Expression of different types of [FeFe]-hydrogenase genes in bacteria isolated from a population of a bio-hydrogen pilot-scale plant

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A B S T R A C T

[FeFe]-hydrogenases are the enzymes responsible for high yield H2 production during dark fermentation in bio-hydrogen production plants. The culturable bacterial population present in a pilot-scale plant efficiently producing H2 from waste materials was isolated, classified and identified by means of 16S rDNA gene analysis. The culturable part of the mixed population consists of nine bacterial species that include non-hydrogen producers (Lactobacillus, Enterococcus and Staphylococcus) and several Clostridium that are directly responsible for H2 production.

An extensive analysis of the expression of [FeFe]-hydrogenases in the three best producer strains was achieved by RT-PCR, covering the complete set of known genes for each species.

This revealed that during H2 production there are several different [FeFe]-hydrogenases simultaneously expressed, with genes belonging to the same phylogenetic and structural classification sharing similar transcriptional profiles.

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Introduction

The characterisation of bacterial consortia involved in H2 production in pilot-scale and operative plants with dark fermentation is of applicative and theoretical importance. In fact, dark fermentation is recognised as a feasible method for sustainable hydrogen production combined to treatment and valorisation of waste biomasses [1,2].

A crucial point for a biotechnological approach is that of devising methods to optimise and stabilise the microbial consortia that ensure high productivity [2,3], as well as elucidating the biochemical pathways that can be tuned or controlled for the hydrogen production purpose. A particular interest concerning the precise mode and timing of hydrogen production is relevant for bio-hytane production, a very promising fuel that can be obtained in two-phase dark fermentation plants by suitably combining bio-hydrogen and bio-methane streams [4].
The characterisation of the bacterial populations, including both hydrogen producers and non producers, provides information that can support the improvement of hydrogen production. In fact, hydrogen producers can have very different metabolic strategies resulting in variable yields of hydrogen. The study of the consortia still lacks some crucial points, such as a complete listing and characterisation of the diversity of genes – and proteins – that in hydrogen producers are the first responsible for high hydrogen productivity, namely hydrogenases. In this respect, there are recent works on the non-hydrogen producers [2,5–8], while concerning hydrogenases diversity in strains found in dark fermentation plants the available papers focus only on few genes and proteins. In particular the diversity of [FeFe]-hydrogenases has not yet been fully investigated in the perspective of achieving optimised performances in dark fermentation plants. Thus, the study of new [FeFe]-hydrogenases, their time and mode of expression, leads to a better knowledge of the mechanisms of catalysis for improved biotechnological applications [9–15].

[FeFe]-hydrogenases are the best performing hydrogen producing biocatalysts [16–21], with turnover frequency of up to 10^3 s^{-1}. These enzymes have a modular structure: in addition to the catalytically active domain (named H-domain), several other accessory domains may be present [17,22]. Moreover, the functional enzymes can be monomeric, dimeric, trimeric or tetrameric, having the various domains on the same polypeptide chain or separated over one or more subunits. The detailed analysis of the sequences of [FeFe]-hydrogenases allows the definition of different groups and subgroups on the basis of the phylogenesis and domains composition [23].

The main feature common to all known [FeFe]-hydrogenases is the highly conserved part of the H-domain surrounding the active site. This region is characterised by three signature sequences containing the four cysteines that coordinate the active site and are named motifs L1 (TSCCPxW), L2 (MPCxxKxxE) and L3 (ExMACxxGCxxG) [23].

The detailed analysis of the [FeFe]-hydrogenases observed into the genus Clostridium revealed a very large number of enzymes with an exceptional diversity [23,24]. Even within the same strain up to 8 genes encoding for different [FeFe]-hydrogenases can be detected, all belonging to different groups, thus suggesting different roles, interactions with several physiological redox partners and a complex regulation of the biological function of each enzyme.

So far, the high diversity of [FeFe]-hydrogenases has been studied both in natural environments [25–27] and several pilot-scale plants for H₂ production [28–30].

Here the [FeFe]-hydrogenases diversity is investigated in the bacteria isolated from a pilot-scale plant for H₂ production fed with waste materials that was previously described [31]. The high efficiency observed suggested the involvement of highly efficient H₂ producing microorganisms and highly efficient hydrogenases.

With the aim of characterising the vital form of the bacterial species present in the plant, the cultivable part of the population present at the time of highest percentage in hydrogen produced was isolated, giving specific, although partial, insights into the composition of the consortium responsible for the wastes degradation and H₂ production. Extensive RT-PCR analysis of the [FeFe]-hydrogenase genes expressed by the isolated hydrogen-producing strains is presented.

The novelty of this work is that RT-PCR was performed covering the complete set of known [FeFe]-hydrogenase genes as available in databases and classified in recently published literature [21,23,24], while several previous studies in various microorganisms have focused on a single [FeFe]-hydrogenase gene only [29,32–38].

Materials and methods

Pilot-scale plant set up and operation

Dark anaerobic fermentation was carried out in a continuous stirred tank reactor (CSTR) of 35 L, equipped with pH, oxidation/reduction potential and temperature control system. The temperature control was done through the inlet of hot water in the heating jacket; to avoid stratification, the pilot reactor was provided with an external recirculating system through a peristaltic pump.

The substrate used for the fermentative process was a mixture composed of 80% of fruits and vegetables wastes and 20% of manure (20 mg/mL total suspended solids – 2% total suspended solids), supplemented with a micronutrient medium previously described [31].

The inoculum used (10%v/v) was a digested sludge coming from the municipal wastewater treatment plant of Torino, chemically pre-treated with HCl for 24 h at pH 3 to inhibit the methane forming bacteria and enrich the H₂ producing flora [39–41].

The operating volume was 25 L, nitrogen gas was injected till no oxygen was detected to create adequate anaerobic conditions, pressure inside the reactor was set at values around 20–30 mbar. The experiment was carried out at pH 5–6 and temperature between 25 and 30 °C.

The volume of gas produced was measured through a drum-type gasmeter (Dr.-Ing. Ritter Apparatebau GmbH & co.) whereas the gas composition (H₂ and CO₂ mainly) was determined by using a micro gas chromatograph (Varian, CP4900).

Isolation of cultivable bacteria

Bacteria were isolated from a sample of the digestate present in the plant during its operation. Un-treated samples were diluted and directly plated on the rich non selective “Clostridial nutrient” medium (Sigma–Aldrich) agar plates. Alternatively, to select for spore-forming microorganisms, the sample was heat-treated at 80 °C for 10 min prior to dilution. Plates were incubated under anaerobic conditions into “Anaerogen” bags (Oxoid) at 37 °C over night.

Forty single and isolated colonies with different morphology were randomly selected and pure cultures were obtained by an additional passage on plate.

Morphological characterisation was obtained using Gram staining. Discrimination between obligate and facultative anaerobes was obtained detecting the growth on plates incubated under anaerobic or aerobic conditions.
16S rDNA amplification, RFLP analysis and sequencing

Genomic DNA was extracted from colonies of each isolate by rapid freeze-thawing from sterile water [42]. 16S rDNA gene was amplified from genomic DNA by PCR using the proof-reading polymerase “KOD Hot Start DNA polymerase” (Merk Millipore) and the two universal primers 27F (5’-AGAGTTTGTATCCTGGCTCAG) and 1492R (5’-TACGGYTACCTTGTTACGACT) [42].

Restriction fragment length polymorphism (RFLP) analysis was performed using the four restriction endonucleases HaeIII, HhaI and TaqI (Thermo Scientific) that were previously reported to have a high discriminatory power [43]. The amplified 16S rDNA fragment was digested following the manufacturer’s instructions. DNA fragments were separated by electrophoresis on 1.5% w/v agarose gels in 1× TAE stained with SYBR® Safe (Invitrogen) using “PerfectSize DNA Molecular Weight 100 bp XL Ladder” (5Prime) as a reference. After classification, two isolates per group were identified by sequencing the entire PCR fragment from an external company (Eurofins MWG Operon, Germany). The sequences were compared with the NCBI database using BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi). Exact matches were found for each sequence with 100% identity.

Quantification of H2 production yield of the strains

Pure cultures of each species were grown in sealed glass vials under anaerobic argon atmosphere at 37 °C with shaking. The vial contained a medium that was previously reported [42] containing 100 mM potassium phosphate, 17 g/L tryptone, 3 g/L peptone papain digest of soybean, 10 g/L glucose, initial pH 7.0.

The gas phase was sampled with a SampleLock Gastight syringe (Hamilton) and analysed by gas chromatography, at least in three replicates. The gas chromatograph (Agilent Technologies 7890A) was equipped with purged packed inlet, Molesieve 5A column (30 m, ID 0.53 mm, film 25 mm) and thermal conductivity detector; argon was used as carrier gas.

RT-PCR

Samples were taken from the pure cultures at the time points shown in Fig. 4. Total RNA was extracted and purified using the kit “Total RNA isolation Nucleospin® RNA II” (Macherey-Nagel). Since genomic DNA contamination was often observed, the total RNA was also subjected to an additional DNaseI treatment (Sigma–Aldrich).

Reverse transcription of total RNA into cDNA was achieved with random hexamers using “Maxima H Minus First Strand cDNA Synthesis Kit” (Thermo Scientific) after normalisation of the concentration.

Specific primers were designed for each target gene using the tool “primer3” (http://emboss.bioinformatics.nl/cgi-bin/emboss/primer3) and the genome sequences deposited in the NCBI database: Clostridium beijericini (NC_009617.1), Clostridium butyricum (PRJNA54843) and Clostridium perfringens (NC_008261.1). Specificity was tested by checking the sequence of each primer against the sequence of the other [FeFe]-hydrogenase genes; moreover the PCR products have been sequenced showing the specific amplification of the target gene only.

Detection of each transcript was obtained by PCR using “KOD Hot Start DNA polymerase” and the following conditions: 95 °C for 2 min, 38 cycles of 95 °C for 20 s, 53 °C for 10 s, 70 °C for 14 s.

During RT-PCR, opportune controls were always performed as follows: genomic DNA (positive control), no template (NTC) and no reverse transcriptase (RT-). The 16S rDNA gene was also included in the RT-PCR analysis to validate the quality of the cDNA and to be used as a reference for the semi-quantitative estimate of the other transcripts.

Results

Hydrogen production via dark anaerobic fermentation in a CSTR reactor

The pilot-scale plant for H2 production from waste materials, designed and built within the “BioEnergy Lab” of the “Environment Park” in Torino, Italy, has already been described [31]. This plant showed to be efficient in the production of H2 from a number of different wastes; the biogas recovered could be directly used to generate electricity [31].

For the present study the plant was fed with a mixture of vegetable wastes from food markets supplemented with 20% manure. The inoculum was composed of sludge from a wastewater treatment plant that was pre-treated with HCl to inhibit the methanogenic microorganisms.

After a lag time of 30 h and a batch phase of 60 h, the plant was operated with a hydraulic retention time (HRT) of 30 h. The production of hydrogen and CO2 was followed as a function of time. Hydrogen percentage ranged between 20 and 50%, with a maximum value of 54–55% (Fig. 1). This value is interestingly higher in comparison to previously published percentages, typically in the range 20–40% [44,45].

![Fig. 1](image_url) - Amount and rate of hydrogen production in the pilot plant. The left vertical axis is referred to the gas production rate; the right axis is referred to the percentage of hydrogen on total gas produced (the main other gas is represented by CO2). The dashed vertical line indicates the time when the sampling of the digestate was performed for studies of microbial flora isolation.
The average hydrogen production rate was 0.9 L/h with a maximum yield of 72.6 mL H₂/g volatile suspended solids and a productivity of 0.72 (LH₂/(L culture)⁻¹ day⁻¹. No methane production was observed, demonstrating that acid pre-treatment of inoculum is an effective method to avoid methanogenesis during fermentation.

A sample of the digestate was taken (Fig. 1) at the maximum hydrogen production percentage (54% of hydrogen, 46% of carbon dioxide) with a close-to-average hydrogen production rate (0.9 L/h) for the isolation of the culturable bacteria.

**Isolation and characterisation of the culturable bacterial population**

All the isolates were classified on the basis of cell morphology after Gram staining and the ability to grow also in the presence of oxygen, thus discriminating the obligate from the facultative anaerobes (Table 1).

Subsequently, the isolates were classified in more details on a molecular basis, by means of 16S rDNA gene restriction fragment length polymorphism (RFLP) analysis which allowed the subdivision of all the isolated strains into 9 different groups (Fig. 2).

Each group was then identified by sequencing the entire 16S rDNA gene amplicon and searching against the NCBI database; exact matches were found per each sequence with 100% identity. The 9 groups were identified as 9 different species of the genus Clostridium, three species of the genus Enterococcus, one species of the genus Lactobacillus and one species of the genus Staphylococcus.

The ability to produce H₂ was assayed to investigate the direct involvement in H₂ production of each species (Table 1). This showed that only the four species of the genus Clostridium are able to produce H₂. Among them, the bacterium showing the highest hydrogen production yield is C. beijerinckii SM10, followed by C. butyricum SM32, C. perfringens SM09 and Clostridium bifermantans SM11.

**[FeFe]-hydrogenase genes expression during H₂ production**

Fermentative hydrogen production in the genus Clostridium requires the catalytic activity of hydrogenases, redox enzymes that use protons as the final electron acceptors in the cellular energy metabolism [23,46]. Analysis of the known Clostridium species genomes revealed that each species does not posses a single gene encoding for this enzyme but many genes, usually between 2 and 8, encoding for different hydrogenases [23,24]. Most of the genes are annotated to encode for [FeFe]-hydrogenases, usually involved in H₂ evolution, and only few of them encode for [NiFe]-hydrogenases, usually involved in H₂ uptake. Thus, the genome sequence analysis revealed a complex framework with many genes and enzymes potentially involved in H₂ production that so far is supported by very few experimental data concerning the physiological role of the various hydrogenases.

The genomes of the three species with the highest hydrogen production yield isolated here have been previously sequenced: C. beijerinckii NCIMB 8052, C. butyricum 5521 and C. perfringens ATCC 13124. Altogether, they have annotation of 14 genes encoding for different [FeFe]-hydrogenases (Fig. 3) that belong to different modular structure and phylogenetic classification [23,24]. Most genes encode for monomeric enzymes, while only three of them encode for heterotrimERIC enzymes. All the gene products are predicted to be cytoplasmic, with the exception of that belonging to cluster A3 (structure TR(M2)). It has to be noted that only C. beijerinckii has also a gene annotated to encode for a [NiFe]-hydrogenase.

Interestingly, most of these genes or enzymes have never been studied in details before. C. butyricum and C. perfringens have a gene encoding for an [FeFe]-hydrogenase of the M3 structure; enzymes from this group (CpI and CaHydA) are known to be directly involved in H₂ production and they have been previously characterised in details in Clostridium pasteurianum [47–50] and Clostridium acetobutylicum [20,33,51,52] respectively. On the contrary, C. beijerinckii lacks enzymes of this well known type.

In order to study the transcriptional levels of each of these genes in hydrogen production in the three strains isolated here, their expression at mRNA level was studied by RT-PCR during the growth of pure cultures.

Interestingly, H₂ evolution occurs mainly during the exponential growth phase in C. beijerinckii SM10 and C. butyricum SM32, while it is mainly observed during the late exponential and stationary phase in C. perfringens SM09 (Fig. 4), thus suggesting important differences in the role of H₂ production in the metabolism. On the basis of the relationship

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**Table 1 – Summary of the culturable strains identified in the pilot plant. Each species was identified by 16S rDNA gene sequencing. Cell morphology and the presence of spores were determined by Gram staining of the isolated strains. The ability to grow in the presence of oxygen allowed discrimination between obligate and facultative anaerobes. The ability of each species to produce H₂ was determined. Strains 1–4 were isolated from heat-treated samples, while strains 5–9 were isolated from un-treated samples.**

<table>
<thead>
<tr>
<th>ID</th>
<th>Species</th>
<th>Strain</th>
<th>Cell morphology</th>
<th>Spores</th>
<th>Anaerobe</th>
<th>mL H₂/g glucose</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Clostridium beijerinckii</td>
<td>SM10</td>
<td>Bacilli, Gram +</td>
<td>–</td>
<td>Obligate</td>
<td>247.2 ± 9.6</td>
</tr>
<tr>
<td>2</td>
<td>Clostridium butyricum</td>
<td>SM32</td>
<td>Bacilli, Gram +</td>
<td></td>
<td>Obligate</td>
<td>136.3 ± 4.6</td>
</tr>
<tr>
<td>3</td>
<td>Clostridium perfringens</td>
<td>SM09</td>
<td>Bacilli, Gram +</td>
<td>–</td>
<td>Obligate</td>
<td>130.6 ± 3.0</td>
</tr>
<tr>
<td>4</td>
<td>Clostridium bifermantans</td>
<td>SM11</td>
<td>Bacilli, Gram +</td>
<td></td>
<td>Obligate</td>
<td>97.1 ± 9.2</td>
</tr>
<tr>
<td>5</td>
<td>Lactobacillus plantarum</td>
<td>SM21</td>
<td>Bacilli, Gram +</td>
<td>–</td>
<td>Facultative</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>Enterococcus sp.</td>
<td>SM01</td>
<td>Cocci, Gram +</td>
<td>–</td>
<td>Obligate</td>
<td>0</td>
</tr>
<tr>
<td>7</td>
<td>Enterococcus devriesi</td>
<td>SM03</td>
<td>Cocci, Gram +</td>
<td>–</td>
<td>Facultative</td>
<td>0</td>
</tr>
<tr>
<td>8</td>
<td>Enterococcus sp.</td>
<td>SM08</td>
<td>Cocci, Gram +</td>
<td>–</td>
<td>Facultative</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>Staphylococcus hominis</td>
<td>SM17</td>
<td>Cocci, Gram +</td>
<td>–</td>
<td>Facultative</td>
<td>0</td>
</tr>
</tbody>
</table>

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between growth and H₂ production, time points were selected for the study of [FeFe]-hydrogenase expression.

Specific primer couples targeting each gene were designed (Table 2) on the basis of the sequenced genomes and tested with the genomic DNA of *C. beijerinckii* SM10, *C. butyricum* SM32 and *C. perfringens* SM09. PCR fragments with the expected length could be amplified; DNA sequencing of the PCR product demonstrated the specificity and efficiency of the amplification and also the presence of each gene in the genome of the strains isolated from the plant (data not shown).

RT-PCR (Fig. 3) showed that transcripts of all the [FeFe]-hydrogenases analysed could be detected. Moreover, the transcriptional profiles were interesting: most genes were constitutively expressed and only few of them showed a modulation.

**Discussion**

The culturable part of the bacterial population present in a bio-hydrogen pilot-scale plant at the maximum H₂ production percentage was characterised, revealing the presence of microorganisms belonging to the genera *Clostridium*, *Lactobacillus*, *Enterococcus* and *Staphylococcus*.

The involvement of *Clostridium* species in H₂ production is well known and they are reported to be able to use several waste matrices [1,53]. Various strains of *C. beijerincki* were isolated from several sources but, interestingly, the hydrogen production yield of the strain isolated here is higher than those previously reported [54–56]. Moreover *C. beijerincki* is one of the major producers of butanol, an important solvent that can be used as biofuel [57], thus suggesting another possibility to recover simultaneously different valuable products from wastes.

In the culturable part of the bacterial population, the presence of several species that are not directly involved in H₂ production reveals that a complex consortium of microorganisms with different roles was spontaneously selected in the pilot plant, giving place to efficient matrix degradation and H₂ production.

*Lactobacillus plantarum* has been found in a wide range of natural and artificial fermentation processes, in particular in plant-derived matrices [58]. It is a very versatile microorganism that produces large amount of lactic acid and causes acidification of the fermentation medium [59], thus favouring the hydrogen producing microorganisms versus the methanogenic ones.

Also *Enterococcus* species are found in fermentation processes, including food products, and have a metabolism that results in lactic acid production and acidification [60]. Interestingly, a consortium for hydrogen production between *C. butyricum* and *Enterococcus saccharolyticus* was previously reported [61].

It is also interesting to consider that *L. plantarum* produces various bacteriocins that can inhibit the growth of several microorganisms, including *Clostridium*, *Enterococcus* and *Staphylococcus* species; importantly, many strain-to-strain differences have been reported [59,62–64]. Also various *Enterococcus* species are reported to produce bacteriocins [60,65].

In this case, the coexistence of *L. plantarum* and *Enterococcus* species with various *Clostridium* species suggests that the strains isolated here interact favourably between them and, conversely, that the bacteriocins potentially produced by *L. plantarum* [59,62] and *Enterococcus* sp. [60] might act to select the specific strains within the consortium.

The involvement of [FeFe]-hydrogenase in hydrogen production in *Clostridium* is known [23], but the specific investigation of the role of different [FeFe]-hydrogenases has never been performed before. Here, the study of the expression at the mRNA level by RT-PCR was performed on pure cultures of the isolated *C. beijerincki* SM10, *C. butyricum* SM32 and *C. perfringens* SM09 during the growth, showing that all the known
genes (6, 4 and 4, respectively) are expressed. The constitutive expression of [FeFe]-hydrogenase genes at the mRNA level is not simple to interpret, especially for the time points where the amount of accumulated H₂ does not change, suggesting a post-transcriptional down-regulation of the functionality or an equilibrium between H₂ evolution and uptake.

*C. beijerinckii* SM10 shows the most complex behaviour, with two genes constitutively expressed in all the four time points: Cbei_0327 (M3α' structure) and Cbei_4110 (TR(M3) structure).

Interestingly, the other four genes of *C. beijerinckii* are variants of the M2 modular structure (M2a, M2b, M2c and TR(M2)) and they share a common behaviour in terms of transcripts: they are all down-regulated during the stationary phase.

Cbei_4110 was previously reported to be expressed in *C. beijerinckii* RZF-1108 [38] but surprisingly, in a recent genome-

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### Table 3

<table>
<thead>
<tr>
<th>Expression</th>
<th>Gene</th>
<th>Structure</th>
<th>Enzyme domains composition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>16S</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cbei_1901</td>
<td>M2a</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cbei_4000</td>
<td>M2b</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cbei_1773</td>
<td>M2c</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cbei_0327</td>
<td>M3α'</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cbei_3796</td>
<td>TR(M2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cbei_4110</td>
<td>TR(M3)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

Fig. 3 – [FeFe]-hydrogenase genes expression in *C. beijerinckii* SM10, *C. butyricum* SM32 and *C. perfringens* SM09. The expression was studied by RT-PCR, using the 16S gene as a reference. The modular structure classification and the domains composition refer to the enzyme in accordance with the classification proposed before [23].

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wide transcriptional analysis of C. beijerinckii NCIMB 8052, there is no mention of any hydrogenase genes expression levels [66].

C. butyricum SM32 has three genes constitutively expressed: CBY_2300 (M3 structure), CBY_3049 (M3a structure) and CBY_2047 (TR(M3) structure), while CBY_2676 (M2a structure) is gradually down regulated.

CBY_2300 belongs to the M3 structure (group A2) and the encoded protein has high sequence similarity with the well characterised CpI and CaHydA (67.5% and 66.2% identity, respectively); CBY_2300 expression was previously reported both in experiments with pure [36] and mixed [29,34] cultures. Moreover, CBY_2300 has already been demonstrated to encode for a functional [FeFe]-hydrogenase enzyme that can be heterologously expressed both in Escherichia coli and C. acetobutylicum [37].

Cbei_4110 and CBY_2047 share a common transcriptional regulation and a high sequence identity (63.8%). The enzymes encoded by these genes are considered to include also other two subunits found in the same operon [23], resulting to be similar to the so-called heterotrimeric bifurcating hydrogenases, recently identified in other anaerobic species [67–69]. These enzymes can synergistically couple the oxidation of NADH with the oxidation of ferredoxin, yielding H2 as a final product, and allowing a fine tuning of the electron fluxes within the cell [67]; nevertheless, it must be noticed that bifurcating [FeFe]-hydrogenases were also reported to be able to catalyse the inverse reaction of H2 uptake coupled to ferredoxin and NADH reduction [68]. Thus the exact physiological role of Cbei_4110 and CBY_2047 remains to be experimentally demonstrated.

Despite the different kinetics of H2 production, C. perfringens SM09 genes are similar to those of C. butyricum SM32 in terms of classification and they share a similar transcriptional profile: CPF_2655 (M3 structure), CPF_2900 (M3a structure) and CPF_1076 (M2d structure) are constitutively expressed, while CPF_0270 (M2a structure) is gradually down regulated.

Like CBY_2300, also CPF_2655 belongs to the M3 structure (group A2) and the encoded protein has high sequence similarity with the well characterised CpI and CaHydA (68.8% and 68.7% identity, respectively). The enzyme was never studied, but the gene was characterised in details: it is always expressed during the growth on glucose, in accordance with the results presented here, and found into two different transcripts [32]. Moreover, the disruption of CPF_2655 abolishes H2 productivity in C. perfringens str. 13, demonstrating its direct involvement in H2 production [32].

It is to be noticed that genes belonging to the same modular structure and same phylogenetic group share similar transcriptional levels even in different microorganisms, such as

Table 2 – List of the primers designed for RT-PCR with the size of the amplified fragment.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward</th>
<th>Reverse</th>
<th>Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cbei_1773</td>
<td>GTATTGGATGCGGAGCTTGT</td>
<td>TCCTGGTTCAAAGCCAAAAG</td>
<td>260 bp</td>
</tr>
<tr>
<td>Cbei_0327</td>
<td>TCATTTTCAACAGGCTCCAT</td>
<td>CCACCCTGAGATCTTTACTGG</td>
<td>770 bp</td>
</tr>
<tr>
<td>Cbei_1901</td>
<td>CAGTTTACATTCTGCTGCTC</td>
<td>TCTAGCTATTCTTTATCTGG</td>
<td>484 bp</td>
</tr>
<tr>
<td>Cbei_4110</td>
<td>CAGCGAATCTTCTGCTGCTC</td>
<td>CCACCACCTGATACTACC</td>
<td>381 bp</td>
</tr>
<tr>
<td>Cbei_3796</td>
<td>AGCCAGTCTCTGTCTGCTGCT</td>
<td>TCTAGCTATTCTTTATCTGG</td>
<td>683 bp</td>
</tr>
<tr>
<td>Cbei_4000</td>
<td>GAGCCAGTCTCTGTCTGCTGCT</td>
<td>TTATTACCCTGCTTCTTAC</td>
<td>279 bp</td>
</tr>
<tr>
<td>CBY_3049</td>
<td>CCACCAGTCTCTGTCTGCTGCT</td>
<td>ACCTATTGCAACACCCCTTC</td>
<td>246 bp</td>
</tr>
<tr>
<td>CBY_2300</td>
<td>ATGGGATGCTGCTGCTGCTGCTGCT</td>
<td>AGTGTGCTTTCTCTTCTTAC</td>
<td>312 bp</td>
</tr>
<tr>
<td>CBY_2047</td>
<td>CAGACAGTCTCTGTCTGCTGCTGCT</td>
<td>CCAAAATTTTACATTTGTTGAG</td>
<td>401 bp</td>
</tr>
<tr>
<td>CBY_2676</td>
<td>GACTGTGCTCTGTCTGCTGCTGCTGCT</td>
<td>TAACTTTTTTCTGACCACCA</td>
<td>514 bp</td>
</tr>
<tr>
<td>CPF_2655</td>
<td>AACCGAGTCTCTGTCTGCTGCTGCTGCT</td>
<td>TTACTACTACGCGAAG</td>
<td>281 bp</td>
</tr>
<tr>
<td>CPF_1076</td>
<td>GAGGCAGTCTCTGTCTGCTGCTGCTGCT</td>
<td>CCTCTCTCTTACCAGATTA</td>
<td>331 bp</td>
</tr>
<tr>
<td>CPF_0270</td>
<td>ATGGGATGCTGCTGCTGCTGCTGCTGCT</td>
<td>CCAACACCTGCTTCTTCTTAC</td>
<td>500 bp</td>
</tr>
<tr>
<td>CPF_2900</td>
<td>ATGGGATGCTGCTGCTGCTGCTGCTGCT</td>
<td>CTTGCAAGCTTTCTTCTTCTTCT</td>
<td>769 bp</td>
</tr>
</tbody>
</table>
efficient matrix degradation and H₂ production. Spontaneously selected within the plant, participating to a complex consortium of bacteria belonging to the genera C. beijerinckii, Lactobacillus, Enterococcus and Staphylococcus was isolated and characterised, revealing that a percentage was isolated and characterised, revealing that a functional hydrogenase for each strain, as presented here, brings the simultaneous study of the expression of all the [FeFe]-hydrogenase genes for each strain, as presented here, brings a novel contribution to the field and gives the first insights on the complex metabolic network of the [FeFe]-hydrogenase function.

Conclusions

The culturable part of the bacterial population present in a bio-hydrogen pilot-scale plant at the maximum H₂ production percentage was isolated and characterised, revealing that a complex consortium of bacteria belonging to the genera *Clostridium*, *Lactobacillus*, *Enterococcus* and *Staphylococcus* was spontaneously selected within the plant, participating to efficient matrix degradation and H₂ production.

The characterisation of the isolated *Clostridium* strains and the simultaneous study of the expression of all the [FeFe]-hydrogenase genes for each strain, as presented here, brings a novel contribution to the field and gives the first insights on the complex metabolic network of the [FeFe]-hydrogenase function.

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References

[27] Schmidt O, Drake HL, Horn MA. Hitherto unknown [FeFe]-hydrogenase gene diversity in anaerobes and anoxic

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