for Pielou’s
evenness (p = 0.556; Fig. 4(d)). However, all parameters were influenced by the factor of time
(OTU count: p = 0.001, Shannon-Wiener diversity: p = 0.001, Chao1 richness: p = 0.001,
Pielou’s evenness: p = 0.001) indicating an adaptation of the microbiota from the start of the
experiment to the end, which had been expected after analysing the physico-chemical parameters.
All diversity measures were significantly lower after 22 d of operation, pointing towards an
ongoing specialisation of the microbiota towards a community that is perfectly optimized to
thrive (and produce biogas) at the prevailing process conditions.
To compare the microbial communities in all reactors with each other, a PCA was conducted (Fig. 5(a)). Samples from day 1 are located on the right side, those from day 22 on the left side of the scatter plot. This means that the factor “time” had a huge impact on the microbial composition, particularly since PCA axis 1 explained a high proportion of the total variation (44%). Samples from the start of the experiment (day 1) were distributed along the y-axis, whereas the reactors with biochar addition were closer related to each other than to the control reactors. This trend, however, is minor, since PCA axis 2 only explains 6% of the total variation. On the other hand, samples from the end of the experiment (day 22) were clearly clustered and no deviation was detected. This clearly strengthens the assumption of an ongoing specialisation of the microbiota to the process conditions, being the same for all investigated bioreactors, irrespective of biochar addition. To prove this also statistically, an RDA with forward selection was applied. The factor “time” explained 74.1% of the distribution, whereas the biochar addition only did 2.6%.

The same conclusions can be drawn when looking at the PCA computed solely from archaeal data (Fig. 5(b)). Again, the factor “time” evoked the majority of the variation, whereas the addition of biochar had little to no impact. When going into detail, it turned out that *Methanobrevibacter* and an unknown Woesearchaeia genus were most important for characterizing the microbial community of day 1. At the end of the experiment (day 22), however, *Methanosarcina* and *Methanoculleus* were found to be the characteristic methanogens. Despite the trend being very weak, *Methanosphaera*, as well as Candidatus *Methanoplasma*, were indicators for the samples amended with biochar.
To compare the similarity/dissimilarity of the samples for each taxonomic level, Venn plots were constructed. Between day 1 and day 22, reactors shared a large proportion of phyla (31 out of 36, 86%), classes (50 out of 62, 81%), orders (100 out of 120, 83%), families (161 out of 208, 77%) and genera (256 out of 344, 74%) (Fig. 6(a)). The other taxa were found either exclusively at day 1 or exclusively at day 22, whereas none of them predominated. This shows that there was a relatively large core microbiota during the whole duration of the experiment, at least when looking solely on the presence/absence of taxa. When comparing the two time points based on abundances, however, microbial communities were clearly distinguishable as shown with PCA. Taking this together led to the assumption that the distinction between both groups was mainly based on a small number of organisms that occurred at high abundances whereas the majority of bacteria/archaea was only detected at a small abundance, not decisive for distinguishing between the groups.

When comparing the different treatments (Control, Low conc., High conc.) at day 22, Venn plots again revealed a huge proportion of shared taxa for each taxonomic level (phylum: 28 out of 33, 85%; class: 43 out of 54, 80%; order: 86 out of 109, 79%; family: 136 out of 184, 74%; genus: 193 out of 298, 65%) (Fig. 6(b)). As discussed above, this points towards a large core community when considering the overall presence/absence of taxa. In this case, however, the proportion of shared genera was considerably lower as when comparing the two time points. This is particularly interesting since the PCA and RDA results clearly indicated that the factor “time” was much more important for shaping the microbial community structure than the addition of biochar. We explain this again with the presence of a few, highly abundant organisms (a member of the core microbiota) that are dominating the microbial community irrespective of
the addition of biochar. Apart from the shared taxa, the Venn plots showed that the reactors amended with a high concentration of biochar included slightly more unique taxa than the controls and the reactors amended with a low biochar concentration (e.g. at genus level: control: 15; Low conc.: 11; High conc.: 22). This trend, however, was minor and may not be generally valid for other similar experimental setups.

A deeper analysis of the bacterial community in the reactors on family level revealed that at the beginning of the experiment, Ruminococcaceae, Rikenellaceae, Dysgonomonadaceae and “Family XI” (order Clostridiales) dominated (Fig. 7). All four groups together included 46% of all detected reads (Control: 49%, Low conc.: 44%, High conc.: 44%), clearly corroborating the above assumption of a microbiota heavily dominated by only a small group of organisms. This is common for AD systems treating cattle slurry and was already reported before [38]. Irrespective of biochar addition, Ruminococcaceae were the most abundant family at the start of the experiment (as well as at the end; 16% of all reads). This is in line with various other studies describing the family as members of the AD core microbiome, particularly with animal wastes as main substrate [67-69]. Ruminococcaceae are important hydrolytic bacteria mainly degrading cellulose and hemicellulose to acetate and other VFAs [70]. Rikenellaceae is another important family frequently found in AD systems [67, 71]. It typically occurs in the gastrointestinal tract of animals and humans and thrives anaerobically. Glucose and other carbohydrates can be converted fermentatively to VFAs, alcohols, H₂ and CO₂ [72], and Nakasaki et al. [73] reported that members of this family might also be involved in the degradation of glycerol. In this study, Rikenellaceae were almost exclusively found at the beginning of the experiment, while their abundance dramatically decreased until day 22. Dysgonomonadaceae are other common
members of AD systems [71], thriving preferably between 35°C and 45°C. They utilize carbohydrates and peptides and ferment it to mainly acetate and propionate [74]. The pseudo clade “Family XI” includes various genera, for which a precise taxonomic assignment was not possible. At the start of the experiment, mainly the two genera *Tissierella* and *Sedimentibacter* were detected. *Tissierella* are obligatory anaerobic and grow optimally at 37°C. Moreover, they are non-fermentative, and degrade glucose and/or proteins down to acetate, propionate, butyrate, ammonia and CO₂ [75, 76]. *Sedimentibacter*, on the other hand, are involved in the acetogenesis step, where VFAs are degraded to acetate and H₂ [77, 78]. Wang et al. [79] reported that the organisms are vulnerable to high ammonia loads. This may explain their occurrence in this study, where the main substrate cattle slurry was diluted and hence ammonium loads were reduced, whereas in previous studies using the same (but undiluted) substrate and reactor setup, *Sedimentibacter* was absent [4, 38]. After 22 d, “Family XI” members were only detected at very low numbers anymore. We assume that this might be explained at least partially with the increasing ammonium levels over the duration of the experiment.

At the end of the experiment (day 22), Ruminococcaceae still dominated the microbial communities, irrespective of the addition of biochar (18-21% of all reads). The second most abundant family, however, was Burkholderiaceae, which was detected only at very low counts at day 0 (1-2%). At day 22, 12% of all reads were assigned to this family within the control reactors, whereas even higher abundances were found when biochar was added (Low conc.: 18%, High conc.: 17%). Burkholderiaceae is a very diverse family in terms of both, ecological and physiological characteristics. The most abundant genus in our reactors was *Comamonas*, which was also detected by other studies in AD reactor sludge [80-82]. Generally, this group is known
for being involved in denitrification processes [83]; in AD systems, however, *Comamonas* are particularly important for the degradation of a broad range of organic acids during acidogenesis and acetogenesis [77, 84]. Apart from *Comamonas*, *Parapusillimonas* and *Verticia* were the most important genera among the Burkholderiaceae family, although at considerably lower abundances. However, so far, only little information is available on their precise role in AD; even their broad occurrence in other bioreactors or AD systems is not well documented yet. *Heliobacteriaceae* is another frequently detected family at day 22, which was not found at all at the beginning of the experiment. So far, only little is known about their role in artificial biogas reactors; however, recent studies found them in rice field soil [85], paddy soil, and flooded desert soil [86]. Peng et al. [86] proved with their metatranscriptomic-based study that *Heliobacteriaceae* are involved in syntrophic acetate oxidation (syntrophic acetate oxidation bacteria, SAOB), where methane is produced from acetate in association with hydrogenotrophic methanogens. In addition, Noll et al. [87] speculated that *Heliobacteriaceae* might be involved in the degradation of propionate in rice soils when the temperature is high. This is reflected in our data, where the appearance of *Heliobacteriaceae* at the end of the experiment came along with a considerable decrease of both, acetate and propionate in the reactor sludge.

Looking at Archaea showed that over the whole experiment the vast majority of all reads was assigned to *Methanosarcina* (Fig. 8). Even though the counts were still very high, there was a decrease from the start of the experiment (Control: 98%, Low conc.: 92%, High conc.: 92%) to the end (Control: 83%, Low conc.: 75%, High conc.: 84%), whereas the addition of biochar showed no significant influence. A clear domination of *Methanosarcina* at 45°C is known from previous studies [4, 38], pointing towards highly preferable growth conditions for these
methanogens, at least when cattle slurry and straw are used as main substrates. Physiologically, the organisms are highly versatile, producing methane from various substrates. *Methanosarcina* are the only methanogens capable of using both, the hydrogenotrophic as well as the acetoclastic pathway. Which pathway is active is generally determined by environmental factors such as pH and the ammonium content. Jiang et al. [88] reported that hydrogenotrophic methanogenesis prevails at high ammonium levels, whereas acetoclastic methanogenesis is preferably used at low concentrations. The relatively high ammonium loads in our reactors (typical for reactors fed with animal manure/slurry) together with the numerous occurrence of SAOBs suggests that most of the methane is produced via hydrogenotrophic methanogenesis by *Methanosarcina* in association with syntrophic Heliobacteriaceae species. The addition of biochar had no influence and did not affect the production of methane, neither in a positive nor in a negative direction. Apart from *Methanosarcina*, low numbers of *Methanobrevibacter* and *Methanoculleus* were detected. The first occurred more at the beginning of the experiment (3-4%), whereas the latter appeared more towards the end (5-10%). However, their abundances were small and we assume only a minor contribution of these genera to the total amount of produced biogas.

Summing up, the microbiota in the bioreactors underwent an adaptation process from the start to the end of the experiment. A firstly Rikenellaceae- and Dysgonomonadaceae-dominated consortium shifted towards a community, where Burkholderiaceae and Heliobacteriaceae were highly abundant. The addition of biochar did not show an effect on the microbial community composition when looking at the core microbes; however, less abundant taxa might be affected. Our results demonstrated that stable biogas production was possible, irrespective of biochar addition. We attribute this to the presence of a balanced and properly working microbiota in all
reactors. Nevertheless, further studies are needed on stressed or imbalanced AD systems, where we expect a higher impact of biochar addition, also in terms of microbial community composition. Biochar might indeed function as a balancing agent, allowing for satisfying methane yields even under unfavourable process conditions. Moreover, studies on continuously operated bioreactors with high loading rates are needed in order to reveal the immobilizing potential of biochar for process-relevant microbes [89], particularly slow-growing organisms like methanogens [90].

4. Conclusions

This study monitored the start-up phase of lab-scale bioreactors fed with cattle slurry and corn straw at 45°C, and tried to reveal the effect of biochar addition to the system. Our data showed that biochar had no influence on both, amount and composition of the produced biogas. Likewise, none of the observed stability indicators was affected, except for a reduced H₂S content at the beginning of the experiment. Further studies are needed to prove our data also for the long term in continuously running AD systems. Moreover, future studies should focus on disturbed systems (e.g. organic overload, H₂S problems or acidification) in order to fully reveal the potential of biochar as balancing agent in AD reactors operated at 45°C. Our microbiome analysis gave vivid insight into the bacterial/archaeal dynamics during reactor start-up. As expected, the addition of biochar did not evoke considerable changes in the microbiota. Thus, for well-balanced reactor systems biochar does not offer operational advantages, if it does in unbalanced systems still needs to be investigated.
**Data Deposition**

All NGS data are uploaded to the European Nucleotide Archive under the accession numbers ERS3858284 to ERS3858301 within the study PRJEB34762 ([https://www.ebi.ac.uk/ena/data/view/PRJEB34762](https://www.ebi.ac.uk/ena/data/view/PRJEB34762)).

**Acknowledgements**

This work was supported by the Austrian Agency for International Cooperation in Education and Research (OeAD) within the “Scientific & Technological Cooperation” program. We thank the Abenthum family for providing cattle slurry, and Thomas Klammsteiner and Anna Winkler for their highly appreciated assistance during slurry collection and sieving.

**Author Contribution**

S.M. (Postdoc) created the experimental design, conducted the experiment and wrote the manuscript. O.F. (Ph.D. student) helped interpreting the NGS results and revised the manuscript. H.I. (Professor) created the experimental design and revised the manuscript. C.M. (Professor) revised the manuscript. A.W. (Associated Professor) contributed the VFA analysis and revised the manuscript. R.M. (Ph.D. student) contributed the NH$_4^+$ analysis and revised the manuscript. S.H. (Postdoc) created the experimental design, conducted the experiment, carried out the NGS data analyses with CoMA, performed statistical tests, wrote the manuscript and conducted the revision of the manuscript.
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**Fig. 1.** Accumulated biogas production (a), CH$_4$/CO$_2$ ratio (b), accumulated methane yield (c) and hydrogen sulphide content (d) of lab-scale bioreactors operated over a period of 22 days at 45 °C. Different colours indicate the addition of low (0.25 g g$^{-1}$ VS) and high (0.5 g g$^{-1}$ VS) concentrations of biochar. Reactors without biochar addition served as control. All plots show the means ± standard deviation (n = 3).
Fig. 2. The pH (a) and ammonium levels (b) in lab-scale bioreactors operated at 45°C over a period of 22 days. Different colours indicate the addition of low (0.25 g g\(^{-1}\) VS) and high (0.5 g g\(^{-1}\) VS) concentrations of biochar. Reactors without biochar addition served as control. The plot shows the means ± standard deviation (n = 3).

Fig. 3. Acetate (a) and propionate levels (b) in lab-scale bioreactors operated at 45 °C over a period of 22 days. Different colours indicate the addition of low (0.25 g g\(^{-1}\) VS) and high (0.5 g g\(^{-1}\) VS) concentrations of biochar. Reactors without biochar addition served as control. The plot shows the means ± standard deviation (n = 3).
Fig. 4. OTU richness (a), Shannon-Wiener diversity \( H' \) (b), Chao1 richness (c) and Pielou’s evenness \( J' \) (d) calculated from 16S sequencing data. Different colours indicate the addition of low (0.25 g g\(^{-1}\) VS) and high (0.5 g g\(^{-1}\) VS) concentrations of biochar. Reactors without biochar addition served as control. The plot shows the means ± standard deviation (n = 3).

Fig. 5. Principal component analysis (PCA) calculated based on bacterial and archaeal (a) as well as solely on archaeal (b) 16S sequences. Different colours indicate the addition of low (0.25 g g\(^{-1}\) VS) and high (0.5 g g\(^{-1}\) VS) concentrations of biochar. Reactors without biochar addition served as control. Different markers indicate the time point.
Fig. 6. Venn plots for comparing the microbiota at the beginning and end of the experiment (a) as well as to reveal the effect of biochar addition after 22 days of operation (b).
Fig. 7. Heatmap depicting the bacterial community composition of lab-scale bioreactors on family level based on 16S rRNA amplicon sequencing data. The plot shows mean values (n = 3) for two different time points and different amounts of biochar addition. Control: 0 g g\(^{-1}\) VS. Low conc.: 0.25 g g\(^{-1}\) VS. High conc.: 0.5 g g\(^{-1}\) VS.
**Fig. 8.** Heatmap depicting the archaeal community composition of lab-scale bioreactors on genus level based on 16S rRNA amplicon sequencing data. The plot shows mean values (n = 3) for two different time points and different amounts of biochar addition. Control: 0 g g⁻¹ VS. Low conc.: 0.25 g g⁻¹ VS. High conc.: 0.5 g g⁻¹ VS.