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Monitoring the Physiological State in the Dark Fermentation of Maize/Grass Silage Using Flow Cytometry and Electrooptic Polarizability Measurements

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Abstract

Dark fermentation (DF), a key biohydrogen-producing process, is generally operated as a *black-box*, by monitoring different operative macroscopic process parameters without evaluating or tracking the physiology of the biotic phase. The biotic phase in DF is constituted by a large variety of microorganisms, mainly fermentative bacteria. The present study uses two (electro)optical techniques, flow cytometry (FC) and frequency-dependent polarizability anisotropy (FDPA) measurements, to gain insights into the physiology of open mixed consortia throughout the DF process. The mixed consortia for DF were obtained from a methanogenic sludge, selecting spore-forming bacteria by means of an acid treatment. Then, DF systems with and without pH control were studied, using as substrate a mixture of maize and grass silage (9:1 w/w). Over the course of fermentation, the butyric pathway was dominant in both systems, and relevant titers of acetate, formate, and ethanol were detected; while hydrogen yields amounted to 20.80 ± 0.05 and 17.08 ± 0.05 NmL/gVS under pH-regulated and non-regulated conditions, respectively. The cytometric pattern analysis of the culture together with microscopic observations made it possible, over the course of fermentation, to identify and track the predominant morphologies in play (i.e., free spore, rod-shaped, and endospore, which are typical of *Clostridium* spp.). Furthermore, the use of the fluorescent dye DiBAC₄(3) in FC and FDPA measurements provided similar information regarding the physiological state (PS) of the mixed consortia during the different phases of the culture.

Keywords Dark fermentation · Biohydrogen · Physiological state · Flow cytometry · Frequency-dependent polarizability anisotropy

Introduction

The shift from oil-based economies to circular bioeconomic models is a growing concern of the biotechnological industry. In particular, the industrially feasible exploitation of different biomass resources and the vast microbial diversity is an

important pillar for the production of bioenergy and biochemicals. Europe accounts for 224 biorefineries [1] that use various feedstock, such as bio-waste, lignocellulosic material, waste fats and oils, and sugar- and starch-rich residues, either for bioenergy or bio-based chemical production.

Dark fermentation (DF) and anaerobic digestion (AD) are consolidated anaerobic fermentative processes that are used for bioenergy (H₂ and CH₄) and the production of biochemicals from organic waste or to stabilize activated sludges of wastewater treatments. Besides DF, biohydrogen can also be produced using photofermentation or other photobiological processes [2].

The DF and AD exploit open mixed cultures, composed of microorganisms belonging to the Archaea and Bacteria domains, where different biochemical cycles take place. The AD can be described as a concatenated succession of the following different phases: hydrolysis, acidogenesis, acetogenesis, and methanogenesis, while DF can be seen as a truncated version where methanogenesis does not take place.

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Fermentative bacteria are the main responsible organisms for hydrolysis and the two fermentative conversion stages (acidogenesis and acetogenesis), while methanogens are the terminal players in the trophic chain [3], converting the intermediates to the final products, CH₄ and CO₂.

The composition of the mixed cultures used in DF and AD has different proportions of hydrolytic fermentative bacteria that first hydrolyze (a) large macromolecules, such as polysaccharides, into small chain carbohydrates, (b) proteins into amino acids, and (c) lipids to glycerol, long-chain and short-chain carboxylic acids. In the first fermentation phase (acidogenesis), hydrolysis products feed different bacterial species, which in turn yield liquid-phase metabolites of interest, such as lactate, butyrate, propionate, acetate, ethanol, and gas-phase H₂ and CO₂. Thereafter, the second fermentation (acetogenesis) converts the primary fermented products into acetic acid, H₂ and CO₂, which are finally used in methanogenesis by hydrogenotrophic methanogens (which consume CO₂ and H₂) and acetoclastic methanogens (which consume acetic acid) and other substances (methylated compounds), to a lesser extent [3].

The DF can be conducted at different scales, using different types of bioreactors, such as continuously stirred tank reactors (CSTR), up-flow anaerobic sludge bed (UASB) reactors, anaerobic filter (AF) reactors, and fixed-bed reactors (FBR) [4]; each configuration presents particular features and dynamics, biohydrogen yields, and dominant bacterial species [5]. The DF can also be exploited in cascade reactor arrangements, for example coupling DF to AD in two-stage AD (TSAD) [6]. The TSAD optimizes process parameters to produce H₂ and a mixed pool of carboxylic acids in the DF bioreactor (fermentative bacteria), while a second methanogenic bioreactor (fermentative bacteria and methanogens) is fed with the liquid output from the first reactor and targets CH₄ production [7]. Alternatively, the liquid effluent of carboxylic acids and solvents from the DF process can be fed into different processes (see Fig. 1), thus increasing value addition and improving the sustainability of the process [8, 9]. This can be achieved by additional energy production as it occurs in microbial fuel cells (MFCs) [10], algae and microalgae cultivations [11], and other multi-cascade reactor configurations [7], as well as in the synthesis of valuable chemicals [12].

Despite the advantages of the use of biohydrogen as clean fuel, there are certain limitations that have prevented production on a consolidated and cost-effective industrial scale. The photofermentation and photobiological processes require very specific cultivation conditions and have low efficiencies of light conversion and relatively low H₂ yields [2, 13]. Enzymatic systems suffer from sensitivity to O₂ [8, 14] and H₂, which reduces the efficiency of these systems, although novel approaches are actively being researched to overcome this drawback [15]. Pure fermentative cultures (natural or engineered strains) require high sterilization costs that are

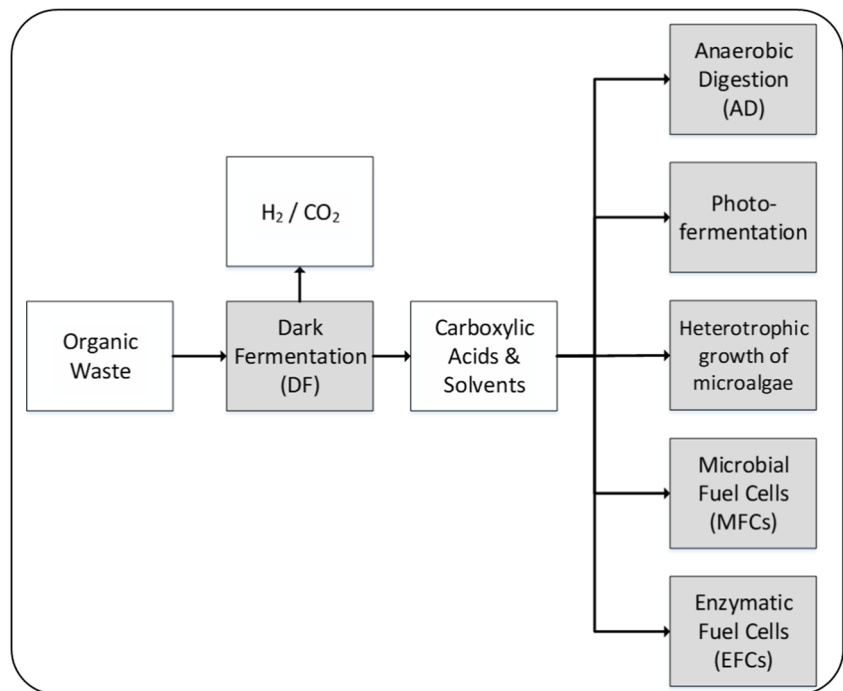
currently unsustainable [9] and can lead to a negative net energy balance [8]. Open mixed fermentative cultures, such as those used in DF, are preferred due to their high production rates and their simplicity of operation [15]. However, they require elevated power input (mixing) to overcome thermodynamic inhibitions for high H₂ productivity, and they are susceptible to metabolic shifts in long-term operated systems (e.g., solventogenesis, methanogenesis) [7].

Due to the inherent complexity and the intricate metabolic networks of mixed cultures in DF and AD, these bioprocesses are often operated as “black-boxes” and are managed through heuristics, empirical knowledge, and macroscopic process parameters [7]. Different molecular techniques are available to decipher the *black-box*, for example by profiling the bacterial communities involved in these processes. These methods can serve to better understand the ecological relationships among the community members or to construct detailed metabolic pathways or genome-based models. However, they are currently expensive, require laborious and time-consuming sample preparations [16], and, therefore, are not suitable for real-time monitoring at industrial level for DF and AD. Despite the large uncertainties of molecular tools [17], bacterial and archaeal communities in DF and AD have shown similar taxonomic and functional compositions, either in laboratory or industrial scale [18]. A study of 20 different laboratory-scale DF bioreactors operated in four different countries shows that the Firmicutes phylum [19] plays a predominant role as key fermentative microorganisms (mainly belonging to the Clostridia and Bacilli classes); while other phyla, such as Proteobacteria, Bacteroidetes, and Chloroflexi, are found in a lesser proportion. An overview of the AD microbiome is presented in [20], which analyzes 134 samples from both laboratory- and full-scale biogas plants fed with a range of different substrates and located in seven different countries; the results indicate that bacterial phyla Firmicutes, Bacteroidetes, Coprothermobacter, Actinobacteria, Termotogae, and Chloroflexi are the most abundant and that the methanogenic community is almost exclusively represented by the phylum Euryarchaeota.

Besides profiling bacterial communities and functional relationships, new process analytical technologies (PATs) can serve to develop process monitoring and control strategies, following the principle of quality by design (QbD). This is of particular interest for the development of advanced monitoring systems in biorefineries [21]. The PATs have proven their potential in measuring critical process attributes, mainly for the pharmaceutical industry and biotechnological processes with monocultures [22], which often include monitoring the physiology of the biotic phase. Nevertheless, this potential can also be exploited in complex microbial cultures to provide insights into key features of these systems.

As early introduced by Malek [23], the physiological state (PS) of the biotic phase is influenced by the physicochemical

Fig. 1 Possible process integration for dark fermentation (DF)



conditions of the environment and can either be a qualitative measure or be assessed through auxiliary parameters to monitor the process performance. The PS can thus be considered as a fitness indicator of culturing systems for their correct function and successful operation, which is a concept applicable to monocultures or mixed consortia. Hence, the compilation of key features of the biotic phase can provide better insights regarding the physiological state, such as morphology (shape and size of cells), the state of nucleic acids, the functionality of certain enzymes or redox cofactors, the status of transport mechanisms, and the membrane potential, among others [24].

Recent developments in (electro)optical techniques have allowed developing versatile PATs, which can be used for monocultures or microbial communities, even if microbial communities exhibit a higher level of complexity. One important platform, which is available for high throughput analysis of cells, is flow cytometry (FC). FC is a laser-based technique, which collects optical characteristics of cell suspensions, such as light scatter and fluorescence. Microbial morphological heterogeneities can be examined, in terms of relative cell size and granularity, by using the forward scatter (FSC) and side scatter (SSC) signals. Additionally, fluorescence detectors can further collect information of qualitative and quantitative nature, regarding the PS of several thousand cells per second, using specific dyes (bound to certain molecules and/or structures) or exploiting auto-fluorescent properties of different cell types [25]. The FC is widely used in medical and diagnostic applications [26] and has also been proposed as a tool to determine the heterogeneity of microbial monocultures due

to physicochemical gradients appearing within culturing systems when grown in industrial-scale fermenters [27]. Recently, it has been used to characterize microbial community dynamics by analyzing the cytometric print (i.e., morphology-based clusters of virtual populations generated from the FSC/SSC data) [28]. However, only a few experimental data are available regarding the application of FC for bioenergy production. For instance, FC has been employed to monitor the culture dynamics in an AD system under substrate overload (i.e., increasing the organic loading rate) [29] to estimate the stability of the microbial community by analyzing segregated cell abundance in different clusters. The results showed that perturbations created abundance changes in three acidogenic subcommunities instead of variations in the microbiome phylogenetic composition. In addition, recent studies have also used the FC platform to evaluate morphological traits of *Clostridium beijerinckii* monocultures, in particular, monitoring different cell stages and the share of sporulating cells [30]. A practical application of PS monitoring of a thermophilic DF system has been reported in [31], using propidium iodide staining in FC measurements.

Another electrooptical technique to monitor the PS of microorganisms uses the relative changes in light absorption of cell suspensions under the application of an electric field at different frequencies, which has been termed frequency-dependent polarizability anisotropy (FDPA). The application of an electric field induces a rotational moment on charged particles that can be described by means of polarizability tensors. Following the electrophysical model, certain bacterial cells can be considered axially symmetrical ellipsoids of

revolution, which accumulate electric charges (i.e., membrane potential) at the interface between two media with different dielectric properties: the cytoplasm and suspension medium (segregated by the non-conductive layer of the lipid membrane). The mismatch of the electric properties at the cell-medium interface due to the Maxwell-Wagner polarization can be quantified using the changes in the absorption of orthogonally located light beams, $\alpha_{\parallel}(w)$ and $\alpha_{\perp}(w)$, and used to calculate the FDPA as $d\alpha(w) = \alpha_{\parallel}(w) - \alpha_{\perp}(w)$ [32]. Relative changes in this parameter can be considered an integral property reflecting the PS of microorganisms, as a result of the different levels of ion accumulation in the biotic phase. FDPA measurements have previously been used as an indicator of the PS of cell suspensions, although most of the cases pertain to monocultures. For instance, FDPA levels have been used to monitor the PS of aerobic *Escherichia coli* cultivations at different temperatures [33], to follow the acidogenic-solventogenic metabolic shift in *Clostridium acetobutylicum* [34] and to investigate the performance of *Lactobacillus plantarum* under different stress conditions, such as non-pH-regulated conditions, different substrate concentrations, nutrient limitations, and suboptimal cultivation temperature [35]. Recently, the applicability of FDPA to monitor the PS of mixed cultures has been tested; Habermann et al. [36] measured the FDPA of different AD systems using corn silage as feedstock and obtained decreasing FDPA values under acidification stress, suggesting the PS of the biotic phase in AD systems and sewage sludges can also be monitored through FDPA measurements.

In the present study, a simple method is used for the selection of an open mixed culture for the production of biohydrogen (hydrogen-producing bacteria, HPB) from a methanogenic AD sludge. Then, the HPB culture is used as inoculum of mesophilic DF tests, using as substrate a mixture of maize and grass silage, under pH-regulated and non-regulated conditions. During the course of fermentation, a batch-to-batch monitoring is performed following key parameters, such as pH evolution, biohydrogen yields, carboxylic acids, and ethanol titers in the liquid phase; while insights into the PS of the biotic phase are gained through FC and FDPA measurements.

Materials and Methods

Preparation of the HPB Inoculum

The biohydrogen-producing inoculum was obtained following a selection procedure that aims to suppress methanogenic activity from an AD sludge. Although several efficient strategies can be applied to obtain hydrogen-producing bacteria (HPB) from AD sludges [37, 38], the combination of acid/aerobic stresses was chosen to promote sporulation of Firmicutes, in particular,

Clostridium spp., which are anaerobic microorganisms, capable of forming spores. A sludge from an on-going AD plant digesting agricultural wastes was treated at pH = 3.0, using a 10 M HCl solution, and placed under aerobic conditions for 24 h at 35 °C in an incubation chamber (Multitron II-Infors HT, Bottmingen, Switzerland). The pH was then set to 6.0, and the resulting broth (spore-enriched due to the acid treatment) was then pre-cultivated under anaerobic conditions for 12 h at 35 °C, using a 30 g/L glucose media, prior to the inoculation of the DF systems to run the tests (the “[Experimental Set-Up of the DF System](#)” section).

Preparation and Pre-Treatment of the Substrate

The substrate for the DF tests was a mixture of agricultural products composed of maize and grass silage, mixed in a proportion of 9:1 (w/w). The proximate analyses of these feedstocks are reported in Table 1. Fresh materials were grinded with a laboratory batch mill (IKA-A10 Werke, Staufen, Germany) to decrease the particle size and subsequently diluted with tap water (4:11 w/w) to facilitate mixing during the fermentation process. The substrate media was pre-treated with a 30% (w/w) NaOH solution, until pH = 12 was reached and incubated at 35 °C for 24 h in the aforementioned incubation chamber.

Table 1 Raw substrate proximate analyses

	Corn silage		Grass silage	
	(g/kg _{FM})		(g/kg _{FM})	
Dry matter (DM)	371.0	± 17.3	491	± 35.1
Organic matter (OM)	356.9	± 23.6	443.4	± 37.0
Moisture	629.0	± 8.1	509	± 41.4
Total ash	14.1	± 0.3	47.6	± 1.8
Crude protein (CP)	23.4	± 0.5	82	± 3.0
Crude fiber	76.4	± 5.6	118.3	± 4.9
Carbohydrates	123.9	± 10.9	–	± –
Crude fats	10.4	± 0.8	15.7	± 0.2
ELOS	252.3	± 6.1	317.2	± 7.9
ADF	89.4	± 4.6	143.9	± 5.0
NDF	154.0	± 2.8	232.7	± 17.1
NFC	169.2	± 12.0	112.9	± 8.7
Avail. CP	46.7	± 3.5	66.3	± 2.5
Ammonia	0.6	± 0.1	1.8	± 0.1
NH ₃ -N of total N (%)	12.6	± 0.6	11.5	± 0.7
BGP (m ³ biogas/t _{FM})	234.8	± 7.9	286.1	± 7.8
BMP (m ³ CH ₄ /t _{FM})	123.3	± 8.3	150.2	± 11.7

FM, fresh matter; DM, dry matter; OM, organic matter; CP, crude protein; ELOS, enzymatic soluble organic matter; ADF, acid-detergent fiber; NDF, neutral-detergent fiber; NFC, non-fibrous carbohydrates; BGP, biogas potential; BMP, biomethane potential

Experimental Set-Up of the DF System

The DF experimental tests were conducted in 1 L STR EloFerm bioreactors (EloSystems GbR, Berlin, Germany) with a working volume of 400 mL. The empty vessels were sterilized (121 °C for 20 min), filled with the pre-treated substrate, and inoculated with the pre-cultivated HPB inoculum in a ratio of 10% (v/v). Bioreactors were operated at mesophilic conditions (35 °C) and under anaerobic conditions achieved by flushing N₂ for 10 min into the fermentation broth at the beginning of the tests. The bioreactors were stirred magnetically, at a rate of 200 rpm. Two bioreactors were operated in parallel: in the first fermenter (A), a pH = 6.0 was fixed as set point and controlled with an integrated control loop of the EloFerm system using a 30% (w/w) NaOH solution, while in the second bioreactor (B), the pH remained uncontrolled throughout the course of fermentation.

Analytical Measurements

Gas Analysis

The produced gas of each bioreactor was first dried in a silica-gel bed. The composition was measured online by means of a gas analyzer arrangement. It consisted of three in-series connected gas sensors (BlueSens, Herten, Germany): BPC-CO₂ and BPC-CH₄, based on infrared (IR) absorption for CO₂ and CH₄, respectively, and BPC-H₂, based on thermal conductivity difference to estimate the amount of H₂. The total gas flow was measured using the water-displacement method: connecting the output of the gas analyzer arrangement to acidic water (pH = 2.0), which displaces water-filled gas-meter flasks, ending up to collector bottles placed on analytical balances. The biohydrogen yield of each DF system is reported in normal milliliters per gram of volatile solid (NmL/gVS), which are the standard units used in the field of biogas production that allow comparing systems at different conditions (e.g., pressure/temperature) for complex feedstocks (i.e., based on the organic matter or volatile solid content, see Table 1 for the proximate analysis).

Quantification of Short-Chain Carboxylic Acids and Solvents

Raw samples were first filtered with 125 µm nylon filters (Carl Roth GmbH, Karlsruhe, Germany) and then using syringe filter units (i.e., pore size 0.45 µm) and subsequently diluted (1:1 v/v) with ultra-deionized water provided by an Easypure II RF device (Barnstead, Iowa, USA). High-performance liquid chromatography (HPLC) was used to analyze the liquid phase, in particular short-chain carboxylic acids and solvents. The HPLC system (1200-series Agilent Technologies, Waldbronn, Germany), equipped with a refractive index detector (RID) and a 300 × 7.7 × 8 mm HyperRez™ XP Carbohydrate H⁺ column (Fisher Scientific,

Schwerte, Germany), was operated at 65 °C using 5 mM H₂SO₄ solution as eluent, at a flow rate of 0.6 mL/min for a cycle time of 50 min [39]. A four-point calibration was performed for the detection of formate, acetate, propionate, butyrate, ethanol, propanol, and butanol in the operative range of the DF.

Flow Cytometry Analysis

For the FC analysis, samples were collected at the beginning of the fermentation in order to set the optimal detector sensitivities (by voltage adjustment) for each channel and with unstained cells (negative control). Samples collected from each bioreactor (A and B) were vacuum filtered using 0.2 µm paper filters, previously wetted with phosphate-buffered saline solution (PBS composition: 136.90 mM NaCl, 2.70 mM KCl, 8.10 mM Na₂HPO₄·2H₂O, and 1.80 mM KH₂PO₄, pH = 7.2). Five rinsing cycles of 1 mL PBS solution were applied to the solids retained in the filter paper to remove debris and impurities. The filter papers, containing the retained biotic phase, were immersed in 10 mL PBS after the washing steps (5 × 1 mL PBS), briefly vortexed and diluted until a cell concentration of 1·10⁶ cells/mL prior to staining. The fluorochrome BOX, also called DiBAC₄(3) or bis-(1,3-dibutylbarbituric acid) trimethine oxonol, was employed to stain the cells. Aliquots of 200 µL were stained with BOX, at a final concentration of 0.5 µg/mL together with dissolved EDTA (~29.75 µg/mL), at room temperature for 4 min to assess the relative amount of depolarized cells. Positive controls to assess the suitability of BOX-staining for the mixed consortia (HPB) were performed by applying a thermal treatment to the biological material of 80 °C for 1 h using the BOX/EDTA mixture and the same staining conditions (4 min). Finally, the samples were analyzed by means of a MACSQuant Analyzer (Miltenyi Biotech, Bergisch-Gladbach, Germany), equipped with three lasers (violet 405 nm, blue 488 nm, and red 635 nm), several filters and mirrors, two photomultiplier tubes for the detection of scattered light (FSC, SSC), and eight photomultiplier tubes for the detection of fluorescent signals. When bacteria samples present suitable characteristics to be hydrodynamically focused by FC, the FSC signal aims to discriminate cells based on the scattered light in the same axis as the incident light beam (i.e., often used an indicator of the *cell length*, although the more general interpretation considers FSC as a measure of cell size or volume), while SSC collects scattered light at an angle of 90° to the incident beam, which provides information about cell complexity, granularity, and/or cell surface.

Frequency-Dependent Polarizability Anisotropy Measurements

Samples were first filtered with 125 µm nylon filters (Carl Roth GmbH, Karlsruhe, Germany) in order to remove large

debris and inert particles. Subsequently, filtrate aliquots (10 mL) were centrifuged at 4 °C and 600 rpm for 5 min. The supernatant was diluted with ultra-deionized water (approx. 1:10 v/v) to adjust the initial electrical conductivity of the samples to an adequate range compatible with FDPA measurements ($< 40 \mu\text{S}/\text{cm}$). FDPA measurements were conducted at line with samples extracted from bioreactor A and B at the following different time points: 1, 20, and 33 h of fermentation using an EloTrace device (EloSystems GbR, Berlin, Germany). The FDPA levels of each sample were acquired at different frequencies of the low and medium frequency spectrum ranges, namely 210, 400, 900, and 2100 kHz, based on the type of analyzed fermentation broth and previous exploratory tests. The mean values are presented together with the standard deviation of the triplicates measured for each sample. The FDPA values are displayed using arbitrary units of the equipment, scaled by a factor of $5 \cdot 10^{-31} \text{ F m}^2$ (i.e., farad square meters) [40], for easier representation and comparison with literature data.

Statistical Analysis

The statistical significance was evaluated using SigmaPlot 10.0; the Student's *t* test was performed as pairwise comparisons (systems A and B) at different timepoints for the FDPA and FC tests. Additional details are provided in the [Supplementary Material](#).

Results and Discussion

DF Cultivation Performance

The DF systems were set following the procedure described in the “[Experimental Set-Up of the DF System](#)” section; the bioreactors were operated with (A) and without (B) pH control. The acid-treatment applied to the AD sludge to obtain the HPB inoculum proved to be effective, since methanogenic activity was not detected during the tests. Throughout the DF experiments, no methanogen growth was appreciated in microscopic observations nor was methane production detected at the off-gas analyzer arrangement (see the “[Gas Analysis](#)” section). As a matter of fact, the continuous analysis of gaseous matrices using multiple gas sensors has been recently tested as a useful strategy for the course analysis of DF systems. Particularly, methane sensors can be useful to identify if populations (e.g., Archaea) and metabolic shifts occur in the DF systems [41].

Figure 2a shows the pH courses for both systems while the cumulative H_2 yield is reported in Fig. 2b. During the tests, the pH value shows the following three different phases:

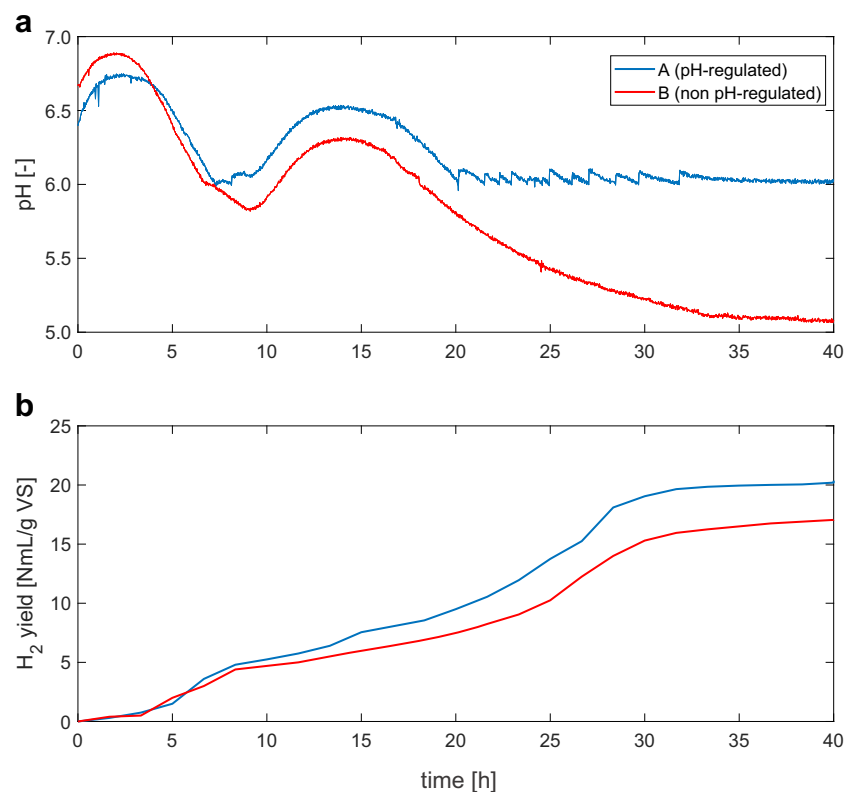
- i) an oscillating behavior around the initial pH value of the systems during the first 20 h of the fermentation;
- ii) a regulated pH signal at a pH = 6.0 for the system (A) and a strong decrease for the non-regulated system (B) in the interval 20–33 h; and
- iii) a constant pH signal at 6.0 and 5.1, for (A) and (B), respectively, until the arrest of the fermentation processes (33–40 h).

In the first 5 h, the microorganisms seem to adapt to the new substrate and environmental conditions. These initial oscillations can be linked to the hydrolytic step during the first hours [42] and can be explained by considering two concomitant occurring phenomena: a slight pH decrease due to the production of moderate titers of volatile fatty acids (VFA) and a pH increase because of the dissolution and reaction of the CO_2 produced by decarboxylation reactions and consequently a basification due to the formation of bicarbonate (coupled to a low production of H_2 as shown in Fig. 2b). During this phase, a similar gas evolution in both systems was recorded. Hence, microbial activity was predominantly hydrolytic, i.e., oriented towards the production of enzymes and other biologically active molecules, necessary to attack the slowly degradable fraction [40]. The latter amounted to $\sim 21\%$ of the organic matter for corn silage and to $\sim 27\%$ for grass silage (Table 1), and this phase was accompanied by a slow H_2 production rate and low carboxylic acid production. During the following hours (~ 15 h), the pH oscillatory behavior was still present. Nevertheless, the biohydrogen production rate increased for both systems. Then, an exponential growth phase of about 12–13 h took place, which yielded different carboxylic acids titers in the liquid phase and gas production ($\text{H}_2 + \text{CO}_2$) with a decreasing pH trend in (B) and constant pH value for (A), at pH = 6.0 (due to the pH control). Finally, the cells entered the stationary phase (after 33 h), and the evolution of gas ceased, while the pH values remained constant.

Cumulative hydrogen yields amounted to 20.80 ± 0.05 and $17.08 \pm 0.05 \text{ NmL}/\text{gVS}$ for the system (A) and (B), respectively (see Fig. 2b). The gas production followed a similar trend for both systems. However, the mean H_2 production rate was higher for the pH-regulated system, while the concentrations of the gas phase were about 28.3 ± 0.5 and $26.0 \pm 0.8\% \text{ v/v}$, with a correspondent CO_2 fraction of 71.7 and 74.0% v/v, for (A) and (B), respectively. As noted by [43], a pH control in the DF systems promotes higher H_2 production using either suspended or immobilized DF microbial systems; in the present case, about 18% more H_2 was obtained for the pH-regulated system.

For simple sugar feedstocks and monoculture systems, the maximum biohydrogen yield can be calculated using stoichiometry; while for complex feedstocks and open mixed cultures, the yield strongly depends on the selected inoculum, the

Fig. 2 Course of pH value (a) and hydrogen yields (b) during the DF tests



culturing conditions, and the feedstock pre-treatment. The obtained yields for system A and B are thus compared with some literature references for similar matrices, highlighting the source of the acidogenic inoculum (HPB), the pre-treatment of the feedstocks, the pH and culturing temperature, and the biohydrogen yields in each case (see Table 2). In these studies, the acidogenic inocula are selected from methanogenic sludges using different selective pressures (e.g., heat shock, substrate overload), the process conditions are mesophilic or thermophilic, and the feedstocks (grass and maize silages) are pre-treated using different strategies (mechanical, acid, basic, and enzymatic treatments). Besides, the biotic phase is not studied in detail, while the optimization of the process in certain of these studies is performed based on macroscopic process conditions, such as optimal pH and temperature, to target enhanced biohydrogen yields.

The pH value is a key parameter not only governing the microbial activity towards H₂ productivity, but also influencing the production of other metabolites by activating different biological pathways [37, 43]. For instance, the analysis of supernatant samples (Fig. 3) revealed some differences between the system with (A) and without pH control (B). Despite the different pH conditions, the butyric acid fermentation was predominant for both systems with three-fold higher titers compared with the rest of detected carboxylic acids and ethanol. The pH-controlled system produced a higher amount of acetic acid and ethanol towards the end of

the fermentation (20–33 h), while the yield of these compounds in the non pH-regulated system was rather modest. Moreover, for system (A), formic acid levels remained almost constant (± 2 mM), while for system (B), a slight formic acid consumption was observed in the exponential H₂ production phase (20 h). Lastly, propionic acid remained almost unaltered for both systems; hence, nor production or consumption was observed in the systems. In fact, the measured titers of propionic acid remained < 2 mM, which is well below inhibition thresholds reported in the literature for similar DF systems [50].

In acid fermentations, particularly those where Firmicutes and specially *Clostridium* spp. take part, low pH levels are believed to induce the shift from acidogenesis to solventogenesis. The final pH for the regulated system (A) was 6.0, while for the non-regulated systems was ~ 5.1 , which are higher values compared with the solventogenic range of *Clostridium* spp. ($\text{pH}_{\text{solventogenesis}} < 4.5$ [51]). This shift has been studied in pure and mixed cultures, and among other reasons, it represents a strategy to overcome pH stress by partial reutilization of a share of the produced acids [52].

Flow Cytometry

The FC allows the monitoring of the morphology of the biotic phase during the process. First, the morphology of the mixed consortia was analyzed using the bivariate

Table 2 Comparison among the present results and DF studies of maize/grass silage from literature

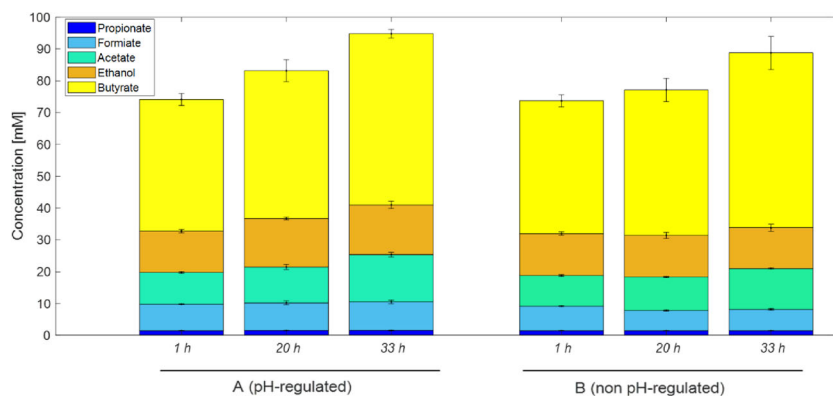
Reference	Feedstock	Feedstock pre-treatment	Inoculum for DF	Yield (Nml/ gVS)	Initial pH (-)	Temperature (°C)	Mode
[44]	Grass silage	None	Thermally treated AD sludge (100 °C for 0.5 h)	15.50–19.80	7.0	37	Batch
[44]	Grass silage	Thermal + acid pre-treatment	Thermally treated AD sludge (100 °C for 0.5 h)	64.20–72.70	7.0	37	Batch
[45]	Grass silage	Mechanical grinding	Thermally treated AD-sludge (boiling for 0.5 h)	0.6–17.7	4.0, 5.0, and 6.0	35, 55, and 70	Batch
[46]	Grass silage	Mechanical cutting	Thermally treated AD-sludge (boiling for 0.5 h)	5.01–6.27	6.0	55	Batch
[46]	Grass silage	Mechanical cutting + basic pre-treatment	Thermally treated AD-sludge (boiling for 0.5 h)	5.83–7.09	6.0	55	Batch
[40]	Corn silage	Mechanical grinding	Thermally treated AD-sludge (80 °C for 1.5 h)	22.22–27.89	5.5	35	Semi-continuous
[47]	Maize silage	Dried at 60 °C and milled	Load shock-treated AD sludge (OLR increased up to 10 g VS/L/d)	19.00–52.00	5.0–5.5	38	Semi-continuous
[48]	Maize silage	n.a.	Granular sludge from a USBA plant	48.90–62.30	6.5	37	Batch
[48]	Maize silage	n.a.	Granular sludge from a USBA plant bioaugmented with <i>Pirromyces rhizinflata</i>	55.30–63.50	6.5	37	Batch
[49]	Comstalk wastes	Mechanical grinding + acid pre-treatment + enzymatic hydrolysis	Thermally treated natural river sludge (100 °C for 0.25 h)	71.39–140.64	5.0, 6.0, 7.0, and 8.0	36	Batch
This study	Maize/grass silage (9:1 w/w)	Mechanical grinding + basic pre-treatment	Acid-treated AD sludge + aerobic stress (pH = 3.0 for 24 h in aerobic conditions)	17.03–20.85	6.0	35	Batch

n.a., not available

FSC/SSC density plots complemented with the correspondent histograms. The information provided by the FSC/SSC channels of FC served to differentiate among morphological traits of different subpopulations (i.e., relative cell length and granularity, see the “Flow Cytometry Analysis” section). As presented in Fig. 4a, the acid treatment of the AD sludge to select HPB resulted in a spore-enriched broth, which presents a cytometric print consisting mainly in one differentiable population, with unimodal histograms for the FSC and SSC channel (Fig. 4b). The pre-cultivation phase (pre-culture = growth of the inoculum for the DF tests) allowed a fraction of these initial free-spore to germinate, into the characteristic rod-shaped morphology (i.e. which is typical for many acidogenic bacteria, such as *Clostridium* spp.) and endospore/forespore form (Fig. 4a and c). This was reflected in a bimodal distribution in the SSC channel and three populations in the FSC channel (Fig. 4b). The FSC/SSC print can be associated, crossing the information with optical microscopy observations (Fig. 4c), to (a) the non-germinated fraction of free spores, (b) rod-shaped vegetative (germinated) cells, and (c) endospores. This assessed morphology distribution after pre-cultivation coincides with the first measurement point of the DF ($t = 1$ h) and is common for both systems. A similar cytometric print was found for the second measuring point ($t = 20$ h), although modest changes can be appreciated. A virtual clustering technique was applied to keep track of the changes in the cytometric prints, hypothesizing that they correspond to the observed morphologies as shown in Table 3. The trend of these sub-populations (virtual clusters) indicated that the relative fraction of free spores did not vary significantly, increasing by 4.8% for system (A) and about 6.1% for (B). Concomitant reductions in the share of rod-shape cells were observed, amounting to 4.8% for the system (A) and 7.74% for (B). Interestingly, the population of endospores remained almost unaltered for system (A), while an increase of 1.66% was reported for (B) (see Table 3). At the end of the DF tests ($t = 33$ h), the cytometric print showed a significant shift from vegetative cells towards free-spores (i.e., sporulation) in both systems, which resulted in final amounts of free spores of more than half of all detected cells, which were used as a calculation base for FC analysis (Fig. 4d): 55.50 and 54.05% for (A) and for (B), respectively. This shift in the microbial population is consistent with the arrest of the biohydrogen production and probably reflects the response of the mixed consortia to starvation conditions.

Another set of tests was performed using the FC platform by staining samples with DiBAC₄(3) in order to track relative changes in the number of (partially) depolarized cells within the mixed HPB consortia. The

Fig. 3 Carboxylic acids and ethanol titers measured during the DF tests



comparison between the negative control (not heat-treated and unstained samples) and the positive control suggests that the SSC/BOX-A cytometric fingerprint could be used as an indicator of the PS of the mixed HPB fermentative microorganisms, since a clear shift in the relative fluorescence intensity in the channel for BOX detection (i.e., excitation at 488 nm and emission at 525 nm) can be observed (Fig. 4e). Different staining protocols can be adopted for the evaluation of the PS through FC analysis of bacteria: either monocultures or mixed populations. A commonly measured parameter for the PS is the membrane potential. Cationic dyes accumulate in polarized cells, while anionic dyes (e.g., DiBaAC₄(3)) accumulate in depolarized cells [53]. Additionally, the staining step included the addition of EDTA (see the “Flow Cytometry Analysis” section), which was aimed to increase the range of action of the dye and improve the reliability of the analysis for complex microbial communities (e.g., mixed bacteria, either Gram-positive or -negative) [54]. The calculations regarding the percentage of depolarized cells were performed using the triggering threshold shown in Fig. 4e, followed by the differentiation and comparison of the signal intensities of positive and negative (unstained cells) controls and the samples for each timepoint. Only a low amount of debris after the washing cycles with PBS (<2%) was obtained and the instrument background was relatively low (data not shown). For the first

timepoint measurement ($t=1$ h), systems (A) and (B) show no significant difference in the number of depolarized cells, which amounted to 23.06 ± 4.60 and $23.68 \pm 3.36\%$, respectively. For the second point ($t=20$ h), the system without pH control exhibits a share of depolarized cells, which is twice ($55.57 \pm 12.41\%$) the value of the one measured for the pH-regulated system ($26.99 \pm 8.39\%$). Finally, at the end of the cultivation ($t=33$ h), the percentage of depolarized cells increased, reaching up to $30.31 \pm 1.34\%$ for the pH-regulated system, while the share for system (B) ($48.90 \pm 5.11\%$) slightly decreased compared with $t=20$ h. These differences among the pH-regulated and non-regulated DF systems, in terms of relative levels of depolarized cells, as well as the different detected morphologies of the sporulation dynamics, seem similar with literature references [55].

Frequency-Dependent Polarizability Anisotropy Measurements

The results of the FDPA measurements, for both systems, are presented in Fig. 5. First of all, it must be noted that FDPA measurements are very sensitive to the presence of ions. Hence, an optimization of the sample preparation for accurate measurements was performed with varying dilution factors, biomass concentration, and centrifugation steps. Several tests were carried out (data not shown) to optimize the analytical

Table 3 Percentual time-resolved evolution of hypothesized FSC/SSC virtual clusters (sub-populations) from FC data

Virtual cluster	pH-regulated system (A)			Non pH-regulated system (B)			Associated morphology
	$t=1$ h	$t=20$ h*	$t=33$ h*	$t=1$ h	$t=20$ h*	$t=33$ h*	
Group A	31.72 ± 0.02	36.53 ± 0.15	55.51 ± 0.34	28.07 ± 0.06	34.14 ± 0.11	54.05 ± 0.26	Free spore
Group B	50.78 ± 0.14	45.98 ± 0.05	32.11 ± 0.21	52.53 ± 0.98	44.79 ± 0.50	30.66 ± 0.19	Rod-shaped
Group C	17.50 ± 0.07	17.49 ± 0.07	12.38 ± 0.18	19.40 ± 0.11	21.07 ± 0.16	15.29 ± 0.31	Endospore

*The different time-evolution of the systems A and B for timepoints II and III is statistically significant at $p < 0.05$. For additional information, see the [Supplementary Material](#)

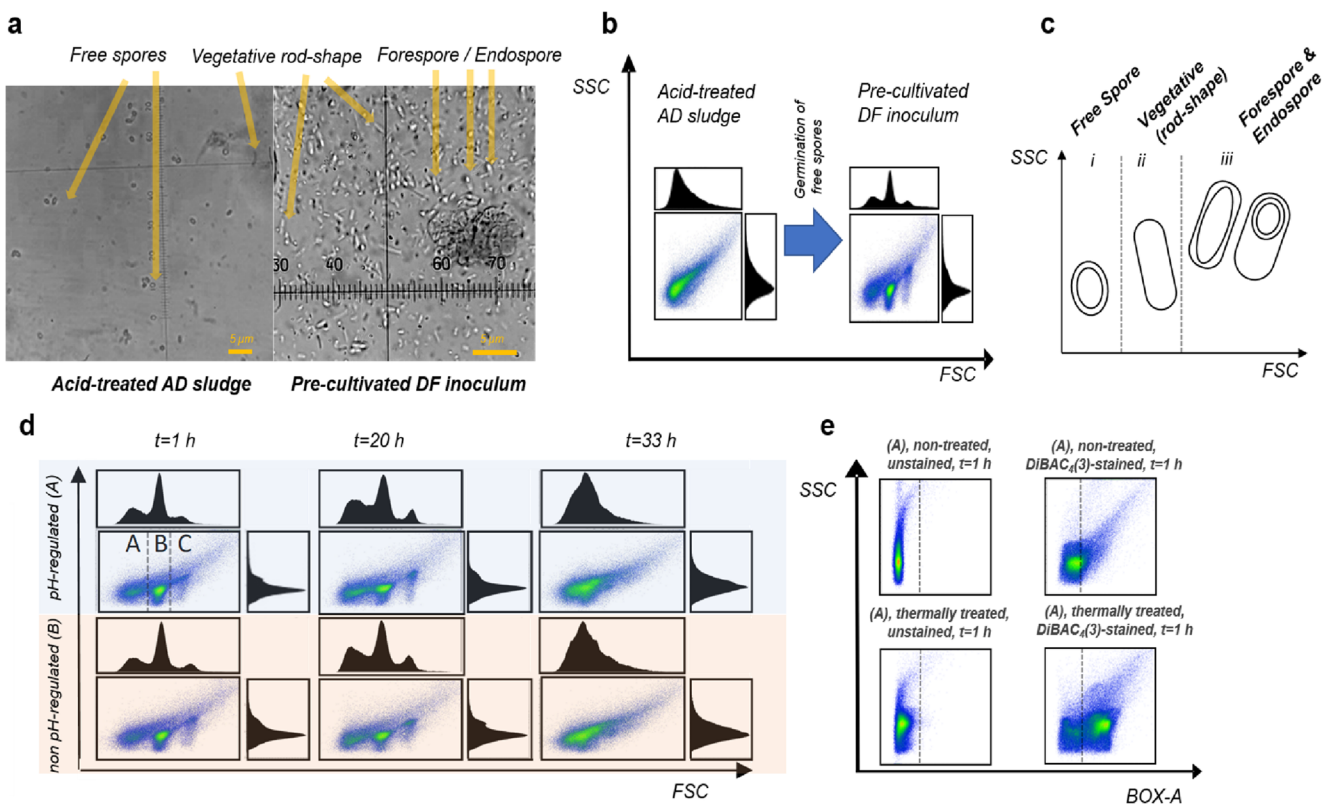


Fig. 4 Analysis of the DF culture: **(a)** microscopy of the acid-treated AD sludge and the pre-cultivated DF inoculum; **(b)** cytometric print (FSC/SSC) during the pre-culture phase; **(c)** coupling virtual clusters (FC) and observed morphologies; **(d)** time evolution of the cytometric print (FSC/

SSC) of the DF systems under pH-regulated and non-regulated conditions; **(e)** gating and triggering strategy (BOX-A/SSC) for the quantification of depolarized cells

protocol, and the final protocol is reported in the “[Frequency-Dependent Polarizability Anisotropy Measurements](#)” section. In this respect, typical AD or DF broths exhibit high conductivity values (i.e., in the range 10–40 mS/cm). The conductivity is a parameter that can be periodically measured to monitor process performance in AD [56]. However, the electrical conductivity of the fermentation broth, especially at high frequencies, could have an important effect on the FDPA measurements. Since FDPA measurements pertain to the biotic phase, the selection of ultra-deionized water for the suspension of microbial cells during the measurements resulted to be the best strategy.

Figure 5 shows that at the beginning of fermentation tests ($t = 1$ h), the FDPA levels were high and similar for both systems under analysis, particularly at low frequencies. This fact probably suggests that the biotic phase presents a high activity due to the necessity to adapt to the new environment and that higher FDPA levels can be associated to the hydrolytic phase. Both systems (A and B), however, exhibited a decreasing trend of FDPA values along the course of fermentation due to substrate depletion and a concomitant decrease in hydrogen production. A possible explanation of these trends is that the systems reached a stable conformation in acidogenic conditions, where the PS (i.e., mean membrane cell

polarizability in this case) is controlled by substrate availability, after the lag phase and hydrolysis. However, in the second measuring point ($t = 20$ h), bioreactor (A) showed a higher FDPA level than system (B), most probably due to the controlled pH condition, which guarantees a constant environment for the mixed culture to grow and allow cells to maintain a constant membrane potential and well-functioning membrane transporter systems. These lower values of the FDPA level for the non-regulated pH system (B) are in accordance with the lower cumulative biohydrogen yields in Fig. 2b. For the last point ($t = 33$ h), where the stationary phase was achieved, FDPA values are also similar for both systems (A and B), although for system (B), a slight increase compared with the 20 h sample was observed.

In general terms, bacteria cells are believed to exert control mechanisms to maintain constant values of the proton-motive force (pmf), balancing both components membrane potential ($\Delta\Psi$) and homeostasis (ΔpH) [57], or to change the operative pmf values due to metabolic shifts or to reduce pmf thresholds when exposed to different stress conditions. For instance, pmf values for *Clostridium acetobutylicum* have been estimated in the range of -99 and -84 mV when cultivated at pH conditions of 5.0 and 6.0, given by the ΔpH components of 59 and 28 mV and the $\Delta\Psi$ component of -40 and -56 mV,

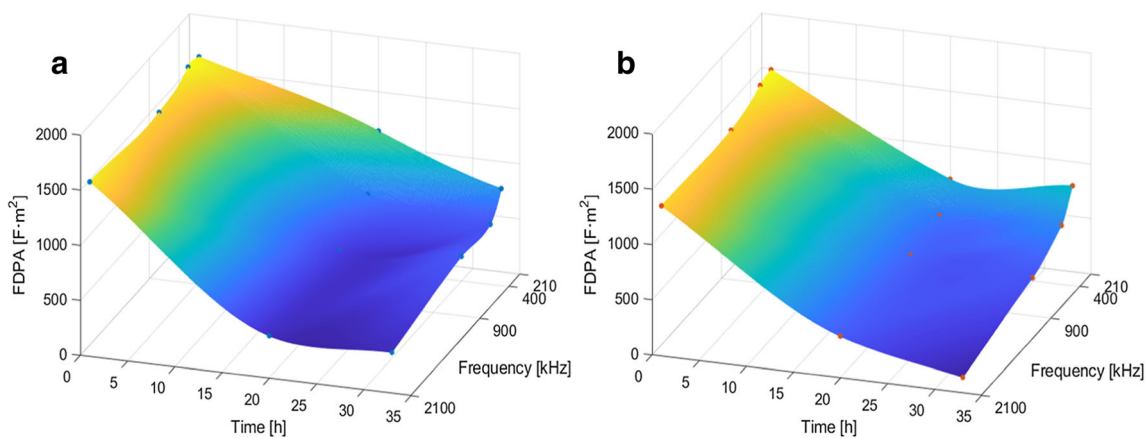


Fig. 5 Electrooptical measurements of FDPA at different frequencies, for the pH-regulated (a) and non-regulated (b) systems

respectively [58]. In the present case, the biotic phase was constituted by a mixture of acidogenic bacteria (based on the butyrate titers, presumably *Clostridium* spp. played the predominant role in the mixed consortium), and the pH stress was used as probe parameter. Although metabolic products (Fig. 3) did not significantly differ among the studied systems, the variations in the FDPA can be taken as a lumped indicator for the PS of the biotic phase based on the changes of the internal Maxwell-Wagner polarizability [59]. Additionally, it should be noted that the equilibrium of the organic acids of interest is shifted towards the non-dissociated form at low pH (i.e., under acidic conditions), which could additionally have stressed the non-regulated pH system (B). In this case, cells might spend a higher fraction of energy to counteract pH stress, in order to maintain homeostasis, rather than for growth, which can also partly explain the lower hydrogen yields in the gas output and reduced titers found in the liquid broth for system (B).

Physiological State Correlation Between FDPA and FC

Although there is no standard method for measuring the PS of mixed bacteria populations, the tested techniques, FC and FDPA measurements, proved to be helpful to track the PS dynamics of the open mixed culture used in the DF process. Both techniques are based on different physical measurement principles. On one hand, FDPA is a volumetric method that is based on the ionic properties of the cell suspension and its variations do reflect valuable information regarding the physiological state of the biotic phase. In the present case, different FDPA levels were useful to differentiate the phases of the culture (i.e., hydrolysis, acidogenesis, and starvation), as well as to compare the pH-regulated with the non-regulated system. For monocultures, the cell size can also be determined using the relaxation times between the relaxed and excited states during FDPA measurements, which is an approach that might be extended for mixed cultures. On the other hand, the FC platform offers different levels of information regarding the microbial community structure. The FSC/SSC fingerprint

can be useful to identify morphological changes and the relative abundance of key groups or sub-populations within microbial communities, while the selection of different fluorochromes can target, at monoculture and mixed culture level, different parameters to characterize the physiological state. The present results show an inverse correlation for both systems (A, B) between the obtained in the FDPA (Fig. 6a) and the percentage of depolarized cells calculated using FC (Fig. 6b). The FDPA values at 210 kHz for system (A) showed an

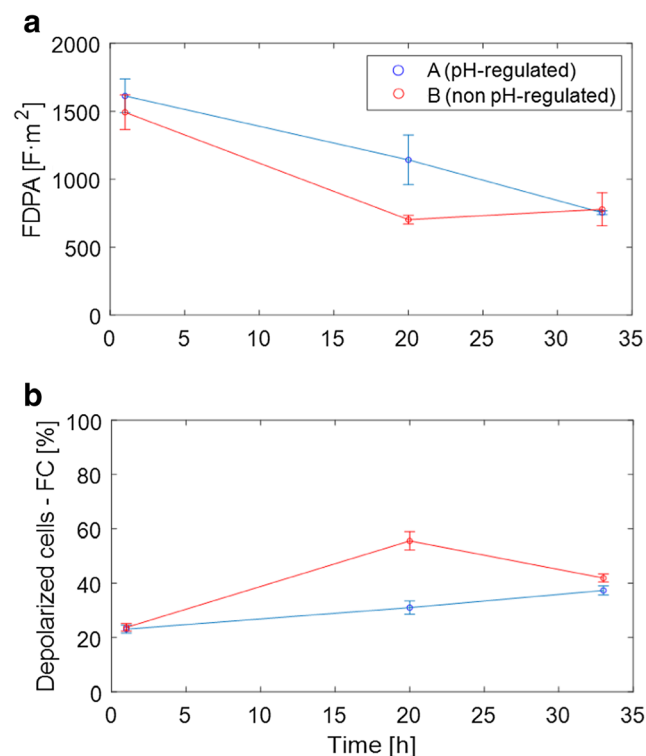


Fig. 6 Physiological state* estimation over the course of DF through FDPA measurements (a) and DiBAC₄(3)-stained FC samples (b). *The different time-evolution of the systems A and B for $t = 20$ h is statistically significant at $p < 0.05$. The FC measurements are also statistically significant at $p < 0.05$ for $t = 33$ h. For additional information, see the [Supplementary Material](#)

almost linear decrease over time, while values of system (B) had a minimum at 20 h. Importantly, FDPA measurements provide a mean value pertaining the whole cell population. Generally, microorganisms can either consume energy for dividing themselves (i.e., increase biomass) or to improve their intracellular status (i.e., increase polarizability, maintenance). At the beginning of the fermentation, the biotic phase is adapting to the actual environmental conditions and is therefore consuming energy basically for maintenance (i.e., high polarizability in both cases). As the cultivation evolves, there is a certain biomass growth over time and thus a steadily decline of the mean cell polarizability. Without pH control, microorganisms face the acidification stress, and therefore, they consume more energy for maintenance (i.e., higher polarizability, lower biomass, lower H₂ production) when compared with system (A) at controlled pH at 6.0. These trends in FDPA measurements are consistent with the trends obtained by the FC measurements. For system (A), the percentage of depolarized cells increases continuously over time (although modestly), while for system (B), there is a maximum in the number of depolarized cells at time point 20 h. Moreover, the cumulative-specific hydrogen evolution for each system (see Fig. 2) is also in agreement with the resulting trends of FDPA and FC measurements. At the timepoint 1 h, the yield was similar for both systems, while the measurements at 20 h showed that system (B) only achieved a 77% of the yield of system (A) (i.e., the most critical condition as shown by FDPA and FC), and for the stationary phase, system (B) achieved 82% of the yield obtained in (A). Finally, it should be taken into consideration that the stress parameter for these tests was the influence of pH regulation, which might have impacted or stressed the biotic phase similarly, whether assessed by the FDPA or the FC technique. However, it is worth noting that for other stress conditions, the information independently provided by each technique can rather be supplementary.

Conclusions

The DF of complex feedstocks, such as agricultural products, is of great interest because it offers a great potential for process integration, with recovery of bioenergy in gaseous form (biohydrogen), and also as a liquid metabolite pool of chemicals of high valorization potential. The use of PATs can improve process performance by facilitating process integration and changing the paradigm regarding the physiology of open mixed cultures in DF and other *black-box* biotechnological processes. This approach can help to avoid process failures, such as substrate and/or product inhibitions, undesired population shifts, low metabolic activity, and suboptimal product yields. Here, the pH influence on the PS was useful to monitor the behavior of the DF systems in terms of

biohydrogen and carboxylic acid productions as well as to assess physiological information in terms of morphology and key parameters related to the membrane polarizability by means of flow cytometry and electrooptical polarizability measurements. Although some industrial processes use sporulation-suppressed microbial strains, open mixed cultures for the production of biohydrogen and biochemicals might include sporulating strains, which can affect the process performance. Hence, morphological observations can be useful to determine groups and physiological states and to improve cellular performance. Additionally, the use of FC with DiBAC₄(3) staining and the measurements of the FDPA levels seem to be useful indicators for complex microbial cultures, and they can potentially facilitate monitoring of their PS during complex fermentations. A natural progression of this work is to analyze and assess the potential of these techniques to be used in continuous and large-scale processes and the development of control strategies for different biorefinery processes.

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