Physiological characteristics of the extreme thermophile \textit{Caldicellulosiruptor saccharolyticus}: an efficient hydrogen cell factory

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Abstract

Global concerns about climate changes and their association with the use of fossil fuels have accelerated research on biological fuel production. Biological hydrogen production from hemicellulose-containing waste is considered one of the promising avenues. A major economical issue for such a process, however, is the low substrate conversion efficiency. Interestingly, the extreme thermophilic bacterium \textit{Caldicellulosiruptor saccharolyticus} can produce hydrogen from carbohydrate-rich substrates at yields close to the theoretical maximum of the dark fermentation process (i.e., 4 mol \text{H}_2/mol hexose). The organism is able to ferment an array of mono-, di- and polysaccharides, and is relatively tolerant to high partial hydrogen pressures, making it a promising candidate for exploitation in a biohydrogen process. The behaviour of this Gram-positive bacterium bears all hallmarks of being adapted to an environment sparse in free sugars, which is further reflected in its low volumetric hydrogen productivity and low osmotolerance. These two properties need to be improved by at least a factor of 10 and 5, respectively, for a cost-effective industrial process. In this review, the physiological characteristics of \textit{C. saccharolyticus} are analyzed in view of the requirements for an efficient hydrogen cell factory. A special emphasis is put on the tight regulation of hydrogen production in \textit{C. saccharolyticus} by both redox and energy metabolism. Suggestions for strategies to overcome the current challenges facing the potential use of the organism in hydrogen production are also discussed.

1. Introduction

Anthropogenic CO\textsubscript{2} emissions have generally been recognized as the major contributor to global warming and associated climate changes. Therefore, several measures are being taken to decrease the CO\textsubscript{2} emission. In recent years, much effort has been devoted to rendering biofuel production economically competitive to that of fossil fuels, since this will contribute significantly to the reduction of energy-linked environmental impacts. In this quest, the choice of the raw material is of central concern. First-generation biofuels are produced from sucrose and starch-rich substrates, which may compete with human consumption - inevitably driving up market prices. As a remedy, more focus should be directed to second-generation biofuels, produced from non-edible lignocellulosic materials, the most naturally abundant raw material [1], as well as domestic and industrial wastes. The accompanying significant cost reductions should make biofuels more competitive.

Biohydrogen is a typical example of an environmentally friendly biofuel, with no CO\textsubscript{2} emission resulting from its combustion. It can be produced from both lignocellulosic and waste materials [2-5], through biological conversion processes, such as dark fermentation and photofermentation. In the latter, biohydrogen can be produced using purple sulphur or non-sulphur bacteria that convert organic acids to \text{H}_2 in photon-driven reactions [6,7]. Although a combination of these two processes is an interesting approach for maximum conversion of the energy contained in carbohydrate-rich substrates into \text{H}_2 [8], only dark fermentative \text{H}_2 production is covered in this review.

In total, 12 \text{H}_2 molecules can be obtained per mole of glucose, based on the overall number of electrons that can be generated in the complete oxidation of the latter. However, dark fermentation is limited to a maximum
H₂-production efficiency of 33%, i.e., maximally four molecules of H₂ can be acquired per molecule of glucose with acetate and CO₂ as the other fermentation end products [9]. Yet, this is only possible when the H₂ partial pressure (P_{H₂}) is kept adequately low [10], e.g. by continuous stripping of the produced H₂ with an inert gas. However, for a cost-effective dark fermentation process it is vital to obtain significantly high H₂ yields at relatively elevated P_{H₂} due to the high impact of central costs of feedstock and gas upgrading [11]. Generally, mesophilic (co-)cultures reach H₂ yields of ≤ 2 moles/mol hexose [12], thus exemplifying conversion efficiencies of merely 17%. In addition, these yields are obtained at low P_{H₂} only [6]. On the other hand, based on thermodynamic aspects, thermophilic bacteria and archaea may produce up to the theoretical maximum of 4 mol H₂/mol hexose [13]. In general, the low H₂ yields obtained in practice by different organisms, in addition to the requirement for low P_{H₂}, are major obstacles that need to be overcome before biohydrogen production can be industrially feasible [6].

*Caldicellulosiruptor saccharolyticus* is an extreme thermophilic bacterium that can produce high H₂ yields [14,15], and at the same time is relatively insensitive to high P_{H₂} [16]. This organism has recently gained increased interest due to its ability to produce thermostable cellulolytic and xylanolytic enzymes [17-21], to grow on complex lignocellulosic carbon sources [2,22,23], and to co-metabolize a wide spectrum of monosaccharides including both pentose and hexose sugars [24]. Although *C. saccharolyticus* possesses these desirable traits and several research articles have been recently published on this heterotrophic strict anaerobe, it has not yet been the subject of a review article. Herein, the physiological properties of *C. saccharolyticus* will be analyzed to unravel the characteristics required for a superior H₂ cell factory. To further underline current insights in this topic, the physiology of *C. saccharolyticus* will be compared with that of other H₂-producing microorganisms.

2. Characteristics and natural habitat of *C. saccharolyticus*

*C. saccharolyticus* is a strictly anaerobic, extreme-thermophilic, Gram-positive bacterium that belongs to the Clostridia class. There are to date 13 characterized species in the genus *Caldicellulosiruptor*, of which *C. saccharolyticus* is described best, including its whole-genome sequence [25]. *C. bescii*, formerly known as *Anaerocellum thermophilum*, is the other species in the genus with its genome also fully sequenced [26,27]. *C. saccharolyticus* was isolated from a piece of wood in a hot spring in the Rotorua-Taupo thermal area in New Zealand, of which the waters have neutral pH, and are low in sulphate, salt content and free sugars [28]. Instead, The majority of the sugars present in this environment, such as glucose and xylose, are contained in complex (hem)cellulosic polymers, which are difficult to access.

*C. saccharolyticus* is well adapted to the selective pressure of this habitat through the production of various hydrolyzing enzymes, possession of a wide range of high-affinity transport systems, and absence of glucose-based catabolite repression [24,25,29], which are all attractive properties for using the organism for industrial purposes. However, being sensitive to high osmotic pressures [30,31] is one of the undesirable consequences of adaptation to such environment. Interestingly, *C. saccharolyticus* is non-motile and non-sporulating [32], although it possesses a complete set of genes for flagella biosynthesis and several genes required for sporulation [25], signifying a historical genetic record of characteristics that could have been required in past habitats.

3. Genome analysis and carbohydrate utilisation by *C. saccharolyticus*

The genome of *C. saccharolyticus* has been sequenced and a microarray-based protocol for genome-wide transcriptome analysis has been developed, which is a major convenience for studying and understanding its physiology [25]. *C. saccharolyticus* has a relatively small genome of 2,970 kb and 2,760 protein-coding open reading frames (ORFs). From the genome-curation project, several new insights were gained such as the unusually high number of transposons and transposable derivatives *C. saccharolyticus* possesses, as compared with other organisms [25]. Since transposition increases the chance of genetic alterations, a large number of transposons can enhance the adaptability of this organism and may widen the spectrum of environmental conditions it can grow in, which is beneficial from an industrial point of view. On the other hand, this may also have negative repercussions since it can jeopardize stable fermentation processes. How to balance both impacts in practice needs to be further investigated.

A general favourable property of members of the Clostridia class is their ability to degrade complex polymers such as cellulosic and lignocellulosic materials [33]. Based on growth analysis, *C. saccharolyticus* can metabolize various carbon sources ranging from monomers, such as xylose, arabinose, glucose, fructose and galactose to α- and β-linked di- and polysaccharides, such as maltose, lactose, sucrose, starch, pullulan, threhalose, xylan and cellulose [32]. *C. saccharolyticus* can also grow and produce H₂ from complex lignocellulosic materials, both pre-treated, such as Miscanthus hydrolysate [2], sugar beet juice [34] and paper sludge [23], and untreated, such as wheat straw [35], pine wood [22] and bagasse [35].
The fermentation of these raw materials by *C. saccharolyticus* has yielded H2, CO2 and acetate as the main metabolic end products [2,23,34,35], with a maximum H2 yield per consumed hexose of 3.0-3.8 mol/mol [2,34,35]. Interestingly, cultivations on Miscanthus hydrolysate were accompanied by complete depletion of oligosaccharides and simultaneous oligosaccharide and monosaccharide consumption [2]. *C. saccharolyticus* degrades complex polymers by producing several cellulolytic and xylanolytic enzymes [21]. Properties of the cellulase and hemicellulase enzymes of this organism as well as their gene structures have been extensively studied [21]. In contrast to cellulolytic clostridia [33], *C. saccharolyticus* does not contain a cellulosome complex. *C. saccharolyticus* is unusual with respect to its possession of a multi-functional, multi-domain organisation for the majority of its β-glucanases [29]. The majority of the xylanases in *C. saccharolyticus* are clustered in a large gene cluster, and another cluster of genes involved in xylose metabolism is in close proximity [21]. Detailed information on glycoside hydrolases and other carbohydrate active enzymes encoded by *C. saccharolyticus* genome can be retrieved from the Carbohydrate Active Enzymes database (CAZy; http://www.cazy.org; [36]).

The manual curation of the genome sequence of *C. saccharolyticus* indicated that it possesses an unusually high number of ATP-binding cassette (ABC) transporter genes (i.e., at least 177 ABC transporter genes) [25]; many of which are responsible for the transport of monomeric and dimeric sugars. *C. saccharolyticus* has only one sugar phosphotransferase system (PTS), which is believed to be used for fructose transport [24,25]. When *C. saccharolyticus* was cultivated on a mixture of monosaccharides, they were consumed simultaneously but at different rates, i.e., fructose > arabinose > xylose > mannose > glucose > galactose. Unlike the case with disaccharides, the ABC transporters were not substrate specific for these monosaccharides. Instead, the affinity for the different sugars varied, with a clear preference for pentose sugars. Overall, the genome and transcription analyses revealed that *C. saccharolyticus* is well equipped for utilizing complex carbohydrates, such as starch and lignocellulosic polysaccharides [25].

4. Hydrogen yields

For cost-efficient biohydrogen production, it is critical to achieve high substrate-conversion efficiencies due to the central cost of the raw material and the need to reduce the chemical oxygen demand (COD) in the waste stream [6]. Strategies to improve the overall process yield include the application of hybrid two-step systems, consisting of a dark fermentation step combined with either a light fermentation step [37], or a methanogenesis step [4,38]. In this way, the products of the first fermentation process, consisting mainly of volatile fatty acids, can be converted in the second step to either H2 and CO2 or CH4 and CO2, respectively. The pros and cons of these two strategies were evaluated in a recent review [6] and will not be treated herein. Instead, we will focus on H2 yields of the dark fermentation step.

4.1 Metabolic pathways for hydrogen production

During fermentation, H2 is produced by anaerobic heterotrophs as a means to reoxidize their reducing equivalents. The maximum obtainable yield in a dark fermentation process is 4 moles H2 per mol of hexose, the so-called Thauer limit [9]. However, many H2 yields reported in literature are less than 50% of the Thauer limit (Table 1).

The maximum H2 yield obtained during sugar oxidation depends primarily on the catabolic pathway employed. *C. saccharolyticus*, with its H2 yields close to the Thauer limit [9], employs the Embden-Meyerhof-Parnas (EMP) pathway to oxidize glucose to pyruvate [14]. The genome sequence and 13C-NMR analysis revealed a complete gene setup for the EMP pathway and the non-oxidative part of the pentose phosphate pathway (PPP) [14,25], which enables the organism to convert different sugars into precursor metabolites and energy. No essential genes for the Entner-Doudoroff (ED) pathway have been found [14,25]. Pyruvate can be oxidized further to acetyl-CoA by pyruvate:ferredoxin oxidoreductase (PFOR), with the generation of reduced ferredoxin (Fdred). Molecular H2 can be obtained from both NADH (two moles of which are generated through the glyceraldehyde phosphate dehydrogenase (GAPDH)-catalyzed reaction in the EMP pathway) and Fdred (total 8 mol H2/mol hexose). Unlike pyruvate oxidation to acetate. The produced Fdred (total 8 mol H2/mol hexose) can subsequently donate electrons to hydrogenases that reduce protons to H2 [39].

On the other hand, enterobacteria are only able to produce maximally 2 moles H2/mol hexose, since H2 is generated via the hydrolysis of formate being produced from glucose through the pyruvate formate lyase-catalyzed pyruvate oxidation (Figure 1C). Interestingly, even the introduction of a synthetic ferredoxin-mediated pyruvate-to-H2 pathway in *E. coli* resulted in low H2 yields (i.e., ~2 mol/mol hexose) [40]. Some of the possible
reasons why mesophilic bacteria can produce only 2 moles H2/mol hexose, despite being equipped with the relevant metabolic pathways, are revealed in the following sections.

4.2 Thermodynamic aspects

One of the reasons to which the low H2 yields observed in mesophilic cultures have been attributed is the thermodynamically unfavourable NADH-dependent H2-evolving reaction at \( P_{H2} > 0.039 \text{ kPa} \) and at standard conditions (1 M concentration of the reactants, 25°C, pH 7) [41]. This is because H2 has a lower formal potential than NADH, hence for the reduction of protons by NADH to occur spontaneously, the \( P_{H2} \) has to be low. However, these conditions are often not encountered even when the reactor is sparged with an inert gas at a high flow rate (i.e., 400 mL/min), because the H2 concentration in the liquid will exceed the critical \( P_{H2} \) a 1000-fold [42]. Therefore, in addition to H2, mesophilic anaerobes normally form other reduced compounds, such as acetone, butanol, ethanol, and lactate, resulting in a H2 yield that usually does not exceed 2 mol/mol hexose (Table 1).

A strategy to overcome this problem is to use thermophilic hydrogen producers instead, hence this works a decrease in the Gibbs free energy change of the H2-generation reaction in hand (\( \Delta G'_f \); [10]):

\[
\Delta G' = \Delta G^{0'} + RT \cdot \ln \left( \frac{NAD^+}{NADH} \cdot \frac{P_{H2}}{H^+} \right)
\]  

where \( \Delta G^{0'} \) is the Gibbs free energy at standard conditions (kJ/mol), \( R \) is the gas constant (kJ/mol·K) and \( T \) the absolute temperature (K).

In general, extreme thermophiles reach higher H2 yields than mesophiles (Table 1) [13], and also at higher temperatures the critical \( P_{H2} \) is higher. For instance, at a temperature of 70°C the critical \( P_{H2} \) is 0.173 kPa. However, this value is still well below 60 kPa, at which in practice \( C. \ saccharolyticus \) is still capable of producing 3.6-4.0 moles H2/mol glucose [16].

To understand this controversy, it is to be noticed that temperature is not the only factor influencing H2 generation. The \( \Delta G' \) of the H2-producing process is also a function of the type of cofactor involved and the ratio of reduced to oxidized cofactors [43]. Each cofactor has a different formal potential, i.e., the \( E^{0'} \) for Fdred/Fdox \(<\) NADPH/NADP \(<\) NADH/NAD [36,38,39], and unless the cofactor specificity of the involved hydrogenase is known, it is not possible to determine the thermodynamics of the system. \( C. \ saccharolyticus \) possesses two distinct hydrogenases, i.e., one NADH-dependent Fe-only hydrogenase (Csac_1860-1864) and one Fd-dependent, membrane-associated NiFe hydrogenase (Csac_1540-1545) [26]. Given that the formal potential of Fdred/Fdox (\( E^{0'} \) approx. -400 mV) is close to that of H2 (\( E^{0'}_{H2} \) : -414 mV) [41], the Fd-dependent hydrogenase reaction is energetically favourable up to a \( P_{H2} \) of 39 kPa at standard conditions. However, this calculation is based on an approximation since the actual midpoint redox potential of Fd depends on the strain and the

| Table 1 Effect of growth temperature on H2 yield |
| --- |
| **Organism** | **Opt T °C** | **Cultivation** | **Substrate** | **Maximum yield** | **Reference** |
| **Mesophiles** | | | | | |
| *Escherichia coli* SR15 | 37 | Fed-batch | Glucose | 1.82 | [85] |
| *Enterobacter aerogenes* | 40 | Batch | Glucose | 1.0 | [99] |
| *Enterobacter cloacae* | 37 | Packed bed | Starch | 1.4 | [100] |
| *Clostridium butyricum* | 30 | continuous | Glucose | 2 | [101] |
| *Clostridium acetobutylicum* | 34 | Fed-batch | Glucose | 2 | [102] |
| **Thermophiles** | | | | | |
| *Clostridium thermocellum* | 60 | Continuous | | 3.5 | [103] |
| **Extreme thermophiles** | | | | | |
| *Thermotoga neapolitana* | 77 | Batch | Glucose | 3.9 | [104] |
| *Thermotoga elii* | 65 | Batch | Glucose | 3.3 | [15] |
| *Caldicellulosiruptus saccharolyticus* | 70 | Continuous | Glucose | 3.6 | [14] |
| *Caldicellulosiruptor Owensensis* | 70 | Batch | Glucose | 4 | [66] |
| *Thermoanaerobacter tengcongensis* | 75 | Batch | Glucose | 4 | [45] |
| *Thermoanaerobacter pseudethanolicus* | 70 | Continuous | Xylose | 1.2 | [105] |
| **Hyper thermophiles** | | | | | |
| *Thermotoga maritima* | 80 | Batch | Glucose | 4 | [60] |
| *Pyrococcus furiosus* | 100 | Batch | Glucose | 3.5 | [57] |
| *Thermococcales kodakaraensis* | 88 | Continuous | Starch | 3.3 | [106] |

A selection of mesophilic, thermophilic and extreme thermophilic H2-producing strains, showing their optimal growth temperature and H2 yields.
temperature [44]. The actual midpoint redox potential for Fd at 70°C in *C. saccharolyticus* is not known, but should be in the same order of magnitude as found in other related thermophilic bacteria. From the thermodynamic estimation, it is obvious that it is primarily the NADH-dependent Fe-only hydrogenase that is inhibited by elevated $P_H2$. Consistently, the NADH-dependent Fe-only hydrogenase in *Thermoanaerobacter*...
C. saccharolyticus was downregulated at high $P_{112}$, while the Fe-dependent hydrogenase was constitutively expressed, independently of the $P_{112}$ [45].

Recently, Schut and Adams [46] characterized a novel type of Fe-only hydrogenase in *Thermotoga maritima* (TM1424-TM1426) that uses NADH and Fe$_{red}$ simultaneously. This novel bifurcating hydrogenase could therefore catalyze the unfavourable oxidation of NADH to H$_2$ by using the exothermic oxidation of ferredoxin as a driving force. Interestingly, the Fe-only hydrogenase genes in *C. saccharolyticus* (Csac_1860, 1863-1864) show $\geq 50\%$ sequence similarities to the bifurcating hydrogenase genes in *Tm. maritima* (TM1424-TM1426), but it remains to be investigated whether this hydrogenase in *C. saccharolyticus* possesses a bifurcating function. In fact, many of the thermophiles listed in Table 1 and several mesophilic clostridia possess a Fe-only hydrogenase-encoding genes with sequence similarity to TM1424-TM1426 in *Tm. maritima* [46]. This observation could give a plausible explanation to the ability of these thermophiles to produce high H$_2$ yields in well-sparged bioreactors.

The ratio of reduced to oxidized cofactors has also a significant impact on the thermodynamics of the system (Eq. 1), which was confirmed experimentally by Velt et al [43]. In *C. saccharolyticus*, the NADH/NAD ratio changed six folds during growth-phase transition in batch cultures [47], and consequently the value of the critical $P_{112}$ for hydrogen production will also vary with the growth state of the organism. Similar observations were reported for *Clostridium cellulolyticum* [48], *Cl. butyricum* [49] and *Cl. acetobutylicum* [50]. Therefore, based on the reasoning above, the thermodynamic constraint might not be an absolute parameter for determining the critical $P_{112}$ for hydrogen production.

However, the generally higher H$_2$ yields obtained by thermophilic organisms still implies that temperature has a significant impact. An intriguing question that arises is why these thermophiles have evolved for optimized H$_2$ production. Resolving this phenomenon would aid in designing highly efficient H$_2$ cell factories.

### 5. Effect of $P_{112}$ versus dissolved H$_2$ concentration

As discussed above, due to gaps in the knowledge on the redox system in *C. saccharolyticus*, the thermodynamic constraint cannot be used to estimate its tolerance to H$_2$. For practical reasons, the $P_{112}$ in the gas phase is generally used as a measure for H$_2$ tolerance. The critical value of $P_{112}$ (10-20 kPa) as found for *C. saccharolyticus* is often quoted in the literature, and is defined as the partial pressure of H$_2$ at which lactate formation, an alternative way for reoxidizing NADH, is initiated [30]. This critical $P_{112}$ was determined in batch cultures of *C. saccharolyticus* on sucrose using a closed bioreactor without sparging. However, in a similar experimental setup, but with xylose as the substrate, the critical $P_{112}$ was up to 60 kPa, upon which the metabolism shifted to lactate [16]. A possible explanation for the observed discrepancy can be based on the difference in the organism’s metabolic activity on the two substrates, i.e., the volumetric H$_2$ productivity is lower on xylose than on sucrose which results in different concentrations of dissolved H$_2$ [16]. The variation in the observed critical $P_{112}$ values can also be a consequence of a different distribution pattern of catabolic and anabolic fluxes on the two substrates, leading to different metabolite levels that modulate lactate dehydrogenase (LDH) activity [47]. In that sense, the critical $P_{112}$ has no strict value, but will depend on the organism, the substrate and probably other environmental factors of the fermentation process. To make it more complicated, in batch cultures on glucose (5 g/L), lactate formation is triggered at the transition to the stationary phase, even though the $P_{112}$ was low (6.3 kPa) [47]. Therefore, there are obviously other factors that initiate lactate formation [47], making its use as a proper criterion for defining the critical $P_{112}$ questionable. Despite lactate being formed, growth still continues, but rather in a linear than an exponential fashion [16]. In addition, in continuous cultures of *C. saccharolyticus* on glucose, growth and H$_2$ production were still observed at a $P_{112}$ of 67 kPa at a low dilution rate (0.05 h$^{-1}$) [47].

Kraemer et al [51] could not find any coherent correlation between the sparging gas flow rate and H$_2$ yield under different fermentation conditions. Due to this ambiguity, one may wonder whether $P_{112}$ is an appropriate parameter for inferring the critical H$_2$ concentration for growth and hydrogen production. Indeed, Pauss et al [52] demonstrated in various bioreactor systems that the dissolved H$_2$ easily supersaturates the liquid phase and its concentration is far from equilibrium with the gas phase. This can be attributed to the low solubility of H$_2$ in water and that H$_2$ production takes place in the liquid phase. The dissolved H$_2$ concentration is a function of H$_2$ productivity and the mass transfer rate [16], where the latter is a function of gas sparging rate and the stirring rate. A study with *C. saccharolyticus* in a pH and temperature controlled stirred tank reactor confirmed that the dissolved H2 concentration is a more indicative parameter for inhibition of H$_2$ production [16]. The supersaturation of H$_2$ in the liquid enacts a decline in H$_2$ productivity via a metabolic shift to lactate, which may be a mechanism to steer away from exceeding the critical dissolved H$_2$ concentration at which growth is inhibited [16]. To maintain a high productivity the dissolved H$_2$ concentration should be kept...
low through increasing the mass transfer rate in the reactor, e.g. by using an appropriate sparging gas and/or proper reactor design.

Nonetheless, since data on dissolved H₂ concentrations are scarce in the literature and P₁₁₂ is usually the determined parameter, the P₁₁₂ will continue to be used for comparisons in this review.

5.1 Hydrogen tolerance and sulphur reduction

Based on thermodynamics, the heterotrophic archaeon *Pyrococcus furiosus*, with its growth optimum near 100°C [53], should reach a higher H₂ yield at higher P₁₁₂ than that obtained with *C. saccharolyticus* [10]. This organism also has an unusual advantage in that its GAPDH counterpart, glyceraldehyde-3-phosphate: ferredoxin oxidoreductase (GAPOR), generates Fd_red [54], which has an exceptionally low formal potential (E_Fd100°C = -600 mV; [44]). Therefore, based only on thermodynamic constraints (Eq. 1; [41]), *P. furiosus* should be the least H₂-inhibited anaerobe listed in Table 1[41]. Yet, it produces only 2-3 moles H₂/mol hexose, it is highly sensitive to an increase in P₁₁₂ [55], and, when present, it prefers to reduce S⁰ instead of protons. In nature, low-P₁₁₂ environments are usually achievable for H₂ producers living in symbiosis with hydrogenotrophic methanogens [10].

This preference for S⁰ can be explained by examining the physiological properties of its hydrogenases, of which *P. furiosus* possesses three distinct sets. It has two cytosolic NiFe-hydrogenases, which can reduce either protons to H₂ or polysulphide to H₂S, using NADPH as an electron donor. It also has a membrane-bound NiFe-hydrogenase, which can reduce protons to H₂ using Fd_red as an electron donor [51], with the unique capability of generating a proton motive force via proton translocation [56]. Moreover, it has a NiFe-containing hydrogenase complex [51], but its physiological role is partly unknown. As a third set, the organism possesses a ferredoxin:NADP oxidoreductase (or sulphide dehydrogenase) that regulates product formation depending on the P₁₁₂ [51]. In other words, with increasing P₁₁₂, this enzyme becomes more active and decreases H₂ production by transferring electrons from Fd_red to either polysulphide or NADP [51]. NADPH can then be used to reduce pyruvate to alanine, catalyzed by an NADPH-dependent glutamate dehydrogenase and alanine aminotransferase [57].

S⁰ reduction by *Thermoanaerobacter tengcongensis* enhances growth, and in its absence the organism is highly sensitive to H₂ and growth is strongly inhibited at P₁₁₂ above 10 kPa [45]. There are, however, *Thermoanaerobacter* species that are H₂-tolerant, such as the ethanol-adapted *T. thermosulfurigenes* [58]. This adapted strain is able to maintain growth at 1 atm H₂ by redirecting its metabolism to ethanol, but its H₂ yield is not higher than 50% of the Thauer limit [58].

Members of the genus *Thermotoga*, which are isolated from both shallow and deep sea hydrothermal vents rich in S⁰ [59], are normally able to reduce either S⁰ or thiosulphate [60]. The members of this genus display varying degrees of sensitivity to H₂, but H₂ inhibition can be generally relieved by the addition of S⁰, most likely as the result of a “detoxification” reaction [61]. For instance, the *Thermotoga* sp. strains FjSS3.B1 and FjSS. B1 are both highly sensitive to H₂ and the addition of S⁰ can re-establish their growth [61]. Although not generally required for its growth, *Tm. maritima* is able to reduce S⁰, but does not respire it, decreasing its H₂ yield by 40% [62]. In a H₂-pressurized reactor (H₂/CO₂, 80/20%; 300 kPa), however, growth of *Tm. maritima* cells was not observed unless S⁰ was added [61]. Johnson et al [63] demonstrated that *Tm. maritima* was strongly inhibited by H₂, which triggers the cells to enter stationary phase. Consistently, Schröder et al [64], demonstrated that P₁₁₂ levels above 2.3 kPa might be inhibitory to growth of *Tm. maritima*. However, since the organism was cultivated under uncontrolled conditions, other effects, such as the growth-associated drop in pH, might have contributed to growth inhibition. Although *Marinitoga* is closely related to *Thermotoga*, members of this genus can generally tolerate more H₂ than *Thermotoga* species. For instance, *M. hydrogenotolerans* can grow at 1 atm H₂ without any reduction in biomass yield, but then ethanol is formed under these conditions, which drains the electrons required for H₂ production. In addition, in the presence of S⁰, H₂ is not formed [65].

In contrast, members of the genus *Caldericelluloliruptor* are generally unable to reduce S⁰ or thiosulphate [29,62,63]. Moreover, both *C. saccharolyticus* and *C. owensensis* are capable of growing and producing H₂ at P₁₁₂ levels up to 67 and 44 kPa, respectively [16,66]. Similarly, growth of *C. kristjanssonii* was not inhibited at P₁₁₂ levels up to 50 kPa [67].

The information gathered from the literature so far, in addition to our own work, reveals that S⁰-reducing thermophiles are generally more sensitive to H₂ than thermophiles that are unable to use S⁰ as an electron acceptor. This conclusion supports the previously proposed correlation between the ability to reduce S⁰ and the sensitivity to H₂ inhibition [60]. Due to inconsistencies in the experimental design for the different microorganisms studied and the lack of data on dissolved H₂ concentrations, one should be aware that it is difficult to quantify H₂ tolerance. In conclusion, based on this knowledge, it can be suggested to select for non-S⁰-reducing H₂-producing
candidates and future attempts to isolate promising H2 producers should be targeting S0-poor environments.

5.2 H2 removal from the liquid

To obtain higher H2 yields and productivities, high dissolved H2 concentrations should be avoided [51]. Therefore, methods to increase the mass transfer rate of H2 from the liquid to the gas phase should be employed. Several studies have been dedicated to develop different ways for efficient removal of H2 from the liquid phase (Table 2) [31].

Currently, N2 is commonly used as a sparging gas for lab-scale H2 production. However, applying this method at an industrial scale is not cost-effective since N2 is inert, and thus difficult to remove from the effluent gas stream [11]. Instead, CO2 could be an appropriate alternative since it can be more readily separated from H2 and, as a convenience, it is a product of the fermentation process itself [11]. Nevertheless, being chemically non inert, CO2 can influence the metabolism of some microorganisms as well as the composition of the culture medium, and hence may not be viewed as an adequate universal sparging gas. In C. saccharolyticus cultures, CO2 was already shown to negatively influence growth and H2 production rate [31]. In that study, higher partial pressures of CO2 (Pco2) were found to increase the concentration of dissolved CO2, which is hydrolyzed to bicarbonate and protons. This requires the addition of larger amounts of a caustic agent to compensate for the decrease in pH, contributing to an increased osmotic pressure, which inhibits growth in C. saccharolyticus, and a higher environmental burden [31]. Methane is another interesting alternative sparging gas that was previously shown to improve the H2 yield of a mesophilic consortium by 88% [5]. However, the effect of this gas on H2 production by C. saccharolyticus has not been investigated yet.

Other strategies, such as applying reduced pressure to remove H2 and CO2 from the headspace, have been shown to efficiently increase H2 yields in Enterobacter cloacae at lab scale [68]. However, on an industrial scale, this strategy might not be economically feasible due to high energy cost and increased contamination risk. Instead, two other alternative approaches may be pursued: i) improvement of reactor design to decrease the levels of dissolved H2 and CO2, and ii) metabolic engineering of the organism to enable it to either produce higher H2 yields at high P1H2 or to withstand higher osmotic pressures.

6. Sugar transport and energy conservation

Based on its genome sequence and transcriptome analysis, C. saccharolyticus appears to rely primarily on ABC transporters for sugar uptake [23,26]. Bacterial and archaeal ABC transporters consist of five subunits, i.e., a sugar-binding protein, two transmembrane domains and two nucleotide-binding domains [69]. A common characteristic of the sugar binding protein is its ability to capture the sugar outside the cell via a cap-lock mechanism, which determines the high affinity of the transporter for the substrate. Despite the large number of ABC transporters in C. saccharolyticus [26], none of them have been characterized so far. However, based on transcriptional analysis, it has been concluded that, in C. saccharolyticus, the same transporter might translocate different sugars with varying affinities [23,26].

From a physiological perspective, ABC transporters have the disadvantage of relatively high energy demand, compared with the phosphoenol pyruvate (PEP)-dependent phosphotransferase system (PTS) (Table 3) [70]. In the latter system, one mole of the high-energy compound PEP is required for both sugar transport and subsequent phosphorylation [70], whereas two moles of ATP are required for the same steps using an ABC transporter. The fuelling of these energy demanding, high-affinity transport systems in a strict anaerobe such as C. saccharolyticus requires an efficient energy-conserving metabolism. It can be hypothesized that one of these energy-conserving strategies is the oxidation of glucose to acetate and CO2, which results in two extra moles of ATP per mole of hexose on top of the usual ATP yield of 2 mol/mol hexose in the EMP pathway. Thus, with the conversion of glucose to two moles of

| Sparging gas         | Separation | Benefits                      | Drawbacks                                      | Reference |
|----------------------|------------|-------------------------------|------------------------------------------------|-----------|
| N2                   | Difficult  | Not inhibitory                | Difficult to separate from H2, Expensive       | [10,28]   |
| CO2                  | Easy       | Easily separated, Produced in the process | Increases osmolarity, Inhibits H2 productivity | [10,28]   |
| CH4                  | Easy       | Easily separated, Produced in a hybrid bioreactor/biogas process | No results available for C. saccharolyticus | [5]       |
| Under-pressure       | Easy       | Not inhibitory, No dilution of the gas, Cheap separation | Expensive at large scale, Risk of contamination | [68]      |
| No sparging          | Easy       | No dilution of the gas, Cheap separation | Inhibits H2 productivity, Inhibits H2 yield | [16]      |
acetate, maximally four moles of ATP can be acquired, with the net yield being only three moles if an ABC transporter is involved (Table 3). This means that with acetate as the end product, the desired H₂ production becomes advantageous for the microorganism since it serves as an electron sink in the re-oxidation of the reduced cofactors Fdₚₚ and NADH generated in the breakdown of sugars.

To investigate the presented hypothesis further, the genomes of known H₂-yielding anaerobes were analyzed to evaluate whether they possess PTS- or ABC-type of sugar transport. Indeed, there is a clear correlation between high H₂ yields and the use of ABC transporters (Tables 1 & 4). Organisms that were able to produce high H₂ yields generally possess mainly ABC-type transporters, whereas low H₂-yielding bacteria utilize more PTSs for transport. This correlation between H₂ yields and possession of ABC transporters is more pronounced in thermophiles. For example, the extreme thermophile *Thermoaerobacter pseudoethanolicus*, which has a PTS for glucose uptake, is unable to produce more than 2 moles H₂/mol hexose (101), and mainly forms ethanol as a metabolic end product instead of acetate.

### Table 3 Theoretical ATP yield (YATP/S) (mol/mol) of anaerobic glucose oxidation with different transport systems and fermentation products

| Transport  | Fermentation product |
|------------|----------------------|
|            | acetate | lactate | ethanol | butyrate | butanol | propionate |
| ABC        | 3       | 1       | 1       | 2        | 1       | 1          |
| PTS        | 4       | 2       | 2       | 3        | 2       | 2          |

### Table 4 The type of transport system with respect to growth temperature

| Organism                  | Opt T °C | Transporters | Reference |
|---------------------------|----------|--------------|-----------|
| *Enterobacter aerogenes*  | 37       | PTS          | [107]     |
| *Enterobacter cloacae*    | 37       | PTS          | [107]     |
| *Clostridium butyricum*   | 30       | PTS          | This study |
| *Clostridium acetobutylicum* | 34     | PTS          | [108]     |
| *Clostridium thermocellum* | 60      | ABC          | [109]     |
| *C. saccharolyticus*      | 70       | ABC          | [26]      |
| *Thermotoga neapolitana*  | 77       | ABC          | [110]     |
| *Thermoaerobacter tengcongensis* | 75 | ABC | This study |
| *Thermoaerobacter pseudoethanolicus* | 70 | PTS | This study |
| *Pyrococcus furiosus*     | 90       | ABC          | This study |

### 6.1 PPI as an additional energy carrier

Another strategy to conserve energy employed by *C. saccharolyticus* is to utilize inorganic PPI as an additional energy carrier [71]. This strategy is widely applied by anaerobes and other organisms that can acquire only low energy outputs in their metabolism to obtain sufficient ATP [72], of which *Tm. maritima* [73] and various *Clostridium* species [72,73] are a few examples that have been studied to some extent.

The primary sources of PPI are poly-nucleic acid biosynthesis from (deoxy)nucleotide triphosphates and activation of amino acids and fatty acids for protein and lipid synthesis, respectively, which are reactions that operate close to equilibrium. Therefore, subsequent PPI hydrolysis is required for driving these reactions forward [74]. Moreover, studies on *E. coli* have shown that maintaining low PPI concentrations through effective PPI hydrolysis is required for rapid growth [75]. In *E. coli*, as in a majority of other organisms, PPI hydrolysis is catalyzed by a cytosolic inorganic pyrophosphatase (PPase), with the dissociation of the free-energy change of the bound PPI ($\Delta G^0 = -21.8$ kJ/mol; [76]) as heat.

*C. saccharolyticus* lacks a cytosolic PPase. Instead, the energy of PPI hydrolysis can be conserved, for instance through: (i) an active membrane-bound PPase employing the free-energy change of the phosphate bond hydrolysis to establish a proton motive force that could be utilized for transport of nutrients or for ATP synthesis; (ii) a PPI-dependent phosphofructokinase (PPI-PFK) in addition to the ATP-dependent one (ATP-PFK); and (iii) a PPI-dependent pyruvate phosphate dikinase (PPdK) in addition to the ADP-dependent pyruvate kinase (PYK; Figure 2) [71].

Interestingly, there are indications that the PPI-level follows the growth dynamics of *C. saccharolyticus*. When the cells were growing exponentially at the maximum growth rate, the PPI levels were relatively high (4 ± 2 mM), and when the growth rate decreased, the PPI levels declined concordantly. The dynamics in PPI levels were shown to be associated with the low cytosolic PPase activity in *C. saccharolyticus* [71]. Consistently, the PPI levels followed the same growth-dependent trend in *Moorella thermoacetica* and *Cl. pasteurianum*, both of which also lack a cytosolic PPase [76]. In contrast, in *E. coli* possessing elevated cytosolic PPase activity (0.54 U/mg cdw), the PPI levels were low (0.3 mM) and did not fluctuate during growth [76].

Unlike PPI, the ATP levels in *C. saccharolyticus* were very low (0.43 ± 0.07 mM) during the exponential growth phase, but increased 2 folds at the beginning of the stationary phase [71]. Interestingly, the ADP concentration was higher than that of ATP during both phases [71], which is generally considered to be a sign of starvation. Similar low ATP/ADP ratios were observed in...
Figure 2 Proposed catabolic network in C. saccharolyticus. Glucose is transported over the membrane by an ABC transport system (1) [23,26] and oxidized to pyruvate in the EMP pathway [13,26], with the possibility that PPI can potentially replace ATP, i.e., PPI-PFK (2) instead of ATP-PFK (3), or ADP, i.e., PPDK (5) instead of PYK (6) [71]. NADH generated through the GAPDH-catalyzed reaction (4) and Fd_red generated from the PFOR-catalyzed reaction (7) can donate electrons for the generation of molecular H₂ catalyzed by cytosolic NADH-dependent Fe-only hydrogenase and membrane-bound Fd-dependent NiFe-hydrogenase, respectively [26]. Acetate is formed by the consecutive actions of phosphotransacetylase (PTA; 8) and acetate kinase (AK; 9), with the generation of ATP. Alternative routes for NADH reoxidation are the formation of lactate (LDH; 11), ethanol (ADH; 10) or succinate (12) [16]. The solid lines represent metabolic routes, whereas the dashed lines represent metabolite activation (+) or inhibition (−) of enzyme activities. PPI generated from anabolic reactions [71] is a strong inhibitor of both PYK and LDH activities [47,71].
Cl. acetobutylicum [77]. Nevertheless, C. saccharolyticus grew exponentially, suggesting that it relies on other energy carriers, such as PPI, that may contribute to its energy charge in addition to ATP [71].

The use of PPI as an additional energy carrier also introduces greater metabolic flexibility. In addition, it creates a stronger link between anabolism and catabolism, i.e., the net ATP generated in catabolic pathways is used as an energy input in biosynthetic pathways, which generates PPI that can participate in driving catabolism and generating ATP (Figure 3). In this cycle, both ATP and PPI may be key regulators of catabolic and anabolic fluxes.

6.2 Enzyme regulation
Regulation of the classical glycolysis and the flux distribution at the pyruvate node is a very complex and highly tuned process, since it is the principal provider of energy carriers and redox equivalents that need to be precisely balanced under all growth conditions. Regulation takes place at metabolite level as well as at enzyme and gene expression levels. Enzyme activities normally are regulated via direct mass action law (concentrations of substrates or effectors) or via allosteric inhibition or activation [78]. To maintain high growth rate conditions, cells of C. saccharolyticus should obtain optimal energy gain from the substrate to fuel both anabolism and sugar transport, and thus lactate dehydrogenase (LDH) and alcohol dehydrogenase (ADH) should be kept inactive (Figure 2). Indeed, a recent batch-fermentation study has shown that during exponential growth, H2, CO2 and acetate are the only fermentation products in C. saccharolyticus [47]. In this growth phase, neither ethanol nor lactate was formed, although both LDH and ADH activities were detected in crude cell extracts at high levels. However, as the growth rate decreased during transition to the stationary phase, the metabolism was partly directed towards lactate formation. Similar growth phase-related metabolic shifts have been observed in Cl. cellulolyticum [48]. Studying the in vitro kinetics of LDH activity in C. saccharolyticus confirmed the observations of the batch experiments. It revealed that LDH activity in C. saccharolyticus is strongly regulated by the levels of the energy carriers PPI and ATP,

![Figure 3 Schematic representation of the suggested circulation of energy carriers in catabolism and anabolism in C. saccharolyticus during exponential growth (A), and non-growth (B), and its influence on metabolite formation.](image-url)
in addition to the NADH/NAD ratio (Figure 2) [47]. When the cells are growing at the maximum specific growth rate, PPI levels are high and ATP levels are low, keeping LDH inactive and its affinity for NADH low. It further assures that the catabolic flux is directed to acetate and H2 (Figure 2). However, as soon as the anabolic activity declines, the PPI/ATP ratio drops by an order of magnitude [71], which results in an increase in LDH activity as well as its affinity for NADH and hence lactate starts being formed [47].

Due to the strong control of C. saccharolyticus LDH activity at enzyme level, lactate is formed only in special cases, i.e., at elevated NADH/NAD ratios and at higher ATP levels. High NADH/NAD ratios also lead to inhibition of GAPDH activity and consequently the glycolytic flux [16,47]. Since LDH activity secures the continuation of the glycolytic flux, through diminishing the NADH/NAD ratio, the enzyme can be considered essential for balanced growth in C. saccharolyticus. It thus might be questioned whether deleting the ldh gene would improve H2 yields during sugar fermentation. It might be a better approach to control lactate production through maintaining process conditions that promote exponential growth to keep high cellular PPI levels [47]. It is worth noting that deletion of the ldh gene in E. coli only slightly increased H2 yield and productivities [79].

Ethanol is another reduced end product that is formed during growth of C. saccharolyticus, albeit in very low quantities (Figure 2; [13,42]). Based on sequence similarities to the characterized ADH in T. ethanolicus [80], ADH in C. saccharolyticus possesses a NADPH-binding domain instead of a NADH-binding domain (Figure 2) [26]. This observation is supported by a preliminary kinetic study showing that the affinity of the ADH of C. saccharolyticus for NADPH is higher than for NADH [16], indicating that ethanol and H2 formation do not compete for the same reduced cofactor.

13C-NMR analysis of intact cells has also revealed that C. saccharolyticus is capable of reducing either pyruvate or PEP to succinate (Figure 2) in the reductive branch of its incomplete TCA cycle. However, the succinate yield is significantly lower than that of acetate or lactate under regular growth conditions and might only operate under combined elevated P142 and PCO2 levels [31].

Understanding the regulation of the key glycolytic enzyme GAPDH in C. saccharolyticus may also contribute to a better understanding of the organism’s ability to tolerate elevated P142. In general, GAPDH is strongly regulated by the NADH/NAD ratio [81], and GAPDH in C. saccharolyticus proved to be no exception [16]. However, the enzyme in C. saccharolyticus is more resistant to increased NADH levels than GAPDH in most other related bacteria. The NADH concentration required for 50% inhibition of the enzyme in C. saccharolyticus was 0.03 mM, as compared to 0.01 mM in T. thermohydrosulfurificus (formerly known as Cl. thermohydrosulfuricum) [58] and Cl. acetobutylicum [77]. This observation would in theory have a significant impact on the organism’s tolerance to H2, i.e., if GAPDH was less inhibited by NADH, higher NADH/NAD ratios could be tolerated in the cell without affecting the glycolytic flux. As discussed above, this would also render the hydrogenase reaction more energetically favourable (Eq. 1) [10]. Consistently, the ethanol-adapted T. thermohydrosulfurificus strain possesses a GAPDH that tolerates approximately twice the amount of NADH and is more tolerant to H2 than the wild-type strain [58]. Likewise, the NADH/NAD ratio in Cl. cellulolyticum is significantly higher than in C. saccharolyticus, which is possibly due to a more relaxed inhibition of its GAPDH by NADH (50% inhibition at 0.1 mM NADH) [42,43].

7. Challenges for optimizing hydrogen production by C. saccharolyticus

C. saccharolyticus has several desirable characteristics needed for a H2 cell factory, such as its high H2 yields on a wide spectrum of carbon sources, its relatively high tolerance to elevated P142, and its inability to reduce S0, which therefore does not interfere with its H2 production capabilities. However, the organism also lacks several properties, which should be considered before it can be used industrially.

7.1 Sensitivity to high osmolalities

Primarily, the sensitivity of C. saccharolyticus to increased osmotic pressures is a major drawback that should be addressed. The critical concentration of salts and solutes at which the growth of the organism ceases completely was estimated to be 400-425 mM [30]. Osmolalities above 0.218 osm/kg H2O were found to induce cell lysis in C. saccharolyticus, as indicated by increased protein and DNA concentrations in the culture supernatant [31]. This low osmotolerance imposes limitations on sparging the culture with CO2 and on the maximum sugar concentration in the medium [31], with the latter obviously having a negative effect on H2 productivity [30]. On the other hand, the hyperthermophiles Tm. neutolitana and P. furiosus can grow optimally in the presence of 0.46 M NaCl and 0.5 M NaCl, respectively [82,83]. These salt concentrations correspond to an additional 0.92 and 1 osm/kg H2O, respectively, in the culture media, demonstrating that these two hyperthermophiles are better protected against osmotic stress than C. saccharolyticus. This higher osmotic tolerance is due to their ability to produce compatible solutes, or osmoprotectants, such as β-mannosylglycerate, di-myo-insitol phosphate and glutamate [82,83]. It remains to be investigated whether C. saccharolyticus is also able to
efficiently synthesize similar solutes when exposed to osmotic stress. The low salt concentration in its natural habitat [28] may have resulted in the loss of any capability to survive in an unfavourable osmotic environment [82]. However, the presence of biosynthetic genes for some osmoprotectants, such as proline and glutamate, could open the way for directed evolution to render C. saccharolyticus more osmotolerant [31].

7.2 Low productivities
The low cell density of cultures of C. saccharolyticus and other thermophiles, is another major drawback for industrial-scale application. It results in a low volumetric H₂ productivity [83], unless dialysis or cell-recycling methods are used to increase the cell mass [84]. Especial dialysis was found to increase the cell density of Pyrococcus furiosus by an order of magnitude, indicating that the presence of metabolic products prevents the increase in cell densities. In continuous cultures of C. saccharolyticus, the volumetric productivity could be enhanced by increasing the dilution rates, but this comes at the expense of the H₂ yield [13,85]. The decrease in yield could be related to increased dissolved H₂ concentrations resulting from higher H₂ productivities[16]. This can inhibit the hydrogenase activity and ultimately results in a metabolic shift to lactate. The same trade-off between yield and productivity was also observed with other microorganisms. For instance, E. coli produces much lower H₂ yields but its volumetric productivity is almost 100-fold higher than in C. saccharolyticus [13,86]. The cell density is about 200 times higher for E. coli [85] than for C. saccharolyticus [14], which contributes considerably to the higher volumetric productivity. The specific H₂ productivity of E. coli, however, is low, which could be an undesirable consequence of the higher cell number, i.e., it enhances the H₂ mass-transfer resistance [85].

7.3 Biofilm formation and reactor design
Since C. saccharolyticus exhibits a relatively high specific H₂ productivity, its volumetric production rate can potentially be enhanced through increasing its cell density [83] by, for example, inducing biofilm formation. The reactor configuration is an important factor in enhancing biofilm formation. So far, most research on C. saccharolyticus has been carried out in continuously stirred tank reactors (CSTR) [2,13,14,85], which do not allow for efficient biomass retention. Alternative reactors that enhance biofilm formation include trickling-bed bioreactors [11] and fluidized-bed systems with granules [86]. Trickling-bed bioreactors promote biofilm growth by means of packing material with a high surface area within the trickling bed. The liquid passes continuously through the filter, such that it guarantees a good liquid-gas exchange. In the fluidized-bed system, the biofilm is also formed as granules. This reactor system increases the turbulence, thus providing an enhanced mass transfer, which potentially also could decrease the dissolved H₂ concentration, and hence, improve H₂ yields. It should be noted that the biofilm mode of growth also offers the surface-attached and matrix-encased bacteria a means of protection against several stressful environmental conditions [87], which could benefit an organism like C. saccharolyticus in tolerating higher osmotic pressures.

Recently, another Caldicellulosiruptor sp., viz. Caldicellulosiruptor owensensis, was shown capable of biofilm formation in both trickling-bed and fluidized-bed reactors at 70°C [88]. Moreover, the organism retained its ability to grow and produce H₂ in the fluidized-bed system even in complete absence of gas sparging.

8. Conclusions
This review article has focused on C. saccharolyticus as having several desirable characteristics for efficient H₂ production. This is not only due to the high growth temperature of this organism, but there are strong indications that its metabolic constitution is adapted to efficiently extract energy from carbohydrates, thereby allowing the production of H₂ at yields close to the Thauer limit [14]. The energy metabolism of C. saccharolyticus, based on maximum ATP production per sugar unit and the use of PPI as an alternative energy carrier, might be correlated with the vast number of ABC transporters it employs. The efficiency of the energy metabolism is further illustrated by the modulation of LDH activity in C. saccharolyticus through the levels of the energy carriers ATP and PPI [47]; an active LDH will significantly lower the ATP production flux and thus the growth and H₂ production rates, in addition to draining the electrons required for H₂ production.

C. saccharolyticus can simultaneously co-utilize a wide variety of different sugars [24] and produces efficient extracellular hydrolytic enzymes, including cellulases [21]. It has a relatively high tolerance to partial hydrogen pressures that might be related to its inability to reduce S⁰.

The set of capabilities of C. saccharolyticus described above is the foundation to exploit this organism in a commercial process using various raw materials [89]. However, for obvious economical reasons the performance of C. saccharolyticus needs to be further improved, i.e., both high H₂ yields and productivities should be obtained at relatively high P₅₁₂ [6].

Several metabolic engineering strategies can be foreseen to improve H₂ yields of C. saccharolyticus beyond 3.5 mol/mol hexose, without the need for gas sparging. First, the NADH-dependent GAPDH in C. saccharolyticus could be replaced with an Fd-dependent GAPOR,
similar to the one in *P. furiosus* [54] making the corresponding H$_2$-generation reaction spontaneous even at elevated $P_{H2}$ [9,36]. Second, introducing the genes for the oxidative part of the PPP supplemented with an NADPH-dependent hydrogenase could allow the oxidation of one mole of hexose to one mole of acetate and four moles of CO$_2$ [14,90]. This would in theory yield 8 mol H$_2$/mol hexose, but would not help overcome the thermodynamic barrier.

*C. saccharolyticus* exhibits a considerably high specific H$_2$ production rate compared to other H$_2$ producers [83], but poor volumetric productivities are obtained since its maximum cell densities is in the range of 10$^8$-10$^9$ cells/mL [15]. Strategies to increase the cell density by at least an order of magnitude would be through inducing biofilm formation and/or increasing the substrate concentration. The latter requires strains with at least 5-fold increased osmotolerance, which might be obtained through evolutionary engineering. It is noteworthy that *C. saccharolyticus* possesses all the essential genes necessary for biosynthesis of compatible solutes, such as glutamate and proline [31].

As a technical approach, oversaturation of H$_2$ in the liquid phase should be avoided to stabilize the fermentation process. This can be solved by smart reactor design and process configuration. Different membrane reactors have been developed, which are capable of reducing dissolved H$_2$ concentrations and have a positive effect on H$_2$ yields [91-94]. Yet, it remains to be investigated how *C. saccharolyticus* would perform in these reactor systems.

All in all, it is vital to understand the physiology of H$_2$ production, which can be facilitated through applying genetic engineering and state-of-the-art ‘omics’ tools. However, so far, genetic protocols have been developed only for a few thermophilic hydrogen producers [95-98], but not yet for *C. saccharolyticus*. Omics-based technologies are increasingly applied for studying thermophilic H$_2$ producers [25,99,100], including *C. saccharolyticus* [24,25]. Together with genome-wide metabolic models, now being developed for some thermophiles [101,102], the systems biology approach can eventually be applied to accelerate our understanding of the physiology of *C. saccharolyticus* and other thermophiles to increase their potential for process applications.

The insights in the physiology of *C. saccharolyticus*, as covered in this review, should contribute to better strategies for the isolation and/or engineering of future superior H$_2$ producers. Creating a superior H$_2$ cell factory is a quintessential step towards a sustainable biological H$_2$ production.

### 9. Competing interests

The authors declare that they have no competing interests.

### 10. Authors’ contributions

KW conceived of the study, designed and drafted the manuscript. AAZ contributed to the design and content of the review article and helped to draft the manuscript. EWJN helped to draft the manuscript. AAZ and EWJN critically revised and commented the manuscript. All authors read and approved the final manuscript.

### 11. Acknowledgements

AAZ acknowledges support from the Ministry of Higher Education in Egypt. The research on biohydrogen production in our lab is financially supported by the Commission of European Communities, Sixth Framework Program, Priority 6, Sustainable Energy Systems (019825 HYVOLUTION) and the Swedish Energy Agency (Energymyndigheten, 31090-1).

Received: 31 August 2010 Accepted: 22 November 2010 Published: 22 November 2010

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doi:10.1186/1475-2859-9-89

Cite this article as: Willquist et al.: Physiological characteristics of the extreme thermophile \textit{Caldicellulosiruptor saccharolyticus}: an efficient hydrogen cell factory. \textit{Microbial Cell Factories} 2010, 9:89.