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Plasma technology – a novel solution for CO₂ conversion?

Plasma technology could provide a viable solution for the conversion of CO_2 into value-added chemicals while the fate of the world hangs in the balance. The future of the Earth rests in our hands.

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Plasma technology – a novel solution for CO₂ conversion?

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CO₂ conversion into value-added chemicals and fuels is considered as one of the great challenges of the 21st century. Due to the limitations of the traditional thermal approaches, several novel technologies are being developed. One promising approach in this field, which has received little attention to date, is plasma technology. Its advantages include mild operating conditions, easy upscaling, and gas activation by energetic electrons instead of heat. This allows thermodynamically difficult reactions, such as CO2 splitting and the dry reformation of methane, to occur with reasonable energy cost. In this review, after exploring the traditional thermal approaches, we have provided a brief overview of the fierce competition between various novel approaches in a quest to find the most effective and efficient CO2 conversion technology. This is needed to critically assess whether plasma technology can be successful in an already crowded arena. The following questions need to be answered in this regard: are there key advantages to using plasma technology over other novel approaches, and if so, what is the flip side to the use of this technology? Can plasma technology be successful on its own, or can synergies be achieved by combining it with other technologies? To answer these specific questions and to evaluate the potentials and limitations of plasma technology in general, this review presents the current state-of-the-art and a critical assessment of plasma-based CO2 conversion, as well as the future challenges for its practical implementation.

1. Introduction

Environmental and energy applications of low temperature plasmas are gaining increasing interest worldwide. The central research question is whether plasma-based solutions can yield

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Ramses Snoeckx

Ramses Snoeckx, born in 1988, obtained his Master's degree in both Environmental Science and Chemistry. Combining these backgrounds, he successfully obtained his PhD in Chemistry (2017) for his research on the plasma-based conversion of greenhouse gases into value-added chemicals and fuels. He actively collaborated with several (inter)national research institutions, for example during research visits at the Instituto Superior Técnico (Portugal), the

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Annemie Bogaerts, born in 1971, obtained her PhD in Chemistry in 1996, from the University of Antwerp in Belgium. She became professor of physical chemistry in 2003, at this university, and has now been full professor since 2012. She is head of the interdisciplinary research group PLASMANT. The research activities of her group include the modelling of plasma chemistry, plasma reactor design and plasma-surface interactions, as well as carrying out plasma

experiments for various applications, including environmental and medical applications (mainly cancer treatment), as well as nanotechnology and analytical chemistry. In recent years, special attention has also been given to CO2 conversion by plasma and plasma catalysis.

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a valuable alternative to existing thermal processes and whether they can compete with other novel gas conversion technologies. Nowadays, the conversion of CO₂ into chemicals and fuels is a hot topic. The worldwide transition to renewable energy gives plasma processes a clean electricity source, and due to their high operation flexibility, plasmas are very suitable for storing this intermittent renewable energy in a chemical form, i.e. as fuels and chemicals.

1.1. CO₂ mitigation and valorisation

Throughout history, the use of natural resources has played a major role in the rapid development of the human race. Among these resources, fossil fuels in particular have contributed to a fast and unprecedented development in human society. Still, this comes with a great cost, since burning fossil fuels leads to the emission of large amounts of the greenhouse gas CO₂. Because these anthropogenic CO₂ emissions outpace the natural carbon cycle, atmospheric CO2 concentrations have been increasing from 280 ppm since the beginning of the industrial revolution to 400 ppm in 2014.1 With high certainty, it can be said that it is this increase that has led to the current adverse global environmental climate changes, which have a growing detrimental effect on our climate and environment, and that represent a severe threat to our current society and future generations in general.^{2–5}

Therefore, the conversion of this main greenhouse gas into value-added chemicals and liquid fuels is considered as one of the main challenges for the 21st century. 1,2 The aim is not only to tackle climate change, but also to provide an answer to our dependence on fossil fuels. As stated by Goeppert et al.,6 "Whether humankind uses up most of the fossil fuel resources (combined with carbon capture) or uses increasingly alternative energies, the need for transportation fuels and materials that we currently obtain from petroleum and natural gas will remain. With increasing population, products based on carbon from plastics to medicines will also be required in increasing quantities. In order to fulfil the demand for carbon-based products, CO2 will have to be recycled in an anthropogenic version of nature's own carbon cycle. Carbon capture and recycling (CCR) will capture CO2 from any source, and eventually mainly from the atmosphere, and recycle it to new materials and fuels using any alternative energy source.'

Utilization of this waste and converting it into a new feedstock not only complies with the framework of sustainable and green chemistry^{7,8} but also fits within the 'cradle-to-cradle' concept. 9 By generating useful products out of CO₂ we create the possibility to effectively close the carbon loop. This has already resulted in a booming interest in technologies that can convert CO₂ into value-added products, 10-12 since they can effectively convert waste into new feedstocks following the cradle-to-cradle principle.⁹ Besides chemical conversion, which is the principle focus of this review, CO₂ also has other applications in the field of carbon dioxide capture, storage and utilization, such as its fixation and technological utilization, for these and other processes, as well as in carbon capture, for which we refer to several existing reviews in the literature. 12-19

Besides the traditional thermal CO₂ conversion, several alternative technologies are being investigated, such as electrochemical, solar thermochemical, photochemical and biochemical pathways, either with or without catalysts, as well as all their possible combinations. Their advantages and disadvantages are briefly discussed herein, as they form the context for a novel approach considered to have great potential in recent years, which is the approach based on (non-thermal) plasma. 20,21 Several options are being investigated in this regard, including both pure CO2 splitting into CO and O2, as well as the reaction with other gases, like CH₄ (dry reforming of methane), H₂ (hydrogenation of CO₂) or H₂O (artificial photosynthesis), aiming for the production of syngas and valuable oxygenates, such as methanol, formaldehyde and formic acid.

1.2. Solar fuels

The important difference between the traditional thermal conversion approaches and the emerging technologies is their independence of burning fossil fuels to provide the necessary thermal heat to drive reactions. More precisely, biochemical pathways rely on the natural photosynthesis process to convert light (either natural or artificial) into biomass, which can then be harvested and further processed. Other emerging technologies, such as the solar thermochemical and photochemical processes rely on direct solar energy. The average solar flux striking the earth's surface is 175 W m⁻², which represents more energy striking the earth's surface in two hours, i.e. 640 EJ, than the worldwide energy consumption from all sources combined in 2008, i.e. 514 EJ.²² The solar thermochemical process makes use of concentrated solar heat, while the photochemical process relies on the energy of photons. Ultimately though, the electrochemical and plasmachemical processes rely on electricity. Fig. 1 gives an

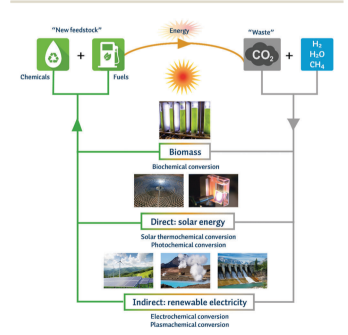


Fig. 1 Overview of the different novel technologies and their principal use of renewable energy for the conversion of CO₂ in a carbon neutral cycle.

overview of the different novel technologies and their principal use of renewable energy for the conversion of CO₂ in a carbon neutral cycle.

The reliance of electrochemical and plasmachemical conversions on electricity at first seems to limit their use as a greenhouse gas mitigation technology, since currently, producing electricity generally results in CO2 emissions. This idea though could not be further from the truth due to the worldwide transition to renewable energy sources, such as solar and wind energy. In 2014, the estimated renewable energy share of the global final energy consumption was already 19.2%, while by the end of 2015, the estimated renewable energy share of global electricity production was 23.7%.²³ It is even at such a stage that the large-scale adoption of these renewable energy sources poses a challenge for the efficient storage and easy transport of the electricity produced, i.e. not only regarding the need for peak shaving, but more importantly the need for technologies to follow the irregular and at times intermittent supply of renewable electricity in a flexible way. While storage in batteries is possible, it is less efficient than chemical storage in fuels.²⁴ Such fuels, often referred to as carbon neutral fuels or solar fuels, offer a much higher gravimetric and volumetric energy storage capacity, have much higher energy densities than electrical storage techniques and they match the existing worldwide liquid fuel infrastructure. 12,24

In the first instance, the reactions in which CO_2 is involved can be divided into two categories: the production of chemicals and the production of fuels. The latter is considered as the most suitable target for the conversion of large volumes of CO_2 since its market size is 12–14 times larger than the former. One of the most interesting compounds is methanol, which is positioned exactly in the middle of these two categories, being at the same time a raw chemical and a fuel, used in both combustion engines and fuel cells. ²⁵

To achieve the transformation of CO2 into value-added chemicals or fuels, the reactions that are of greatest interest involve the conversion of CO2 with a co-reactant that acts as a hydrogen source (like CH₄, H₂ or H₂O). Due to the existing infrastructure, liquid products are preferable to gases, for most applications at least. Two approaches can be considered to achieve this: the indirect oxidative pathway and the direct oxidative pathway. The main product of the former is syngas, a mixture of H₂ and CO, which can be converted to almost any commercial bulk chemical or fuel in a second - albeit very energy intensively - step through methanol and/or Fischer-Tropsch synthesis. 10 In this case, it is of great importance to have a high degree of control over the H₂/CO ratio to be able to steer the synthesis towards the desired products.²⁶ The direct oxidative pathway, on the other hand, tries to eliminate the energyintensive middle man by converting the reactants immediately into hydrocarbons, short-chain olefins (e.g. ethylene and propylene) and oxygenated products (e.g. methanol, formaldehyde, dimethyl ether and formic acid).

Liquid products are more attractive over gaseous hydrogen, since – while in theory a 'Hydrogen Economy'²⁷ would be very attractive – the latter has a number of serious drawbacks due to its physico-chemical properties.²⁸ Furthermore, the infrastructure needed to safely transport, store and dispense hydrogen would be

very expensive to roll out, while liquid chemicals match the already existing worldwide fuel infrastructure. 12,24 Hence, especially in the transportation sector, a transition from liquid-fossil-fuel-derived products (e.g. gasoline, diesel fuel, kerosene) to a renewable and sustainable liquid fuel is highly desirable. Again, methanol is one of the most interesting possible candidates to fulfil these requirements. It is the simplest liquid chemical containing only one carbon. Although to date it is almost exclusively produced from natural gas (and shale gas) for economic reasons, it can easily be obtained from several (future) carbon sources, incl. CO₂, biomass, biogas and landfill gas. Therefore, it has been proposed as a key solar fuel for the above-mentioned anthropogenic carbon cycle under the framework of a 'Methanol Economy'.

To be economically competitive with the existing structures, the efficient production of these solar fuels is critical during the quest to find an effective CO₂ conversion technology with the potential to be commercialized on a large scale.

2. Traditional thermal CO₂ conversion approaches

We first briefly discuss the existing traditional (mainly thermocatalytic) approaches used on an industrial scale. As such, this section will act as a comparison for the novel technologies under development. This section is subdivided into: (1) pure CO₂ splitting and (2) CO₂ conversion in combination with a co-reactant, *i.e.* CH₄, H₂ or H₂O.

2.1. Pure CO₂ splitting

Thermal CO₂ splitting has not been very effective to date. This is not surprising from a thermodynamic point of view; the carbon–oxygen bonds are relatively strong (783 kJ mol⁻¹)²⁹ and the Gibbs free energy of formation $(\Delta G^{\circ} = -394 \text{ kJ mol}^{-1})^{24}$ clearly shows that CO₂ is a highly stable molecule, requiring a substantial energy input, optimized reaction conditions and active catalysts for any chemical conversion to take place. Neither the entropy $(T\Delta S^{\circ})$ nor the enthalpy (ΔH°) term seem favourable for its conversion.² The overall reaction is written as:

$$CO_2(g) \to CO(g) + 1/2O_2(g)$$
 $\Delta H^{\circ} = +283 \text{ kJ mol}^{-1}$ (R1)

Of course, the high value of ΔH° does not mean that its conversion is not feasible. Indeed, strongly endothermic chemical reactions can be found in a large number of industrial processes used worldwide, a classic example being the steam reforming of methane (SMR):^{24,31}

$$CH_4(g) + H_2O(g) \rightarrow CO(g) + 3H_2(g) \quad \Delta H^\circ = +206 \text{ kJ mol}^{-1} \quad (R2)$$

This highly endothermic reaction has found worldwide use. In the fertilizer industry, the $\rm H_2$ is used for the production of ammonia, while in the gas industry, this reaction is responsible for 95% of the worldwide $\rm H_2$ production. This shows that there is no reason to dismiss $\rm CO_2$ splitting just because it is highly endothermic. Hence, a fair amount of research towards this reaction has already been conducted, of which an overview can be found in the work of Rayne.²⁹

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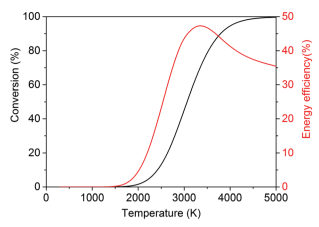


Fig. 2 Calculated theoretical thermal conversion (left axis) and corresponding energy efficiency (right axis) as a function of temperature for the pure splitting of CO₂ into CO and O₂.

It is, however, clear that, without actively removing one of the products (i.e. CO or O_2), the equilibrium of this reaction lies strongly to the left. Thus, thermal CO₂ splitting is thermodynamically and energetically only favourable at very high temperatures, as can be seen in Fig. 2. At 2000 K for instance, the reaction is not very efficient: we can easily estimate that ca. 92 kJ mol⁻¹ would be needed to heat 1 mole of CO₂ from 300 to 2000 K. Furthermore, the reaction enthalpy is equal to 245 kJ mol⁻¹ at 2000 K. Based on a conversion of 1.5% at this temperature, the energy cost for the total conversion is \sim 7.9 MJ mol⁻¹, yielding an energy efficiency of only 4.4% with respect to the reaction enthalpy of 283 kJ mol⁻¹ at 300 K. On the other hand, ca. 184 kJ mol⁻¹ would be needed to heat 1 mole of CO₂ to 3500 K, and at this temperature the reaction enthalpy is equal to 206 kJ mol⁻¹. Hence, based on a conversion of 80% at this temperature, the energy cost of the total conversion is then only ~ 602 kJ mol⁻¹, yielding an energy efficiency of 47% with respect to the reaction enthalpy of 283 kJ mol⁻¹ at 300 K. While the conversion continuing to increase, above 3500 K the energy efficiency starts to decrease. At 5000 K, the conversion is 100% but the energy efficiency is only 35%, as can be deduced from Fig. 2.

Thus, it is clear that the equilibrium production of CO and O2 varies from less than 1% at temperatures below 2000 K up to 45-80% at the temperature range 3000-3500 K.29,30 Therefore, the most pertinent studies regarding thermal CO2 splitting involve membrane reactor systems. Nigara and Cales³² used a calcia-stabilized zirconia membrane and CO as the sweep gas. At a temperature of 1954 K, they were able to reach a conversion of 21.5%, whereas the equilibrium production was a mere 1.2% at the same temperature.³² The overall conversion, however, was much lower due to the permeation of O2 through the membrane, whereby it recombined with the CO sweep gas to form CO2. Itoh et al.33 employed an oxygen permeable yttriastabilized zirconia membrane and used argon as the sweep gas. Unfortunately, despite the removal of oxygen through the membrane, conversions of only up to 0.5% were obtained for a maximum temperature of 1782 K. Fan et al.34 used a solid

oxide (SrCo_{0.5}FeO₃) membrane reactor and methane as the sweep gas. Conversions of up to 10% were found at a temperature of 1213 K, which are one to two orders of magnitude higher than what could be expected to be attained conventionally. Nonetheless, the feed gas was diluted with four parts of helium per one part of CO₂, and when this is taken into account, an effective conversion of only 2% was actually reached.

To summarize, these studies have demonstrated the possibility of producing CO and O2 by the direct thermal splitting of CO₂ - at lower temperatures than the equilibrium predictions by means of the use of semipermeable membranes to extract oxygen. Nevertheless, the attained overall effective conversions of 0.5-2% are too low to be considered practical for successful application on an industrial scale.

Because to date none of the above-mentioned alternative approaches to split CO2 at lower temperatures have yet realized acceptable conversions and energy efficiencies, thermocatalytic CO₂ splitting is currently not applied on an industrial scale. The reason for this is the high energy consumption and, in addition, the lack of effective techniques for separating CO and O₂ at high temperatures to avoid ending up with an explosive mixture. Nevertheless, we presented this brief summary of the initial efforts regarding thermal CO2 splitting to support a complete understanding of the matter, and because this work has laid the foundations for one of the novel technologies, i.e. solar thermochemical CO₂ splitting, discussed later (see Section 3.2).

2.2. Conversion of CO₂ with a co-reactant

Due to the inherent high energy consumption and derived low energy efficiency of thermocatalytic CO2 splitting, the only practical way to reform CO₂ consists of using a co-reactant. Thermodynamically speaking, it is significantly easier to convert CO₂ when it is paired with a co-reactant that has a higher, i.e. less negative, Gibbs free energy.24 Some suitable candidates are CH₄ ($\Delta G^{\circ} = -50.7 \text{ kJ mol}^{-1}$) and H₂ ($\Delta G^{\circ} = 0 \text{ kJ mol}^{-1}$). In essence, these hydrogen-bearing energy carriers give up their intrinsic chemical energy to promote the conversion of CO₂.

As such, it is no surprise that the most widely investigated traditional processes to convert CO2 involve the reaction with either CH₄ or H₂. The former is one of the best known traditional processes for reforming CO₂ into synthesis gas or syngas, which is a mixture of H2 and CO. The reaction with H2 is known as the Sabatier reaction, which is a well-known process to generate CH₄ (and H₂O). Additionally, the combination of CO2 and H2 can also be used to produce methanol through the methanol synthesis process. A final process of interest to mention is the combined conversion of CO2 and H₂O, a technique for which there is no real traditional approach. Nevertheless, we briefly mention it here because water is an interesting co-reactant to pursue for the growing array of novel techniques. After all, H₂O is not only the most ubiquitous and cheapest hydrogen source, compared to CH₄ and H2, but converting CO2 in combination with H2O to produce value-added products using renewable energy would successfully mimic natural photosynthesis. 35,36 It is interesting to note that besides CH₄, H₂ and H₂O, other possible hydrogen

sources could also be considered, such as glycerol. The use of glycerol as a hydrogen donor has already been suggested and successfully implemented for different technologies, such as its use as a hydrogen donor (and green solvent) in catalytic transfer hydrogenation-dehydrogenation reactions of various unsaturated organic compounds;37 its use as a hydrogen donor for the production of CH₃OH from CO₂ in a hybrid enzymatic/photocatalytic approach³⁸ and its use in the plasma-based reforming of glycerol towards syngas, 39 which also led to the suggestion of using glycerol as a hydrogen donor for the possible in situ trapping of oxygen during the plasmachemical splitting of CO2.40 However, in this review, the focus lies on the above-mentioned three most commonly used hydrogen sources CH₄, H₂ and H₂O.

2.2.1. CO₂ + CH₄: dry reforming of methane. The combined conversion of CO2 and CH4, known as the dry reforming of methane (DRM), is named analogous to its sibling conversion, namely the steam reforming of methane (SMR; reaction (2) above) - indicating the replacement of water by carbon dioxide:

$$CH_4(g) + CO_2(g) \rightarrow 2CO(g) + 2H_2(g) \quad \Delta H^\circ = +247 \text{ kJ mol}^{-1}$$
(R3)

This process is, however, not as straightforward as the steam reforming of methane, because CO2 is a highly oxidized, thermodynamically stable molecule, while its reaction partner, CH₄, is chemically inert. Hence, the process needs to be carried out at high temperatures (900-1200 K) in the presence of a catalyst, typically containing Ni, Co, precious metals or Mo₂C as the active phase. 41,42 Fig. 3 illustrates the theoretical thermal conversion and energy efficiency as a function of temperature. At 1500 K, complete conversion is achieved, with an energy efficiency of 60%. However, the maximum energy efficiency of 70% is obtained before this at 1000 K, reaching a conversion of maximum of 83%, but this then decreases with increasing the temperature.

The dry reforming of methane (DRM) has quite a history. It was first studied by Fischer and Tropsch in 1928,42 and has been a challenge for chemical engineering ever since.⁴¹ Since that time, the rationale for investigating this process has

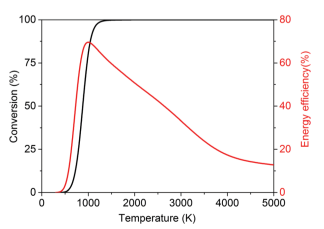


Fig. 3 Calculated theoretical thermal conversion (left axis) and corresponding energy efficiency (right axis) as a function of temperature for the dry reforming of methane

adapted itself several times to the spirit of the age. In its origin, it arose from a desire for alternative ways to produce fuels and chemicals (in combination with the Fischer-Tropsch synthesis) due to the limited supply of fossil fuels during the Second World War. 43 A renewed interest was found in the 1970s in the aftermath of the oil crisis, 43 again to circumvent the need for fossil fuels and with the idea of utilizing cheaper and more abundant natural gas. With the beginning of a new millennium and the increasing concern regarding climate change, DRM was seen as a way to convert the major greenhouse gas CO2 into useful products with the aid of natural gas. 31,44 To date, a true amalgam of environmental and economical motivations exist, such as the conversion of the greenhouse gas CO₂, the capability of using biogas as a feedstock, the search for a convenient way to liquefy CH₄ for easier transport, and the availability of cheap CH₄ through shale gas. 12,18,31,41,42,44-46

Alas, despite all the bright outlooks, there is one major pitfall, namely the process' inherent susceptibility for soot deposition and the detrimental effect this has on the process through deactivation of the catalyst. Due to this drawback, DRM is to date not yet (widely) used on an industrial scale. Of course, a lot of research is still ongoing towards modified catalysts to circumvent this coking issue, which was originally also a big problem for the currently widely adopted steam reforming of methane. Nevertheless, the inability to transform the alluring promises of DRM into reality through the traditional thermal methods - among other reasons- has sparked and fuelled the growing interest for alternative reforming technologies, as is discussed in Sections 3 and 4.

2.2.2. $CO_2 + H_2$: hydrogenation of CO_2 . Both the complete hydrogenation of CO2 to CH4, known as the Sabatier reaction or the methanation of CO2, and the selective hydrogenation of CO₂ to methanol are well-known commercially interesting processes. 47 The catalytic hydrogenation of CO₂ to methane is a thermodynamically very favourable process:⁴⁸

$$CO_2(g) + 4H_2(g) \rightarrow CH_4(g) + 2H_2O(g) \quad \Delta H^\circ = -165.3 \text{ kJ mol}^{-1}$$
(R4)

However, due to the high oxidation of the carbon, its reduction consists of an eight-electron process, significantly limiting the reaction kinetics and requiring a catalyst with high rates and selectivities.48 The process has been extensively studied using various supported nickel catalysts. 48,49 CO2 conversions of >95%, with the methane selectivity going up to 100% at temperatures of 700 K, have already been achieved. 49 However, for industrial commercialization, this process is only viable when the H2 is produced from renewable energy and the CO2 comes from cheap accessible waste streams. 10,49 As mentioned above, 95% of the worldwide H2 production, however, comes from steam methane reforming, leading to a problematic flawed loop. Furthermore, the current cost for CO₂ capture, separation and purification from waste streams is too high. Both reasons make this process economically unfeasible. 49

The selective hydrogenation of CO2 to methanol, on the other hand, is a process that is currently operated on an industrial scale. The annual worldwide production of methanol **Review Article** Chem Soc Rev

is estimated to be around 70 M metric tonnes (2015). The most common commercial catalyst is copper supported on high surface area alumina (often promoted with zinc oxide).⁴⁷ The relevant reactions for the selective hydrogenation of CO₂ are:⁵⁰

$$CO_2(g) + H_2(g) \rightarrow CO(g) + H_2O(g) \quad \Delta H^\circ = +40.9 \text{ kJ mol}^{-1} \quad (R5)$$

$$CO(g) + 2H_2(g) \rightarrow CH_3OH(g) \quad \Delta H^{\circ} = -90.8 \text{ kJ mol}^{-1} \quad (R6)$$

$$CO_2(g) + 3H_2(g) \rightarrow CH_3OH(g) + H_2O(g) \quad \Delta H^\circ = -49.9 \text{ kJ mol}^{-1}$$
(R7)

While the overall reaction (reaction (7)) of CO₂ hydrogenation to methanol is exothermic ($\Delta H^{\circ} = -49.9 \text{ kJ mol}^{-1}$), the rate determining step is the activation of CO2 in the reverse watergas shift (RWGS) reaction, i.e. reaction (5). Obviously, doping metals that function as catalysts in the RWGS reaction will promote CO₂ hydrogenation.⁵⁰ As a result, much effort is still focused on investigating the efficacy of Cu-based catalysts promoted with Pd and Ga,51 with the fundamental material challenge centring on the fact that, generally, CO2 and H2 will only react at high temperatures with multicomponent heterogeneous catalysts.24

One of the main drawbacks of the selective hydrogenation of CO2 to methanol in the above case is the production of water as a by-product (see reaction (7)). A third of the H₂ is thus converted to water compared to the complete conversion to methanol when starting from syngas (see reaction (6)). Furthermore, the thermodynamics for methanol production from H₂ and CO2 are not as favourable as those for the production from syngas (cf. reactions (6) and (7)).⁵¹ Therefore, on an industrial scale, methanol production usually relies on syngas in a 3 to 1 ratio from SMR (reaction (2)), while CO2 is added to deal with the excess H₂ in the feed (compared to reaction (6)), and finally the produced water (reaction (7)) is recycled via the water-gas shift reaction (the reverse reaction of reaction (5)).

Nevertheless, to conclude, we can state that, currently the selective hydrogenation of CO2 with H2 into methanol is the most - if not to say only - industrially successful traditional process for the direct reforming of CO₂ into chemicals and fuels.

2.2.3. CO₂ + H₂O: artificial photosynthesis. Although there is no real traditional approach for the combined conversion of CO2 and H2O, we present here the main overall reactions of interest for the combined conversion of CO₂ and H₂O, mostly for the sake of completeness and because of their interest to the discussion of novel technologies (see Sections 3 and 4):

$$CO_2(g) + H_2O(g) \rightarrow CO(g) + H_2(g) + 1/2O_2(g)$$
 $\Delta H^{\circ} = +525 \text{ kJ mol}^{-1}$
(R8)

$$CO_2(g) + 2H_2O(g) \rightarrow CH_3OH(g) + 3/2O_2(g)$$
 $\Delta H^{\circ} = +676 \text{ kJ mol}^{-1}$
(R9)

These are clearly the most endothermic overall reactions described in this section, partially explaining the absence of a traditional (thermocatalytic) reforming approach. Fig. 4 illustrates that the same high temperatures are needed for this reaction as for the pure CO₂ splitting, while obtaining somewhat lower energy efficiencies. The highest energy efficiency (40%) is

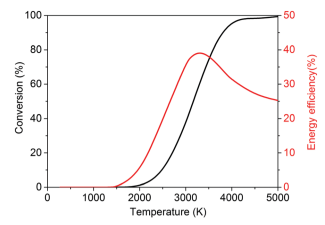


Fig. 4 Calculated theoretical thermal conversion (left axis) and corresponding energy efficiency (right axis) as a function of temperature for the conversion of CO2 and H2O into H2, CO and O2

obtained at 3300 K together with a conversion of 60%. At higher temperatures, the energy efficiency decreases to 25% at 5000 K for a total conversion.

Nevertheless, a process involving the reaction of CO₂, H₂O, O2 and CH4, called tri-reforming, is gaining quite some interest.24 The concept, proposed by Song et al.,31 involves a synergetic combination of dry reforming, steam reforming and the partial oxidation of methane in a single reactor, which can produce syngas in desired ratios (1.5-2.0), while eliminating carbon formation, as demonstrated for a fixed bed flow reactor at 1123 K over supported nickel catalysts.31

3. Novel CO₂ conversion approaches

It stands without doubt that the efficient conversion of CO₂ to useful molecules presents an important challenge and a great opportunity for chemists today. Due to the inability of the traditional thermal approaches to address the worldwide CO₂ and energy challenge, several promising novel technologies are under development. Plasma technology is one such technology, but before elaborating on this specific technology in more detail, first, a summary of its main 'frenemies' in this domain is given. For each technology, first a brief explanation of the working principles and current achievements is given, followed by the major advantages and challenges. From this section, it will become clear that there is, indeed, fierce competition in the quest to find the most effective and efficient CO2 conversion technology with the potential to be used on an industrial scale. It should be noted that only technologies for CO2 conversion are described here. For other (in)direct applications and fixation technologies, we refer to other reviews. 12-19

3.1. Electrochemical conversion

We kick off our discussions with one of the closest competitors to plasma technology, i.e. the electrochemical conversion or reduction of CO2. This closeness derives from the fact that both technologies rely on the use of (renewable) electrical energy, whereas most of the other novel technologies only take direct

advantage of renewable energy, i.e. the sun, either based on its focused radiation heat or its emitted photons. Although significant technical and catalytic advances are still required for its large-scale use, electrochemical conversion is becoming a mature technology for H₂O splitting. For CO₂ reduction, on the other hand, several important challenges remain.7,45,52-55

The electrochemical valorization of CO2 is an innovative technology, in which electrical energy is supplied to establish a potential between two electrodes, allowing CO2 to be transformed into value-added chemicals under mild conditions.^{7,55} This transformation can occur through a wide variety of pathways, which are typically strongly affected by the experimental conditions. The electrochemical reduction of CO₂ can proceed through two-, four-, six- and eight-electron reduction pathways in gaseous, aqueous and non-aqueous phases in different cell and electrode configurations.^{7,53} Fig. 5 shows the three main cell types. Fig. 5(a) and (b) illustrate the principle of a solid proton conducting electrolysis cell (SPCEC) for the combined conversion of CO₂ and H₂O, and of a solid oxide electrolysis cell (SOEC) that could be used for either the pure or combined conversion of CO₂ and H₂O, respectively, while a typical alkaline electrolysis cell for water splitting is shown in Fig. 5(c). The catalyst and/or electrode materials, the reaction medium, electrolyte solution, buffer strength, pH, CO2 concentration and pressure as well as the reaction temperature all influence and determine the wide variety of products that can be obtained.⁵³ The major reduction products obtained include carbon monoxide (CO), formic acid (HCOOH) or formate (HCOO⁻) in basic solution, formaldehyde (CH2O), methanol (CH3OH), oxalic acid $(H_2C_2O_4)$ or oxalate $(C_2O_4^{2-})$ in basic solution, methane (CH_4) , ethylene (C₂H₄) and ethanol (C₂H₅OH).^{7,45,53}

There are a number of reasons why the electrochemical reduction process stands out from the crowd; for instance, the process is controllable by several reaction parameters, including the electrode potential and temperature. 7,45,52-54 Furthermore, a wide variety of valuable products can be made, either in mixtures or more importantly in their pure form. For example, besides the direct electrochemical reduction of CO2 to methanol, it is also possible to produce CO and H2 at the cathode in a H₂/CO ratio close to 2, while at the anode, a

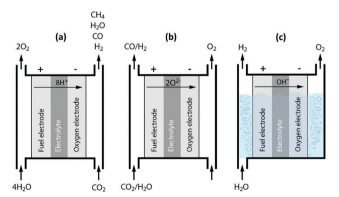


Fig. 5 Principles of a solid proton conducting electrolysis cell (SPCEC) (a), solid oxide electrolysis cell (SOEC) (b) and an alkaline electrolysis cell (c) for the conversion of CO₂ and/or H₂O₃

valuable pure oxygen stream is generated (see Fig. 1-5(b)).45 Another advantage is that electrochemical conversion can make use of a wide variety of (intermittent) renewable electricity sources, i.e. more than just solar energy.⁵³ Finally, the electrochemical reaction systems are compact, modular, on-demand and thus easy to utilize for small or large scale-up applications.⁵³

The accomplishments to date in the electrochemical reduction of CO₂ have been encouraging, and the potential rewards are enormous.⁷ Nevertheless, several challenges remain, such as the high overpotential, which is the difference in electrode voltage between the theoretical thermodynamic and actual real-world values to drive a reaction; 52,54 the low solubility of CO₂ in aqueous solutions;⁵² the formation of product mixtures, thus requiring expensive separation steps;⁵² fouling and catalytic deactivation of the electrodes by impurities, reaction intermediates and by-products;7,52,54 the instability of the electrode material;⁷ the low Faradaic efficiencies, current densities and high energy consumption; 52,53 the kinetic barriers leading to low efficiencies; 53,54 and the non-optimized electrode/ reactor and system design for practical applications.⁵³ In general, it is recognized that the single biggest challenge is the low performance of the electrocatalysts, due to low activity, low selectivity and most importantly insufficient stability. The reported stability tests in the literature are only in the order of, or below, 100 hours, while long-term tests are non-existent to date.⁵³ This makes the development of stable electrocatalyst materials with high activity and selectivity the main priority for this technology.7,53,56

It seems that despite many advances and successful proofof-concepts being reported, the maturity of electrochemical CO₂ reduction technology is still far from reaching the requirements for commercialization, due to the several remaining major technological challenges, as listed above.7,52,53 Particularly for industrial-scale implementation, the low catalyst stability seems to be the major limitation.⁵³ As a result, no electrocatalysts developed to date for the reduction of CO2 would be deemed useful for a large-scale system.⁵⁴ As stated by Qiao et al.,⁵³ "With continued and extensive efforts focused on developing innovative composite and nanostructured catalyst materials to overcome the challenges of insufficient catalytic activity, product selectivity, and catalytic stability, the technology of CO₂ electroreduction will become practical in the near future". Hence, to successfully achieve the transformation of CO2 to liquid fuels and useful chemicals, new methods and approaches for activating the CO2 molecule at lower overpotentials are required.⁷ In the first instance, novel electrodes enabling operation at current densities close to commercially available H2O electrolyzers have to be developed, for which solid oxide electrodes appear to be suitable candidates.⁵² Furthermore, a better understanding of the mechanistic role of metal and metal oxides in the reduction process is needed to open the possibility to design electrodes with certain compositions.⁵² To conclude, efforts to optimize system designs and at the same time to develop durable catalysts still need to be carried out.53 However, the final grand question remains: can all of this be done with inexpensive earth-abundant metals?54

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3.2. Solar thermochemical conversion

Another technology, which has recently made several huge leaps forward, is the solar thermochemical conversion of CO₂. There are several ways to reduce CO2 with the assistance of renewable solar energy, with those using direct solar light irradiation probably the most effective methods because there is no additional extra energy required and no negative influence on the environment.⁵⁷ Two forms of direct solar energy conversion can be distinguished: (i) thermal conversion - described here - where work can be extracted after sunlight is absorbed as thermal energy and (ii) quantum conversion - described in the next section - where the work output can be taken directly from the light absorber (e.g. a semiconductor, molecule or organic compound).⁵⁸ For solar thermochemical conversion, concentrated solar radiation is used in the form of high-temperature heat – as an energy source to drive the highly endothermic reactions.

The single step thermal dissociation of CO₂ (or H₂O) is impeded by the need to operate at high temperatures (>2500 K), as demonstrated in Fig. 2 above, and the need for effective separation techniques to avoid ending up with an explosive mixture of CO/O2 (or H2/O2).58,59 Multi-step thermochemical cycles using metal oxide redox reactions bypass the separation problem and, in addition, they allow operation at relatively moderate temperatures. More specifically, as shown in Fig. 6, solar processes involving heat at \geq 1500 K enable a two-step thermochemical cycle using metal oxide redox reactions for CO₂/H₂O-splitting.^{58,59} The first – endothermic - step is the solar thermal reduction of the metal

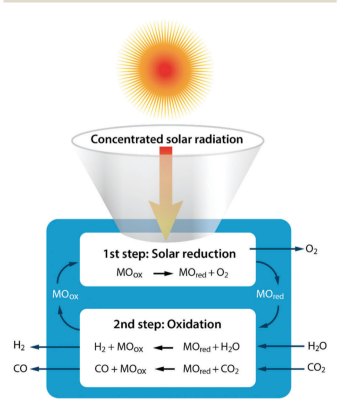


Fig. 6 Schematic of the two-step solar thermochemical cycle for CO₂ and H₂O splitting based on metal oxide redox reactions.

oxide MO_{ox} (where M is e.g. Ce, Zn, or Fe)⁶⁰ to the metal or to the lower-valence metal oxide MO_{red}. The second - nonsolar exothermic step is the oxidation of the reduced metal oxide with CO₂ and/or H₂O to form CO and/or H₂, allowing the (re)oxidized metal oxide to be reused/recycled for the first step. 58,59 In general, two cycle categories can be considered: volatile and non-volatile. Non-volatile cycles utilize metal oxides, which remain in the solid state during reduction, while volatile redox cycles consist of metal oxides, which undergo gas-solid phase transitions. The volatile reactions appear more favourable, but the volatile products must be quenched rapidly to avoid recombination, and to date this issue has not been solved in an energetically efficient fashion.⁵⁹ For the non-volatile cycles, cerium oxide (ceria, CeO2) has emerged as a highly attractive redox active material choice for two-step thermochemical cycling. 60,61 Another promising pathway, which operates at lower temperatures than ceria, is the exploration of doped perovskite oxides.62

The main advantage of the solar thermochemical conversion of CO₂ is obviously the direct use of solar energy. Concentrating solar technologies, which are currently applied commercially for large-scale (megawatt) power generation, can be coupled to high-temperature thermochemical reactors with the potential to achieve high solar-to-fuel energy conversion efficiencies and, consequently, the potential to produce solar fuels on a large scale and at a competitive cost.⁵⁸ To date, solar flux concentration ratios exceeding 2 MW m⁻² are attainable with largescale solar tower and dish systems. Solar thermochemical applications, although not as far developed as solar thermal electricity generation, employ the same solar concentrating infrastructure, with the solar reactor positioned at the focus of the solar tower (for megawatt centralized applications) or solar dish (for kilowatt decentralized applications).58 A recent comprehensive review of solar concentrating technologies for thermal power and thermochemical fuel production was given by Romero et al. 63 Consequently, these cycles inherently have the potential to realize a greater theoretical efficiency than methods using energy vectors or a small part of the solar spectrum and are, in addition, conceptually simpler. 62,64 This potential to achieve high solar-to-fuel energy conversion efficiencies is primarily related to the fact that solar thermal processes inherently operate at high temperatures and utilize the entire solar spectrum, and, as such, provide a thermodynamically favourable path to the production of solar fuels.^{58,59} A thermodynamic analysis based solely on the material properties of e.g. CeO₂ indicated that efficiency values in the range of 16-19% could be attainable, even in the absence of sensible heat recovery. These values are close to the 20% efficiency that is likely to be needed for solar fuels to be considered cost competitive, 65 as will be discussed in more detail in Section 6.3.

Although significant advances have been made in the field of solar thermochemical CO2 conversion technologies using metal oxides, a lack of fundamental research into the behaviour of the metal oxides under the high-temperature conditions present in these cycles has hampered the development of materials. Basic questions relating to oxygen transport, surface

chemistry, structural changes vs. redox reactions, materials' synthesis methods, the effects of thermochemical cycling on the material and the role of supports still have to be addressed.⁶⁶ Furthermore, despite its favourable thermodynamics, both the efficiency and the cycling rates in the reactor can be largely limited by thermal losses, resulting from poor conductive and radiative heat transfer across the porous metal oxide structure.⁵⁸ Finally, the thermochemical conversion rates are higher than, for example, the photocatalytic rates, but although conceptually simple, focusing lenses for sunlight and high-temperature reactors incur high initial investment costs.60

Solar thermochemistry has clearly emerged as a viable path to utilize concentrated solar technology - currently applied commercially for large-scale power generation - for the conversion of CO₂ (and H₂O) into CO (and H₂). Also, solar thermochemical cycles for the conversion of CO2 and H2O via the metal oxide redox reaction have favourable thermodynamics, but the ultimate factor dictating commercial viability is a high solar-to-fuel energy conversion efficiency, and to date efficiencies above 10% are still pending experimental demonstration with robust and scalable solar reactors. 58,59,62 The discovery of new materials with large oxygen exchange capabilities at moderate temperatures and their implementation in efficient solar reactors are thus essential. Additionally, rapid chemical kinetics and material stability over thousands of cycles must be demonstrated for each material considered. 59,67 This is the second key to achieving market viability, because materials must remain active for many thousands of redox cycles in order to avoid the high costs that would be associated with frequent replacement. As such, commercial success is again predicated upon finding appropriate materials composed of earth-abundant elements that can operate at lower reduction temperatures than current systems, together with sufficient activity to achieve high process efficiency.62

3.3. Photochemical conversion

The photochemistry for the photochemical conversion differs from (solar) thermochemistry in the way the solar energy is used: the former uses the energy of a photon in the chemical reactions, 60 while the latter uses the absorbed thermal energy to overcome the activation barriers and to affect the chemical equilibria.⁵⁸

The photoreduction of CO₂ to formaldehyde and methanol in purified water was already reported back in 1979, using the semiconductors TiO2, ZnO, CdS, GaP, SiC and WO3.60 Based on a correlation between the conduction band energy potential and the yield of methanol, it was suggested that the photoreduction of CO₂ proceeds by the photoexcited electrons in the conduction band moving to CO2. This principle mechanism of selective photocatalysts under light irradiation is shown in Fig. 7. Here, the conduction band energy minimum is higher than that for CO₂ photoreduction. 60 Again the efficiency of the photocatalytic materials in their use of sunlight for the conversion of CO₂ to fuel is of critical importance.⁶⁸ This efficiency is influenced by several factors, such as catalyst dosage, reactant ratio, reaction temperature, time, system pressure, pH, light

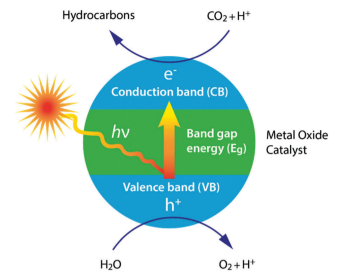


Fig. 7 Principle of the photochemical reduction of CO₂ by water on a photocatalyst.

intensity and wavelength.⁵⁷ A wide variety of reduction products can be obtained, just like with the electrochemical technique, including carbon monoxide (CO), formic acid (HCOOH), formaldehyde (CH₂O), methanol (CH₃OH), methane (CH_4) , ethylene (C_2H_4) , ethane (C_2H_6) and ethanol (C_2H_5OH) .

From a sustainable point of view, solar light is the ideal energy source. In combination with photocatalytic H2O splitting, the solar-driven reduction of CO2 to fuels is a very attractive approach.52 The advantages of photochemical systems include the assertion that they are composed of only a few parts and are therefore theoretically less likely to fail, providing the remaining parts are reliable.⁵⁸ The most extensively investigated catalyst for the photoreduction of CO₂ is TiO₂. 60 Several attempts have already been made to enhance the photocatalytic activity of TiO2, including by the addition of a metal, Rh/TiO2 or Rh/WO3-TiO2,60 the use of highly dispersed active Ti ion species,60 through atomically dispersing TiO2 on zeolites or ordered mesoporous SiO₂ or by doping with Pt, Cu, N, I, CdSe or PbS.⁶⁰ The most significant breakthrough was achieved using nitrogendoped TiO2 nanotube arrays co-catalyzed with Cu and/or Pt nanoparticles, in which water-vapour-saturated CO₂ was reduced to methane and other hydrocarbons without the application of an electrical bias. 60,68

It should be noted, however, that for many oxide semiconductors (incl. TiO2) this electrical bias is necessary, because the conduction band is located below the acceptor level. This is one factor that limits the efficiency of metal oxide materials.⁵⁸ Furthermore, most work is performed using artificial (UV) light sources, 57,68 because the large band gap of metal oxides results in a poor photo-responsiveness to visible light. 52,57,58 Theory dictates that a band gap between 2 and 2.4 eV is optimal, which limits the maximum attainable efficiency to about 17%.68 However, solar energy conversion efficiencies obtained to date are much lower (at present, at < 2%), ⁵⁸ mainly as a result of the energy associated with this electrical bias.58 Furthermore, a remaining challenge lies in the separation and collection of the

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hydrogen and oxygen gas produced (often produced in close proximity). ⁵⁸ Finally, many of the photocatalysts presently being studied are metal complexes employing rare and expensive transition metals, hence efforts must be made using earth-abundant elements that could support large-scale undertakings. ⁵⁴

From the literature, it is clear that encouraging progress has been made towards the photocatalytic conversion of CO2 using sunlight, but nevertheless, the existing techniques are insufficient to date and further efforts are required to increase the solar-to-fuel conversion efficiencies. 54,55,57,58,68 This appears to be a general remark for photochemistry: very few examples exist of chemical processes operating effectively on the basis of photocatalysis technology. Not only do the photon efficiency of materials and the resulting achievable rates remain insufficient, also sub-optimal photocatalytic reactors often induce inefficiencies in operation, which limit their practical application.⁵² The immediate requirement in this technology is to develop visible-light-sensitive photocatalysts, which are prominent in CO₂ recycling.⁵⁷ What we ultimately seek is a means for achieving the high-rate photocatalytic reduction of CO₂, using solar radiation as the only input energy source. Since visible light comprises the majority of the solar spectrum energy, it behoves us to consider photocatalysts sensitive to sunlight.⁶⁸ Immediate research opportunities include uniform co-catalyst sensitization of the entire nanotube array surface for enhanced conversion rates, and the design of co-catalysts to improve and control the product selectivity.68 Although the photocatalytic reduction of CO2 may become an important stepping stone to solar fuel production, much progress remains to be achieved before it could be considered practical as an industrial process.⁵⁴ Based on the highest reported activities, one can conclude that game-changing rates have not yet been achieved. Reported turnover frequencies are far from those required for an efficient catalytic process, and an efficiency improvement of at least 3 orders of magnitude is needed.⁵² It is thus clear that photochemical systems have a long way to go to achieve their full potential and to be able to successfully compete with the alternative approaches to producing fuels from sunlight.58

3.4. Biochemical conversion

Another pathway converting solar energy into chemical energy is by 'natural' photosynthesis for the production of biofuels. ⁶⁹ The biological conversion of CO_2 for producing chemicals or fuels is an attractive route. Nevertheless, the use of first generation biofuels has generated a lot of controversy, mainly due to their competition with agriculture for arable land use for food production, thus impacting global food markets and food security. ^{69,70} The use of microalgae, on the other hand, could meet the conditions for technically and economically viable biofuel production. More specifically, viable biofuel production should be competitive or cost less than petroleum fuels, it should require low to no additional land use, it should enable air quality improvement and should require minimal water use. ^{69,70} Microalgae can typically be used to capture CO_2 from three different sources: atmospheric CO_2 , CO_2 emissions from

power plants and industrial processes, and CO₂ from soluble carbonates.⁷⁰ The pathways for CO₂ fixation have evolved over billions of years and use many diverse mechanisms and enzymes for processing CO₂ by forming C–H and C–C bonds and cleaving C–O bonds.⁴⁷ Furthermore, algae are more photosynthetically efficient than terrestrial plants, making them without doubt very efficient CO₂ fixers.⁷¹

Microalgae are currently considered to be one of the most promising alternative sources for biodiesel; hence, most of the current research and developmental efforts are focused on microalgae, in particular due to their high growth rate and high oil content (up to 77% of the dry cell mass). Algae contain oils, sugars, and functional bioactive compounds that can be used for commercial products. In addition to fuels, the development of appropriate technologies for high-efficiency algal biodiesel production is also applicable to biohydrogen, biogas, bioethanol and biomass-to-liquid (BTL) approaches using fast growing algae. 69,70 Furthermore, valuable co-products, such as biopolymers, proteins and animal feed, can be produced during the process.71,72 Other advantages include very short harvesting cycles ($\sim 1-10$ days), thus allowing multiple or continuous harvests with significantly increased yields. Furthermore, the cultivation can potentially be carried out on marginal or non-arable land, and even the use of waste water for algal cultivation is a viable option. 69,70 To obtain the best performing microalgae strains for biofuel production, one can: (1) screen a wide range of natural isolates, (2) improve them by metabolic (genetic) engineering or (3) obtain them by selection and adaptation. There are algae collections worldwide, which contain thousands of different algal strains that can be accessed.69

It should, however, be emphasized that the significant drawback in all the biochemical techniques is the big share of the cost for cultivation. Among others, the harvesting of algal biomass accounts for the highest proportion of energy input during production, but currently, there are no standard harvesting techniques. 69-71 Therefore, currently algal biomass is not suitable to be cultivated solely for bioenergy applications and instead it must be integrated with the production of other value-added products, e.g. pharmaceutics, cosmetics and food. Unfortunately, processes for the recovery of complex molecules from algal biomass are expensive and significant technological progress is still required before commercial deployment.⁷³ Algae can be grown in many ways, such as in freshwater, saltwater or wastewater, in closed photobioreactors or in open ponds.⁶⁹ One key advantage of algae is that its cultivation does not require cropland, although, on the other hand, other resources are needed. 22 Other inorganic nutrients required for algae production include nitrogen and phosphorus, 70 resulting in the need for unsustainable inputs of nitrogenous fertilizers, which are produced from fossil fuels and that require huge inputs of energy for production.⁷² Furthermore, it has been reported that between 3.15 and 3650 litres of freshwater are needed to produce algal biofuel equivalent to 1 litre of gasoline using current technologies. Thus, the integration of the upstream production and downstream processing of

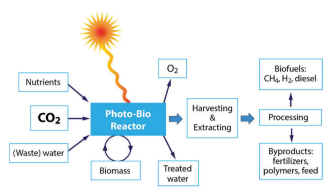


Fig. 8 Schematic of photo-bioreactor application for CO₂ conversion.

microalgae, and the framing of these in the context of water savings and net energy gain is required to build up credibility.⁷² Fig. 8 shows a schematic overview of the different steps for the application of a photobioreactor for CO2 conversion. Finally, optimization of the strain-specific cultivation conditions is complex, with many interrelated factors that can each be limiting factors, including temperature, mixing, fluid dynamics and hydrodynamic stress, gas bubble size and distribution, gas exchange, mass transfer, light cycle and intensity, water quality, pH, salinity, mineral and carbon regulation/bioavailability, cell fragility, cell density and growth inhibition.⁶⁹

Despite its inherent potential as a biofuel resource, many challenges have impeded the development of algal biofuel technology as a commercially viable solution that could support sustainable production and utilization.⁷⁰ Consequently, the large-scale cultivation of algae for biofuel production is still in the research and development phase. The long-term potential of this technology can be improved by the following approaches: (1) identifying and developing cost-saving growth technologies of oil-rich algae; 69,72 (2) utilizing integrated bio-refineries to produce biodiesel, animal feed, biogas and electrical power, thereby reducing the overall cost of production;^{71,72} (3) enhancing the algal biology by genetic modification and metabolic engineering to aid the selection and successful outdoor large-scale cultivation of a robust microalgal strain; ^{69,71,72} (4) identifying area-efficient techniques to capture CO₂ from industrial power plants;⁷² (5) recycling nutrients from municipal sewage and industrial wastewaters to reduce the demand for fertilizers to grow the algae; 69,70,72 (6) improving the economics of microalgae production by generating additional revenues from wastewater treatment and greenhouse gas emissions abatement;⁷² (7) most importantly, by developing cost-effective and energy-efficient harvesting methods to make the whole biofuels production process economical;⁷¹ in this respect, strain selection is an important consideration since certain species are much easier to harvest than others.69

3.5. Catalytic conversion

In Section 2, we already covered the main traditional thermocatalytic approaches used. Of course, there is still a lot of research ongoing towards finding new improved (thermo-)catalytic pathways for the different processes involved in CO₂ conversion. There are two main catalysis types that can be applied for this process: homogeneous and heterogeneous catalysts. The former (e.g. Ru-, Rh- and Ir-based catalysts) are efficient for the formation of formic acid and formates, but are more challenging to be used in commercial applications, while the latter (e.g. Fe-, Cu- and Ni-based catalysts) are more practical for industrial applications, but they frequently suffer from low yields and poor selectivity. As a result, significant improvements in new catalytic systems are necessary to make thermocatalytic CO2 reduction economically feasible.48

Nonetheless, the catalytic conversion is briefly discussed here; especially since it is clear from the sections above that catalytic materials can play an important role in the development and further advancement of most of the novel technologies under study.25 These scientific advances give rise to intriguing new combinations, and corresponding names, such as electrocatalytic, photocatalytic, biocatalytic, as well as their even more advanced hybrids forms, e.g. photoelectrocatalytic and bioelectrocatalytic processes. 12,14,53 Kumar et al. 54 stated quite frankly that virtually every approach under consideration for the conversion of CO2 requires catalysts to facilitate the formation and cleavage of chemical bonds, as illustrated by Fig. 9. In general, these required catalysts fall into three classifications: (1) they already exist and show good performance but are too rare/costly to be scaled up; (2) they already exist but in forms that are not optimal or practical for adaptation to an integrated solar fuels system; (3) they do not exist yet and await discovery.54

From a scientific point of view, the development of catalysts with inexpensive metals, such as iron and copper compounds, which can also be active under mild conditions, is a big challenge. 48 Still, it is evident that due to their reliance on thermal heat - as currently derived from burning fossil feedstocks - the current pure thermocatalytic routes can only make a limited net contribution to CO2 conversion. Hence, techniques based on the use of (in)direct solar energy and other renewable energy sources are needed to contribute to avoiding large volumes of CO₂. ²⁵ One exciting thermocatalytic advancement to circumvent this reliance on fossil fuels - besides the solar thermochemical conversion discussed in Section 3.2 above - is the use of microwaves.⁷⁴ Here, microwave-assisted CO₂ conversion over carbon-based catalysts combines the catalytic and dielectric properties of carbonaceous materials with the advantages of microwave heating, which favours catalytic heterogeneous reactions due to, among other reasons, the generation of hot spots.

Microwave radiation has been shown to have beneficial effects on the reaction rate of heterogeneous (catalytic) reactions. 75,76 The combination of microwave heating and a carbon material acting as both a catalyst and microwave receptor gives rise to enhanced conversions compared to conventional heating, both in the case of CO₂ gasification and in the dry reforming of methane.^{75,76} Fidalgo et al.76 observed conversions that were a factor of 1.6 to 1.9 higher for MW-heating compared to conventional heating. Unlike in conventional heating, the applied microwave energy is transferred

Without catalyst ΔG^{\ddagger} Reactant ΔG_{A}^{\ddagger} With catalyst ΔG_{A}^{\ddagger}

Fig. 9 Reaction coordinate diagram showing the working principle of a catalyst.

Reaction Coordinate

directly to the catalyst without any heat flux. As a result, the temperature inside the material is usually higher than the temperature of the surrounding atmosphere near the surface, and the uniformity of heat distribution is improved with respect to conventional heating, as shown in Fig. 10.⁷⁶ Furthermore, the formation of hot spots, possibly due to the generation of microplasma within the catalyst bed, which may be at higher temperatures compared to the bulk catalyst, have been reported to be responsible for an enhancement in the reaction rate, higher yields and the improved selectivities of heterogeneous (catalytic) reactions.^{75,76}

3.6. Summary

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It should be of no surprise that the field of CO₂ conversion is rapidly evolving. As such, it is not the purpose of this section to cover all the (recent) work performed for these novel technologies, such as CO₂ mineralization and utilization, which converts the chemical energy of CO₂ mineralization into electricity, while producing valuable mineralization products,⁷⁷ or the hybrid enzymatic/photocatalytic approach for the production of CH₃OH from CO₂ with the use of water–glycerol, as mentioned

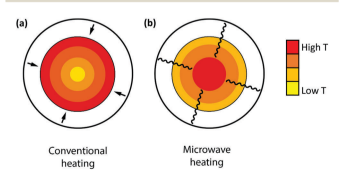


Fig. 10 Comparison of the temperature gradients produced by conventional (a) and microwave (b) heating.

in Section 2.38 Clearly, giving a complete comprehensive overview of all the carbon capture, utilization, and storage (CCUS) technologies is outside the scope of this review. Nevertheless, by discussing the main novel technologies for CO₂ conversion, we were able to roughly sketch out the current landscape, one in which we want to orient another emerging technology with the potential for CO₂ conversion, i.e. plasmachemical conversion. It is evident that, for all technologies discussed above, there are several distinctive up- and downsides. This will not be different for plasma technology - as will become evident from the following section. In the final benchmarking section (Section 6) of this review, we make a visual comparison between all the emerging technologies, based on their versatility, the mentioned distinctive advantages and limitations and - in what appears to be most important from an economical point of view - their solar-to-fuel conversion efficiency.

4. Plasma technology for CO₂ conversion

From the previous section, it is clear that several alternative (non-conventional) CO₂ conversion technologies are, more or less, successfully being investigated. In recent years, another novel technology has started to be considered, but has not spent much time in the spotlight yet: plasma technology. Herein, we first give a brief introduction to plasma technology, and highlight its general advantages and unique features for CO₂ conversion. Next, we describe the different kinds of plasma reactors used for CO₂ conversion, focusing on their advantages and disadvantages. Subsequently, we discuss the possibly fruitful combination of plasma with catalysts, in so-called plasma-catalysis. Finally, in the next section (Section 5), we present a critical assessment of plasma-based CO₂ conversion for these different set-ups, for both pure CO₂ splitting and CO₂ conversion, together with the specified co-reactants.

4.1. Properties of plasma and its unique features for CO₂ conversion

The term 'plasma' was first introduced by Irving Langmuir (1928). Plasma is an ionized gas, which means that at least one electron is unbound, creating positively charged ions. In practice, the ionization degree in plasma can vary from fully ionized gases (100%) to partially ionized gases (e.g. 10^{-4} – 10^{-6}). Besides the various types of ions (both positive and negative), plasma also consists of a large number of neutral species, e.g. different types of atoms, molecules, radicals and excited species. The latter can lead, among other things, to the emission of light. More importantly, all these species can interact with each other, making plasma a highly reactive and complex chemical cocktail, which is of interest to many potential applications. 21,78 Indeed, plasmas can already be found in several applications in materials science (e.g. coating deposition, surface modification, nanomaterial fabrication) and in the microelectronics industry (for microchip manufacturing), but also as light sources, lasers and displays (owing to their light emitting characteristics) as

well as for many emerging environmental and even medical applications (such as sterilization, wound treatment and even cancer treatment). More details about these applications can be found in the above-cited references, as well as in the references therein. In this review, however, we focus only on their application in CO₂ conversion into value-added chemicals and fuels.

Plasma is sometimes referred to as the 'fourth state of matter', owing to the observation that with increasing temperature matter transforms in the sequence: solid, liquid, (neutral) gas and finally an ionized gas or plasma. Although the concept of plasma is less known than the other states of matter, more than 99% of the visible matter in the universe is in the plasma state, mainly attributed to interstellar matter and the stars. Hence, our own sun is a perfect example of a plasma. Furthermore, basically, many - if not to say all - natural occurring weather phenomena emitting light are in fact plasma, e.g. Saint Elmo's fire, lightning, red sprites, auroras (Borealis and Australis), where we must thank the excited species for the emission of these colourful lightshows. Other natural plasmas close to home are the earth's ionosphere, plasma sphere and the outer magnetosphere.

Beside these natural plasmas, we can distinguish between two main groups of man-made plasmas. The first ones are the high-temperature or fusion plasmas, which are in general completely ionized plasmas. Applications include tokomaks, stellarators, plasma pinches and focuses. The second group comprises the weakly ionized plasmas or so-called gas discharges, which are under study and covered in this review.

A second sub-division can be made based on whether the plasma is in thermal equilibrium or not. The temperature in a plasma is determined by the average energies of the different species (electrons, neutrals, ions) and their relevant degrees of freedom (translational, rotational, vibrational and electronic). Since plasma is a multicomponent system, it can exhibit multiple temperatures. When the temperature of all these species is the same in a localized area, the plasma is said to be in 'local thermodynamic equilibrium' (LTE), and these kinds of plasmas are usually called 'thermal plasmas'. When the plasma is characterized by multiple different temperatures and thus is far from thermodynamic equilibrium, the plasma is said to be in 'non-local thermodynamic equilibrium' (non-LTE) and these discharges are usually called 'non-thermal plasmas'.

4.1.1. Thermal plasmas. Thermal plasmas can be achieved in two ways, either at high temperature, typically ranging from 4000 K to 20000 K, depending on the ease of ionization, or at high gas pressure. The latter can be explained as follows. Initially, the electrons receive energy from the electric field during their mean free path in between collisions, and they lose a small portion of this energy during collisions with so-called heavy particles (e.g. gas molecules or atoms). Subsequent collisions of this nature, also known as 'Joule heating', can lead to the temperatures changing to reach equilibrium between the electron and heavy particle temperature. At high pressures, the mean free path becomes smaller, so more collisions occur; hence, leading to a more efficient energy exchange between the electrons and the heavy particles. More specifically, it is the square of the ratio of the electric field (E) to the pressure (p), *i.e.*

 $(E/p)^2$, which is proportional to the temperature difference in gas discharges.21

Thermal plasmas have numerous advantages, compared to traditional technologies, due to their interesting characteristics, including high temperature, high intensity non-ionizing radiation and high-energy density. The heat source is also directional, with sharp interfaces and steep thermal gradients that can be controlled independently of the chemistry. Whereas the upper temperature limit when burning fossil fuels is 2300 K, thermal plasmas can reach temperatures of 20 000 K or more, as mentioned above. As a result, this type of plasma is already being used for a wide range of applications, such as for coating technology, fine powder synthesis, (extractive) metallurgy (e.g. welding, cutting) and the treatment of hazardous waste materials.79

On the other hand, the inherent nature of thermal plasmas makes them unsuitable for the efficient conversion of CO₂. More specifically, the ionization and chemical processes in thermal plasmas are determined by the temperature. As a result, the maximum energy efficiency is limited to the thermodynamic equilibrium efficiency and corresponding conversions of 47% and 80% at 3500 K, respectively (see Fig. 2 above). This is in contrast to non-thermal plasmas, where lab-scale efficiencies of up to 90% have already been reported (see further, Section 5.1).21

4.1.2. Non-thermal plasmas. In its simplest way, a nonthermal plasma is created by applying a potential difference between two parallel electrodes, which are inserted in a reactor filled with gas (or they form the reactor walls). This potential difference creates an electric field, which causes so-called gas breakdown, i.e. the gas 'breaks up' (to some extent) into positive ions and electrons, although the majority of the gas molecules still remain neutral. The electrons are accelerated by this electric field towards the positive electrode (anode). When they collide with gas molecules, it can give rise to ionization, excitation and dissociation. The ionization collisions create new electrons and ions; the ions are accelerated by the electric field towards the negative electrode (cathode), where they cause secondary electron emission. The new electrons - created by ionization or secondary electron emission at the cathode - can then further give rise to further ionization collisions. These processes make the plasma self-sustaining. The excitation collisions create excited molecules, which can decay to the ground-state (or another lower level), thereby emitting light, which is one of the characteristic features of plasmas. Finally, the dissociation collisions create radicals, which can easily form new compounds, and this forms the basis of the gas conversion applications of non-thermal plasmas.

Although this way of plasma creation, i.e. by applying a potential difference between two electrodes, is only one possibility to create a plasma, the principle of gas activation by electrons, creating reactive species (ions, excited species, radicals) is equally valid for other types of plasmas as well, and will be discussed in Section 4.2 below. In general, this gas activation by electrons is also the reason for the good energy efficiency of CO₂ conversion by non-thermal plasmas, as will be explained below.

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In non-thermal plasmas, the electrons are indeed characterized by a much higher temperature than the heavy particles, leading to the non-LTE condition, as mentioned above. As a result of all the different species, the relationship between all their different temperatures can become quite complex, but conventionally the temperature of the electrons (T_e) is the highest, followed by the vibrationally excited molecules (T_v) , while the lowest temperature is shared by the neutral species (T_0 , or simply the gas temperature, T_g), the ions (T_i) and the rotational degrees of freedom of the molecules (T_r) ; hence the temperature order is:

 $T_{\rm e} \gg T_{\rm v} > T_{\rm r} \approx T_{\rm i} \approx T_{\rm 0}^{21}$ In most cases, the electron temperature is in the order of 1 eV (~ 10000 K), while the gas temperature remains close to room temperature. This high electron temperature is due to the small mass of the electrons, allowing them to be easily accelerated by the applied electromagnetic fields, whereas the heavy particles - even the ions are not easily accelerated. Furthermore, due to the large mass difference, the electrons lose less energy during elastic collisions with heavy particles, so they can easily keep their high energy gained from the electric field.

As mentioned above, the electrons can be considered as the initiators of the highly reactive chemical mixture. This is obviously one of the key advantages of non-thermal plasma technology: it allows gases - even as unreactive as CO2 - to be 'activated' at room temperature by the highly energetic electrons. Accordingly, there is no need to heat the entire reactor or the gas, because the discharge and the associated reactions are easily initiated by applying an electromagnetic field. This results in the second key advantage of non-thermal plasma technology for CO₂ conversion, namely that it is a very flexible, or so-called turnkey, process, since it can easily and, more importantly, instantaneously be switched on and off, with conversion and product yield stabilization times generally lower than 30 minutes. Furthermore, its power consumption can easily be scaled and adjusted. As such, it has the inherent ability to be(come) one of the most suitable technologies to utilize excess intermittent renewable energy (e.g. energy originating from wind turbines or solar panels) and for storing it in a chemical form. Indeed, this makes it suitable for both peak shaving and, more importantly, grid stabilization, by being able to adapt to the irregular supply of renewable electricity. Hence, in essence the current transition to renewable energy sources not only gives plasma processes a clean electricity source, but plasma-based CO₂ conversion technology can also provide a solution for the imbalance between energy supply and demand by enabling the storage of electrical energy in a desirable chemical form according to the market needs.

Additional advantages compared to the other emerging technologies for CO₂ conversion are the low investment cost for the reactors, as they do not rely on rare earth materials, and last but not least, their simple scalability from watt to megawatt applications, as already demonstrated by the successful development of ozone generators. 80 Due to their extreme scalability, the applications can vary hugely in both scale and application type, starting from small devices, such as on board vehicles for exhaust treatment or to provide on board fuel cell feeds⁸¹ as well as in household-scale devices for indoor air treatment, 82,83 to medium on-demand installations, such as modular containerized plant concepts⁸⁴ or in in situ CO production, 85 and finally to largescale industrial plants. This scalability and flexibility in applications also gives plasma technology a high locational flexibility. As such, both production on demand and remote production become possible, which is a critical point for carbon capture and utilization (CCU) techniques.

On the other hand, for the comparison made here with other novel technologies, the reliance of non-thermal plasma technology on indirect solar energy in the form of electricity is at the same time a limiting factor for the overall solar-to-fuel conversion efficiency, compared to technologies that can harvest solar energy in a direct form. However, when looking at the broader picture, its ability to rely on multiple energy sources again gives plasma technology a huge advantage regarding locational flexibility, not to mention that it can be operated 24/7, even when the sun is not shining.

Although non-thermal plasma is good at creating a nonequilibrium condition by 'activating' stable molecules, it is absolutely non-selective in the formation of targeted products. More specifically, the reactive species created by the electrons react according to the laws of chemical kinetics, and as such, they recombine into a (large) number of different products, depending on the reaction conditions, not to mention that the formed products can again be destroyed by new electron collisions. For pure CO2 splitting, this is not much of an issue, since CO and O₂ (together with small amounts of O₃) are basically the only products that can be formed. However, when combining CO2 with other reactants, such as CH4, H2O or H2, a wide variety of products can be formed, including syngas (CO and H_2) and higher hydrocarbons (C_2H_x , C_3H_y , C_4H_z) as well as several oxygenates, e.g. methanol, ethanol, dimethyl ether, formaldehyde, acetaldehyde and carboxylic acids. For that reason, the combination with catalysis is also highly desirable for plasma technology when targeting the selective production of specific compounds (see Section 4.3).

4.1.3. Warm plasmas. Conventional thermal and nonthermal discharges cannot simultaneously provide a high level of non-equilibrium and a high electron density, whereas most prospective plasmachemical applications require both, i.e. a high power (as translated in a high electron density) for efficient reactor productivity and a high degree of nonequilibrium to selectively populate certain degrees of freedom, such as vibrationally excited states (see Section 4.1.4 below).86 Recent studies have revealed that a transitional type of plasma, so-called warm discharge or warm plasma, which operates at the boundary – and hence shares properties – of both thermal and non-thermal plasmas, might be very promising for CO2 conversion. These are non-equilibrium discharges, which are not only able to supply (re)active species, but also offer some controlled level of translational temperature. Although this translational gas temperature is still much lower than the electron temperature, it is significantly higher than room temperature and can easily reach up to 2000-3000 K. Hence, these warm plasmas are able to create the advantage of a nonequilibrium condition, while at the same time they can influence the chemical kinetics due to this higher gas temperature.

For the application of CO₂ conversion, however, it is not the increased temperature that makes these discharges most interesting, but rather their characteristic electron energy distribution, which leads to most of the electron energy going into the vibrational excitation of CO2, as will be discussed below. The latter is indeed known to provide the most energy efficient and hence most important channel for CO₂ dissociation. ^{21,87} In fact, in recent modelling studies, it was suggested that the higher gas temperature inherent to these warm discharges is actually an unwanted effect.88-90 As such, the development of 'cooler' warm discharges might be beneficial for further increasing the CO₂ conversion and energy efficiency.

4.1.4. CO₂ dissociation channels. For a better understanding of the difference between non-thermal and warm discharges, and specifically between the different discharge types described in Section 4.2 below, we provide an overview of the different channels of energy transferred by electrons to CO2. As mentioned, the electrons receive their energy from the electric field in non-thermal plasmas, and subsequently, through collisions, this energy is distributed between elastic energy losses and different channels of excitation, ionization and dissociation. Fig. 11 illustrates the fraction of the energy transferred to the different channels of the excitation, ionization and dissociation of CO2, as a function of the reduced electric field (E/n). The reduced electric field is the ratio of the electric field in the plasma over the neutral gas density and has distinctive values for different plasma types. For example, a dielectric barrier discharge (DBD), which is one of the most common types of non-thermal plasmas for CO₂ conversion (see Section 4.2.1), has a reduced electric field in the range above 200 Td (Townsend; 1 Td = 10⁻²¹ V m²), whereas microwave (MW) and gliding arc (GA) discharges (which belong to the category of warm plasmas; see Sections 4.2.2 and 4.2.3) typically operate well below this range (about 50 Td).

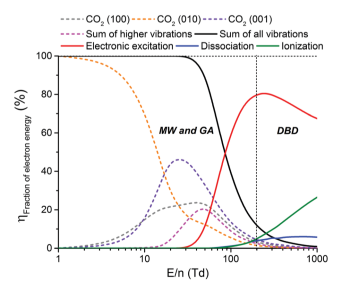


Fig. 11 Fraction of electron energy transferred to different channels of excitation, as well as to the ionization and dissociation of CO₂, as a function of the reduced electric field (E/n), as calculated from the corresponding cross-sections of the electron-impact reactions. The E/n regions characteristic for MW and GA plasma and for DBD plasma are indicated.

From Fig. 11, it is clear that the value of the reduced electric field will have wide implications on the distribution of the electron energy among the different channels. In the region above 200 Td, 70-80% of the electron energy goes into electronic excitation, about 5% is transferred to dissociation, 5% is used for ionization (increasing with E/n), while only 10% goes into vibrational excitation (decreasing with E/n). Around 50 Td, however, only 10% goes into electronic excitation and 90% of the energy goes into vibrational excitations. It is important to keep in mind that the addition of different gases (e.g. Ar, He, N2, H₂O, H₂, CH₄) has an influence on the distribution of these channels. 92 Hence, even during the pure decomposition of CO₂ into CO and O2, there will be an effect on this distribution.

The distribution of energy into different modes, and especially the fraction going into vibrational excitation, is very important, since, as mentioned above, it is known that the vibrational levels of CO₂ can play an important role in the efficient dissociation of CO₂. To achieve direct electron-impact dissociation, an electron needs to have enough energy (>7 eV) to excite CO2 into a dissociative (i.e. repulsive) electronic state, which will lead to its dissociation into CO and O (see Fig. 12). As such, the amount of

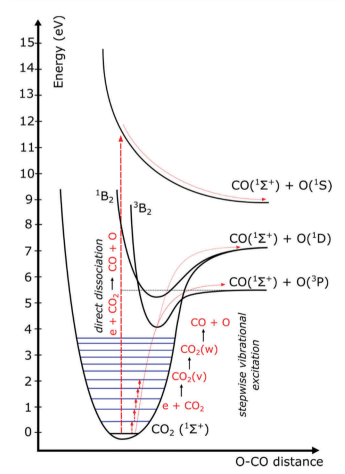


Fig. 12 Schematic of some CO₂ electronic and vibrational levels, illustrating that much more energy is needed for direct electronic excitationdissociation than for step-wise vibrational excitation, i.e. the so-called ladder-climbing process (reproduced from ref. 91 with permission from the Royal Society of Chemistry)

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energy spent is much higher than the theoretical value necessary for C=O bond breaking (5.5 eV). Due to the special nature of the CO₂ molecule, a more efficient dissociation pathway is based on its vibrational excitation. This pathway starts with electronimpact-vibrational-excitation of the lowest vibrational levels, followed by vibrational-vibrational (VV) collisions. This so-called ladder-climbing gradually populates the higher vibrational levels, which eventually leads to dissociation of the CO2 molecule (see also Fig. 12). In this way, it is possible to dissociate CO₂ more efficiently, since only the minimum amount of 5.5 eV for bond breaking is needed, compared to the overshoot in the case of electronic excitation-dissociation.91

4.2. Different types of plasma set-ups used for CO₂ conversion

As mentioned above, in recent years, there has been an increasing interest in the use of plasma technology for CO2 conversion. Experiments have been carried out in several types of plasmas. The most common types reported in the literature are dielectric barrier discharges (DBDs), microwave (MW) and gliding arc (GA) discharges, although other types have been used as well (e.g. radiofrequency, corona, glow, spark and nanosecond pulse discharges). A DBD is a typical example of a 'non-thermal plasma', where the gas is more or less at room temperature, and the electrons are heated to temperatures of 2-3 eV ($\sim 20\,000-30\,000$ K) by the strong electric field in the plasma. The MW and GA discharges are examples of 'warm plasmas' (see Section 4.1.3 above). The gas can reach temperatures of up to 1000 K and more, and the electron temperature is typically up to a few eV. The operating conditions and characteristic features of the three major plasma types for CO₂ conversion are explained here, as well as for some other plasma types that have also been applied for CO2 conversion.

4.2.1. Dielectric barrier discharge (DBD). Dielectric barrier discharges (DBDs), also called 'silent discharges', have been known for more than a century. The first experimental investigations were reported by Siemens in 1857 and were concentrated on the most well-known industrial application of DBDs, i.e. ozone generation. 80 For an extensive overview of the history, discharge physics and industrial applications of DBDs, we refer to the review of Kogelschatz et al.80

A DBD consists of two plane-parallel or concentric metal electrodes and, as its name suggests, it contains at least one dielectric barrier (e.g. glass, quartz, ceramic material or polymers) in between the electrodes. 78,80,86,93 The purpose of the dielectric barrier is to restrict the electric current and thus to prevent the formation of sparks and/or arcs. 93 A gas flow is applied between the (discharge) gap, which can typically vary from 0.1 mm (e.g. in plasma displays), to over 1 mm (e.g. for ozone generators) to several cm (e.g. in CO₂ lasers).^{78,86} Some typical planar (top) and cylindrical (bottom) DBD configurations are shown in Fig. 13. In general, DBDs operate at approximately atmospheric pressure (0.1-10 atm, but usually 1 atm), while an alternating voltage with an amplitude of 1-100 kV and a frequency of a few Hz to MHz is applied between both electrodes.

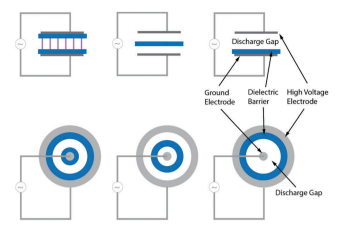


Fig. 13 Basic planar (top) and cylindrical (bottom) dielectric barrier discharge configurations.

To ignite the discharge, or more specifically, to transport current in the discharge gap, an electric field high enough to cause breakdown in the gas needs to be applied. This breakdown voltage (V_b) can be determined according to Paschen's Law, and is a function of the pressure (p) and the gap distance (d):

$$V_{b} = \frac{D \cdot pd}{\ln\left(\frac{C \cdot pd}{\ln(1/\gamma)}\right)}$$

where C, D and γ are the gas (or mixture) specific parameters. Upon breakdown of the gas, most often a non-uniform plasma, consisting of a large number of micro-discharges (or filaments), is observed (indicated as purple in the top-left panel of Fig. 13). This mode is called the 'filamentary mode', and plasma formation is restricted to these micro-discharges. The occurrence of a filamentary mode, as opposed to a homogeneous mode, depends on the type of gas. Most gases, including CO2, give rise to a filamentary mode. It is stated that the volume of the micro-discharges, and hence the plasma volume, comprises about 1-10% of the total gas volume. 94,95 The rest of the gas is not ionized and serves as background reservoir to absorb the energy dissipated in the micro-discharges and to collect and transport the long-lived species created in the micro-discharges.80 For a more detailed physical description on the formation of these micro-discharges, we refer to the review of Fridman et al.86

4.2.2. Microwave (MW) discharge. Microwave discharges operate according to a different principle and belong to the group of warm plasmas. They are electrode-less and as their name suggests, the electric power is applied as microwaves, i.e. electromagnetic radiation in the frequency range of 300 MHz to 10 GHz. There exist several different types of MW plasmas, e.g. cavity induced plasmas, free expanding atmospheric plasma torches, electron cyclotron resonance plasmas and surface-wave discharges. For more details about the different microwave discharges, we refer to several reviews in the literature. 21,78,93,96

In the so-called surface-wave discharge, which is most commonly used for CO2 conversion research, the gas flows through a quartz tube - which is transparent to microwave radiation - intersecting with a rectangular wave guide, where

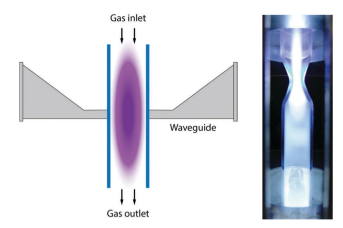


Fig. 14 Schematic (left) and image (right; courtesy of Dutch Institute For Fundamental Energy Research (DIFFER)) of a MW discharge.

the discharge is initiated, see Fig. 14. This system, also called 'guide-surfatron', involves surface waves. The microwaves propagate along the interface between the quartz tube and the plasma column. The wave energy is absorbed by the plasma. Characteristic features are the wavelength (815 MHz or 2.45 GHz), which is comparable to the length of the apparatus, and the short period of the exciting microwave field. Surface-wave discharges can be created both at reduced and at atmospheric pressure, but in the pressure regime above 0.1 atm they approach a state of LTE, 78

4.2.3. Gliding arc (GA) discharge. A gliding arc (GA) discharge is also a warm plasma, combining advantages of both thermal and non-thermal plasma systems.86 It is a transient type of arc discharge. At high currents, the periodic discharge typically evolves during one cycle from an arc to a strongly non-equilibrium discharge. The non-equilibrium GA is a very sophisticated physical phenomenon: this transitional quasi-equilibrium/non-equilibrium discharge is essentially non-uniform in time and in space, and includes an internal transition from thermal to non-thermal mechanisms of ionization.86

A classical GA plasma is an auto-oscillating periodic discharge, where the gas flows between two diverging flat electrodes. When applying a potential difference between both electrodes, an arc plasma is formed at the narrowest gap, which is then dragged by the gas flow towards a rising interelectrode distance, until it extinguishes and a new arc is ignited at the shortest interelectrode gap. More specifically, the length of the arc column increases together with the voltage until it exceeds its critical value (l_{crit}). At this point, heat losses from the plasma column begin to exceed the supplied energy and it is no longer possible to sustain the plasma in its LTE state, resulting in a fast transition into a non-LTE state. The discharge cools rapidly to gas temperature but the plasma conductivity is maintained by a high value of the electron temperature ($T_e \sim 1$ eV, which is most suitable for efficient vibrational excitation of CO₂). After this fast transition, the GA continues its evolution under non-LTE conditions, until the length reaches a new critical value $(l \sim 3 \cdot l_{\text{crit}})$, leading to decay of the discharge, after which the evolution repeats from the initial breakdown. During this cycle,

up to 75-80% of the energy can be dissipated in the non-LTE zone of the gliding arc. It is this effect that permits the stimulation of chemical reactions in regimes quite different from conventional thermal reaction chemistry. When the GA is operated under milder conditions, i.e. lower currents and high frequencies, this transition to the non-LTE phase can occur in the order of nanoseconds. As a result, the GA operates in the non-LTE regime starting almost immediately after its ignition, and a higher fraction of the discharge energy can be consumed by the non-LTE phase. 97–99

The classical GA plasma, however, also exhibits some disadvantages. The flat 2D electrode geometry makes it less compatible with industrial systems, and also the gas conversion is non-uniform and quite limited, because a considerable fraction of the gas does not pass through the active plasma region. Moreover, a high gas flow rate is needed to drag the arc, so the gas residence time is limited, thereby further limiting the conversion. Therefore, a 3D cylindrical GA plasma reactor was developed a few years ago, which makes use of vortex flow stabilization. 100 The gas flows in the reactor through a tangential inlet. An arc is again formed between both electrodes, and is dragged with the tangential gas flow, thereby expanding until it extinguishes, followed by a new cycle. Basically, the arc is again gliding between the anode and cathode, and it is stabilized in the centre of the reactor. Depending on the diameter of the anode tube, a forward or reverse vortex gas flow is created. If the anode diameter is equal to the cathode diameter, the gas can leave the reactor through the anode outlet, leading only to a forward vortex flow. On the other hand, if the anode diameter is smaller than the cathode diameter, the gas cannot immediately leave the reactor, and will first flow upwards close to the walls, in a forward vortex. As this gas is still cold, it creates an isolating and cooling effect, protecting the reactor walls from the warm plasma arc in the centre. When it reaches the upper end of the reactor, it will have lost some speed due to friction and inertia, and when flowing downwards, it moves in a smaller inner vortex, a so-called reverse vortex, where it mixes with the plasma arc, resulting in a more energy-efficient conversion. The reverse vortex flow GA is also called a 'gliding arc plasmatron' (GAP). Fig. 15 shows the difference between these two types of GAs.

4.2.4. Other plasma types used for CO₂ conversion. Besides the above three main discharge types for CO₂ conversion, some research is also being carried out with other plasma types, which are briefly discussed here. These include radio frequency (RF) discharges, several different atmospheric pressure glow discharges (APGD), corona discharges, spark discharges and nanosecond pulsed discharges.

RF discharges usually operate in the 1-100 MHz frequency range, resulting in a corresponding wavelength (300-3 m) much larger than the plasma reactor dimensions. The power coupling can be done through capacitive or inductive coupling, resulting in capacitively coupled plasma (CCP) and inductively coupled plasma (ICP). These kind of discharges are mainly applied to thin-film deposition, plasma etching, the sputtering of materials and as an ion source in mass spectrometry, 96 but some research is also being performed for CO2 conversion applications, as is Review Article Chem Soc Rev

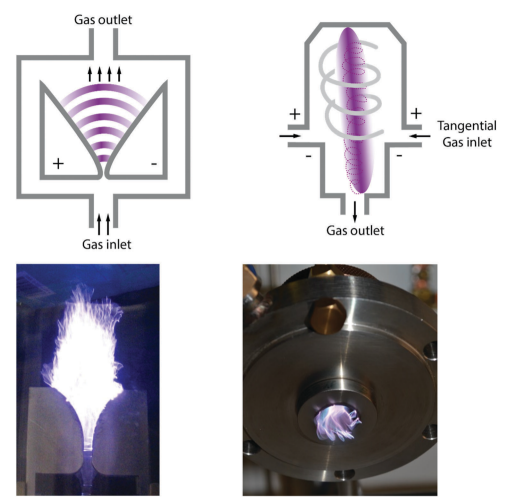


Fig. 15 Schematic of the classical GA (top left) and the GAP (top right) configuration, and pictures of the classical GA (bottom left; courtesy of the University of Manchester) and GAP (bottom right; courtesy of the University of Antwerp).

illustrated in Section 5. An RF source operating at low pressure, both with and without the influence of an external magnetic field, has been used. The advantages for CO₂ conversion are its capability to allow obtaining high electron densities at low gas temperature.

When a sufficiently large electric field is applied, corona discharges occur near sharp points, edges or thin wires when used as an electrode. When a high negative voltage is applied to the electrode, it acts as a cathode and is considered as a negative corona discharge. When a positive voltage is applied, it is considered as a positive corona discharge. Corona discharges are always non-uniform, with a strong electric field, ionization and luminosity close to the wire/sharp electrode, whereas the charged particles are dragged to the other electrode by weak electric fields. They are often operated in a pulsed mode, mostly to increase the power, while inhibiting the transition of the streamer formation into sparks.²¹ An advantage of corona discharges compared to DBDs is that they are relatively easy to establish; however, their performance in CO₂ conversion processes is quite similar, as will become clear from Section 5.

When a streamer is able to connect two electrodes, without the presence of a pulsed power supply (see corona discharge above) or the presence of a dielectric (see DBD), a spark discharge can develop by further growth of the current.²¹ However, when the amount of power provided is restricted, the discharge cannot develop into a stationary (thermal) arc and instead the discharge extinguishes within several hundred microseconds. Hence, spark discharges consist of an initiation of streamers that develop into highly energetic spark channels, which then extinguish and reignite periodically. Lightning is a typical example of a spark discharge.

A wide variety of plasma set-ups fall under APGDs, such as miniaturized direct current (dc) GDs, microhollow cathode dc discharges and RF discharges as well as DBDs. The main advantage of APGDs is the absence of vacuum conditions compared to with regular glow discharges, and also operating without elevated temperatures. Depending on the gas mixture and electrode configuration, the discharge can operate in a stable homogeneous glow or filamentary glow mode. For example, a DBD can operate in APGD mode, and in this case the discharge benefits from the average power densities of a DBD but operating in a uniform homogeneous glow mode, thus benefiting uniform gas treatment. This discharge is interesting for DRM due to its special characteristics of electron density (3 \times 10¹² cm⁻³), electron temperature (\sim 2 eV, suitable for the vibrational excitation of CO₂) and

plasma gas temperature (900 K), which is lower than that of thermal plasma but higher than that of DBDs and corona discharges.

Finally, nanosecond pulsed discharges are basically discharges that rely on repetitively pulsed excitation, through a nanosecond scale pulse rise time and duration. This leads to a highly non-LTE state with very high plasma densities for a relatively low power consumption due to the short pulse durations. Whereas (sub)microsecond pulsed discharges are initiated by Townsend discharges, nanosecond pulsed discharges ignite instantaneously, i.e. without the involvement of secondary electrons. Furthermore, the discharge remains in a glow-like (rather than filamentary) mode despite the high electric field. The interest in shorter pulses is not because of the duration of the discharge itself, but rather because they offer better control of the electron energy than continuous wave discharges, depending on the pulse length. 101 Thus, more energy can be directed towards the desired dissociation channels.

4.3. Plasma-catalysis

As with most of the technologies described in Section 3, plasma set-ups can also be combined with a packing material or catalyst, giving rise to plasma-catalysis. Plasma-catalysis is an emerging branch of plasma processing at the interface of a variety of disciplines, including physical chemistry, material science, nanotechnology, catalysis, plasma physics and plasma chemistry. In short, its objective is to enhance plasma reactions by adding a catalyst to the reaction cycle and vice versa. Theoretically speaking, combining plasma with catalysis offers the best of both worlds. Inert molecules are activated by the plasma under mild conditions, and subsequently the activated species selectively recombine at the catalyst surface to yield the desired products. This will be especially important to further advance and optimize the direct oxidative liquefaction pathway of plasmachemical CO2 conversion, in order to selectively produce the desired liquid products. In this section, we present a brief overview of the different approaches and possible interactions between the plasma and the catalyst. For more extensive details, we refer to several broad reviews on this specific topic that have recently been published. 83,102-104

4.3.1. Approaches. Plasmas and catalysts can be combined in two main configurations, as illustrated in Fig. 16. In the first, the so-called two-stage configuration (Fig. 16(b)), the catalyst is spatially separated from the plasma region, either upstream or downstream, but the main configuration is downstream. In the second, the so-called one-stage configuration (Fig. 16(c)), the catalyst is placed inside the discharge region. In a traditional thermal catalysis experiment, molecules are dissociatively adsorbed onto the catalyst with the energy being supplied in the form of heat. In plasma-assisted catalysis, species are activated by the plasma due to excitation, ionization or dissociation by electrons in the gas phase or on the catalyst surface.

The major difference between the one-stage and two-stage configuration is the kind of species to which the catalyst is exposed. In the two-stage configuration, the end-products and the long-lived intermediates will interact with the catalyst, while in the one-stage configuration, the catalyst can also interact with all the short-lived species, including excited

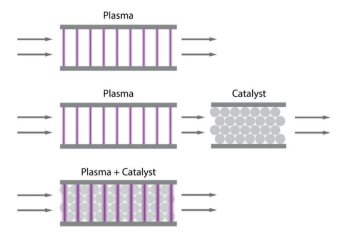


Fig. 16 Schematic of the different plasma-catalyst configurations: (a) plasma alone without a catalyst, (b) catalyst downstream of the discharge, and (c) catalyst directly inside the discharge zone. These examples mainly apply to a DBD. For MW and GA discharges, the catalyst is most commonly placed in the downstream region

species, radicals, photons and electrons. Furthermore, in this configuration the catalyst may also be influenced by the plasma and vice versa (see below). Besides plasma-catalysis, the preparation and modification of catalysts by plasma treatment is gaining increased attention, especially for catalysts with a low thermal stability. 103

The most widely investigated and, for this review, most interesting configuration is the one-stage configuration, in which the catalyst is placed inside the discharge region, either completely or only partially occupying the discharge zone. The catalyst itself can be introduced in the discharge as pellets, (fine) powders, foams, honeycomb monoliths, different electrode materials and electrode coatings or as coated quartz wool. The ease of adding a catalyst into the discharge zone greatly varies depending on the type of plasma reactor used. In general, due to their simple geometry and operation close to room temperature, implementation of a catalyst in a DBD reactor is very easy. Although MWs also have a simple geometry, due to the high gas temperature inside the discharge zone (1000-2000 K compared to 300-400 K for DBD) catalysts are often placed downstream, due to their low thermal stability. Finally, GA discharges have rather complex geometries and the same higher gas temperatures, thus catalysts are typically introduced downstream, although the use of a spouted bed has also been reported.105 If these MW and GA discharges can be operated at slightly lower temperatures (≤1000 K), this would yield new possibilities. It would also open up the way for using sufficiently thermally stable catalysts inside the discharge zone, and at the same time it would allow thermal activation of catalysts inside the discharge zone by the plasma.

4.3.2. Synergetic effects. The resulting interactions when combining a plasma with a catalyst for plasma-based CO2 conversion often yield improved process results in terms of conversion, selectivity, yield and energy efficiency. This surplus effect, as shown in Fig. 17, is a complex phenomenon originating from the interplay between the various plasma-catalyst Review Article Chem Soc Rev

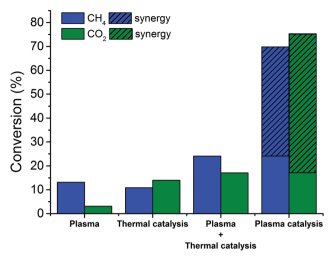


Fig. 17 Demonstration of the synergy of plasma-catalysis for the dry reforming of methane, adapted from Zhang $et\ al.^{106}$

interactions and is often termed 'synergy', when the combined effect is larger than the sum of the two separate effects. ¹⁰² Note that most research to date regarding this effect has been performed for DBDs.

The complex interactions can be subdivided into two categories: the effects of the plasma on the catalyst, and *vice versa*, the effects of the catalyst on the plasma. ^{102,107} Second, it is important to distinguish between the two types of effects: physical and chemical effects. While the physical effects are mainly responsible for gaining a better energy efficiency, the chemical effects can lead to an improved selectivity towards the targeted products. In the case of CO₂ splitting, mainly CO and O₂ are formed, so the primary added value of the catalyst is to increase the energy efficiency, although the conversion can also be improved by chemical effects, such as enhanced dissociative chemisorption due to the catalyst acid/basic sites. When adding a co-reactant (*e.g.* CH₄, H₂O, H₂), the catalyst allows modifying the selectivity towards the targeted products.

An overview of the plasma-catalyst interactions is given in Fig. 18. When adding a catalyst, the following effects on the plasma have been reported: (i) an enhancement of the electric field through the geometric distortion and surface roughness; (ii) the formation of micro-discharges inside the pores of the catalyst material, due to the very strong electric field inside the pores, leading to different characteristics compared to the bulk; (iii) a change in the discharge type, because the presence of insulating surfaces promotes the development of surface discharges; (iv) the adsorption of species on the catalyst surface, which affects their concentration and conversion due to longer retention times. All the effects of the catalyst on the plasma can thus generally be considered as physical effects. The reported effects of the plasma on the catalyst, on the other hand, are of both a physical and chemical nature. They include: (i) changes in its physico-chemical properties, such as a higher adsorption probability at the catalyst surface, a higher catalyst surface area, a changed catalyst oxidation state, the reduction of metal oxide catalysts to their metallic form, reduced coke formation on the surface and a change in the catalyst work function; (ii) the formation of hot spots on the surface due to the formation of strong micro-discharges; (iii) catalyst activation by photon irradiation emitted by the excited plasma species (although this effect is probably of only minor importance due to the low photon fluxes); (iv) lowered activation barriers due to there being more reactive vibrationally excited plasma species and consequently a greater possibility for non-adiabatic barrier crossings; (v) changes in the reaction pathways due to the presence of a wide variety of (re)active species.

While the physical effects have been more extensively studied, such as the plasma behaviour in packed-bed DBD reactors¹⁰⁸ or inside catalyst pores, ¹⁰⁹ the chemical effects are less understood, mainly because they are often correlated with the physical effects. Indeed, in a DBD, the catalyst is mostly introduced as a packed bed, making it difficult to distinguish between the two effects. Hence, more systematic studies on both the physical and chemical effects of the catalyst material are highly needed.¹⁰² Furthermore, as summarized in Fig. 19, the plasma and catalyst both have an influence on the gas composition, and at the same

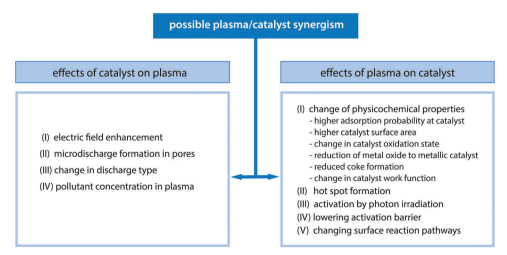


Fig. 18 An overview of the possible effects of the catalyst on the plasma and vice versa, possibly leading to synergism in plasma-catalysis.

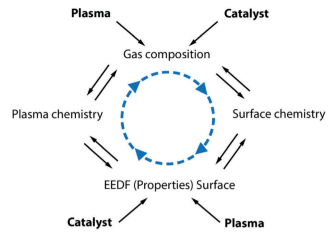


Fig. 19 Visualization of the continuous influence of plasma and the catalyst on the gas conversion process (EEDF = electron energy distribution function).

time the plasma influences the catalyst properties and *vice versa*, as mentioned above. Therefore, both the gas composition and the plasma and catalyst properties continuously influence each other, and at the same time are influenced by the plasma chemistry and surface chemistry. Thus, it is recommended to develop tailored catalysts for plasma-catalysis, rather than simply relying on classical catalysts. This is discussed further in Section 5.

Finally, it is worth mentioning that besides CO_2 conversion, the combination of plasma with catalysts is also promising for nanomaterial synthesis, such as for the growth of carbon nanotubes, nanofibres, and graphene. To date, research on this topic has been mainly focused on pure methane plasmas, instead of CO_2 plasmas. This is, however, outside the scope of the present review, but we refer to another recent review paper, where these applications are discussed in detail.

4.4. Definitions

In this section, an overview of the different expressions for the specific energy input (SEI), conversion (χ), energy efficiency (η) and energy cost (EC) are given, which are used for the critical assessment in Section 5.

4.4.1. Specific energy input (SEI). The specific energy input (expressed in J cm $^{-3}$ or kJ L $^{-1}$) is defined as the plasma power divided by the gas flow rate, and this is the dominant determining factor for the conversion and energy efficiency in a plasma process, as will also become clear in Section 5:

SEI (J cm⁻³) = SEI (kJ L⁻¹)
=
$$\frac{\text{Power (kW)}}{\text{Flowrate (L min}^{-1})} \times 60 \text{ (s min}^{-1})$$
 (1)

The SEI is also commonly expressed in electron volts per molecule:

$$\begin{split} \text{SEI (eV per molecule)} &= \text{SEI (kJ L}^{-1}) \\ &\times \frac{6.24 \times 10^{21} \left(\text{eV kJ}^{-1}\right) \times 24.5 \left(\text{L mol}^{-1}\right)}{6.022 \times 10^{23} \left(\text{molecule mol}^{-1}\right)} \end{split}$$

Note that the value of 24.5 L mol^{-1} is only valid for 298 K and 1 atm.

4.4.2. Absolute and effective conversion. Two types of conversion can be defined. The (absolute) conversion is based on the molar flow rates of the reactants, *e.g.* CO₂, CH₄, H₂O or H₂. This is the typical expression used for 'conversion' throughout this review, and also in general in the literature:

$$\chi_{\text{abs,reactant}_i} = \frac{\dot{n}_{\text{reactant}_i, \text{inlet}} - \dot{n}_{\text{reactant}_i, \text{outlet}}}{\dot{n}_{\text{reactant}_i, \text{inlet}}} \tag{3}$$

where \dot{n} stands for the molar flow rate of reactant species i.

When more than one gas is present in the feed mixture, the effective conversion takes the dilution into account:

$$\chi_{\text{eff,reactant}_i} = \chi_{\text{abs,reactant}_i} \times \frac{\dot{n}_{\text{reactant}_i, \text{inlet}}}{\sum_i \dot{n}_{\text{reactant}_i, \text{inlet}}}$$
(4)

This alternative definition of conversion is important for comparing the conversion of a specific reactant in different mixtures, since it shows how the conversion rate of the reactant is affected, rather than the absolute value of its conversion.

For an easy comparison of DRM mixtures, it is also interesting to determine the total conversion, which is the sum of the effective conversions:

$$\chi_{\text{Total}} = \sum_{i} \left(\frac{\dot{n}_{\text{reactant}_{i}, \text{inlet}}}{\sum_{i} \dot{n}_{\text{reactant}_{i}, \text{inlet}}} \times \chi_{\text{abs, reactant}_{i}} \right) = \sum_{i} \chi_{\text{eff, reactant}_{i}}$$
(5)

4.4.3. Energy efficiency and energy cost. The energy efficiency and energy cost depend on the process under study. The energy efficiency is a measure of how efficiently the process performs compared to the standard reaction enthalpy, based on the specific energy input (SEI):

$$\eta = \frac{\chi_{\text{Total}} \times \Delta H_{298 \text{ K}}^{\circ} \left(\text{kJ mol}^{-1} \right)}{\text{SEI (kJ mol}^{-1})} = \frac{\chi_{\text{Total}} \times \Delta H_{298 \text{ K}}^{\circ} \left(\text{eV molecule}^{-1} \right)}{\text{SEI (eV molecule}^{-1})}$$
(6)

As mentioned in Section 2, $\Delta H_{298\text{K}}^{\circ}$ is 283 kJ mol⁻¹ (or 2.93 eV per molecule) for pure CO₂ splitting and 247 kJ mol⁻¹ (or 2.56 eV per molecule) for DRM.

The energy cost is the amount of energy consumed by the process (generally expressed as kJ per converted mol or eV per converted molecule):

$$EC (kJ \text{ mol}_{conv}^{-1}) = \frac{SEI (kJ L^{-1}) \times 24.5 (L \text{ mol}^{-1})}{\chi_{Total}}$$
 (7)

Note that the value of 24.5 L mol^{-1} is only valid for 298 K and 1 atm.

$$\begin{split} EC \left(eV \; molecule_{conv}^{-1} \right) &= EC \; \left(kJ \; mol_{conv}^{-1} \right) \\ &\times \frac{6.24 \times 10^{21} \; \left(eV \; kJ^{-1} \right)}{6.022 \times 10^{23} \; \left(molecule \; mol^{-1} \right)} \end{split} \tag{8}$$

5. Critical assessment of plasma-based CO₂ conversion

In this section, we address the present state-of-the-art with respect to plasma-based CO2 conversion, focusing on a critical assessment of the advantages and disadvantages of the various set-ups described in Section 4.2. The aim is to unveil their future challenges, risks and opportunities for successful implementation. Several options are being investigated, including both pure CO₂ splitting into CO and O₂, as well as the reaction with other gases, such as CH₄ (the dry reforming of methane), H₂O (artificial photosynthesis) and H₂ (hydrogenation). By adding a hydrogen source, the main aim is to produce valueadded chemicals and/or fuels, e.g. syngas, hydrocarbons and valuable oxygenates, such as methanol, formaldehyde and formic acid. As mentioned above in Section 4, most research on plasma-based CO₂ conversion is performed with DBDs, MW and GA plasmas, with a main focus on improving the energy efficiency of the conversion, as well as the selectivity towards value-added chemicals, in combination with catalysis.

In order for plasma technology to be competitive with traditional as well as emerging novel technologies (as described in Sections 2 and 3) we can define two main goals. First, plasma technology will have to be competitive with electrolysis, which is its main competitor and that also has the same advantage as being able to rely on all sorts of renewable electricity. For electrochemical water splitting, commercial energy efficiencies of 65-75% have been obtained; hence, the goal of plasma technology should be at least to aim for an energy efficiency comparable to this, and better than the thermal equilibrium energy efficiency limits of $\sim 45-50\%$ (see Section 2). Second, when comparing with other novel technologies that can make use of direct solar energy, such as solar thermochemical conversion, we should look at the solar-to-fuel conversion efficiency. As such, for novel technologies, a solar-to-fuel conversion efficiency of approximately 20% is considered industrially competitive, e.g. when looking at the production of syngas, 110 as will be elaborated in Section 6.3 below. When relying on solar energy, taking a solar panel efficiency of 25%111 and a plasma-conversion energy efficiency of 60-80% towards the syngas components CO and H2, this would yield a competitive solar-to-fuel efficiency of 15-20%. Hence, in the following critical assessment of plasma-based CO₂ conversion technology, we should keep in mind that - from an energy efficiency point of view - to be considered as a competitive and worthy alternative, plasma technology should aspire to an energy efficiency of at least 60%, at least when focusing on the production of syngas.

On the other hand, the conversion of syngas into more suitable fuels and chemicals through the Fischer-Tropsch process or for methanol synthesis (and subsequently for methanol or ethanol to olefin) is a very energy-intensive process. As a result, the energy efficiency requirements for the direct production of these compounds is highly dependent on the formed products. Hence, the target energy cost to be competitive can be significantly lower for a direct one-step process. For example, a solar-to-methanol conversion efficiency of 7.1% is already economically

feasible. 110 In this case, a plasma-conversion energy efficiency of 30% (instead of the above 60%) would already suffice. This is very important and it shows that, in contrast to the indirect approach through syngas, for the direct oxidative pathway - which aims to synthesize oxygenated liquid products in one step - the energy efficiency target is much lower, depending on the products formed, due to the circumvention of the energy-intensive step of converting syngas into the desired liquids. The possibility to proceed through this direct oxidative pathway is one of the key benefits of plasma technology, and in theory this pathway is the most promising. For now, however, a lot of research is still needed to understand the underlying processes, in order to improve the yield of the desired (oxygenated) liquid products. 112 Hence, the analysis in this section is based on the production of the syngas components CO and H2 using plasma technology, based on the energy efficiency target of 60%. Nevertheless, it is important to keep in mind that this 60% energy efficiency target applies to syngas production, but this target would change completely, i.e. it would decrease by a factor two to three, if a direct oxidative pathway towards liquids could be successfully realized using plasma technology.

5.1. CO_2 splitting

As outlined in Section 4 above, in the case of pure CO_2 splitting, the dominant products are CO and O_2 in a 2-to-1 ratio. Hence, this is a simple chemical process and there are no concerns regarding the complexity and the wide variety of products that can be formed – such as in the case with an added H-source – and thus one does not have to contemplate how to steer the different selectivities. As such, the research can solely focus on optimizing the CO_2 conversion and the energy efficiencies, and will be judged on those indicators.

5.1.1. DBD plasmas. Although many papers are already published on CO₂ conversion by DBD, detailed systematic studies presenting values for both the conversion and energy efficiency in a DBD appear to be very scarce. The most detailed studies focusing on a wide range of conditions were performed by Aerts *et al.*,⁹⁵ Paulussen *et al.*,¹¹³ Yu *et al.*¹¹⁴ and Ozkan *et al.*¹¹⁵ We combine the most important observations from these works and complement them with various other findings in the literature, to sketch a complete image with the data available to date.

The most commonly used geometry to study CO2 conversion uses a coaxial DBD reactor, 95,113-115 while parallel plate reactors 116 (see Section 4.2.1; Fig. 13) are mainly used to study the system with advanced optical diagnostics. 117 To achieve higher values of CO2 conversion and energy efficiency, several approaches have already been investigated, including changing the applied frequency, applied power, gas flow rate, discharge length, discharge gap, reactor temperature, dielectric material, electrode material, mixing with gases, i.e. Ar, He, N2, and by introducing (catalytic) packing materials. Furthermore, extensive modelling has also been performed to obtain a more fundamental insight into the plasma chemistry in a DBD and in turn to aid in the improvement of future experiments. Fig. 20 summarizes most of the data available in the literature. From this figure, several main trends become clear. First of all, the conversion increases with increasing specific energy input (SEI), while the energy efficiency generally decreases with

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a) 40 Conversion (%) n Aerts [95] b) Paulussen [113] 0 PB Van Laer [137] PB Butterworth [136] Yu [114] 20 Ozkan [115] PB Mei [134] PB Duan [135] Brehmer [117] Energy efficiency (%) Ramakers [123] PB Michielsen [133] Belov [119] Wang [122] Wang [118] 0 0.01 0.1 10 SEI (eV/molecule) c) 20 Energy efficiency (%)

Fig. 20 Experimental data collected from the literature for CO_2 splitting in a DBD, showing the conversion (a) and energy efficiency (b) as a function of the SEI, as well as the energy efficiency as a function of the conversion (c). The open symbols represent the data with a packed bed (PB). Note, some of the data have been recalculated from the original references to take, among others, dilution effects into account.

20

Conversion (%)

10

30

40

increasing SEI, especially above an SEI of 10 eV per molecule. When plotting the energy efficiency as a function of the conversion, it is clear that most of the results are situated below an energy efficiency of 15% and a conversion of 40%, with some exceptions. The highest conversions of 42% were obtained for packed-bed DBDs while the highest energy efficiency of 23% was obtained with a pulsed power DBD. It is important to note that some of the data in

this figure has been recalculated to represent coherent values for the conversions and energy efficiencies. For instance, in plasma research, the mixture is sometimes diluted with He, Ar or N_2 to obtain an more easily ignitable and more stable discharge, but as will be mentioned below, this influences the results – an effect that was, among others, not always taken into account in the original data. A more detailed influence of the different parameters is discussed below.

It is well known that the specific energy input (SEI), which is defined as the plasma power divided by the gas flow rate, is the dominant determining factor for the conversion and energy efficiency. 95,114,115,118 It has also been observed that the same values of SEI defined by different combinations of plasma power and gas flow rate can result in different conversions. It is the gas flow rate, and hence the residence time, which has the most important effect, while the effect of the power is less significant. As such, a lower power with a lower gas flow rate will result in a higher conversion and energy efficiency than a higher power with a higher gas flow rate for the same SEI. 95,115

Some possible geometry modifications are the discharge length and discharge gap. Yu et al. 114 found that varying the discharge length for a fixed SEI has no significant effect on the discharge characteristics, the conversion and the energy efficiency. On the other hand, the discharge gap appears to affect the discharge behaviour, and thus also the $\rm CO_2$ conversion. Aerts et al. 95 found that above a certain gap width (3.3 mm in their case) less streamer formation occurs, leading to a decrease in the effective plasma volume, and hence lower conversions and efficiencies result.

Furthermore, Aerts *et al.*⁹⁵ also reported that the applied frequency seems to have a negligible effect on the conversion and energy efficiency. However, the plasma appears more filamentary at high frequency (75 kHz) compared to at low frequency (6 kHz). Ozkan *et al.*, 115 however, found that both the conversion and energy efficiency decrease slightly upon increasing the frequency from 15 to 30 kHz. This was explained by Paulussen *et al.*, 113 who suggested that the optimum discharge frequency depends on the power input, and as such it cannot be unambiguously stated that higher or lower frequencies give rise to an increased CO_2 conversion and energy efficiency.

In thermochemical reactions, the gas temperature is one of the most important parameters governing the reaction rates. For the plasma-based conversion in a DBD, however, the effect of temperature is not so clear. Paulussen $et\ al.^{113}$ observed a slight and linear increase in the conversion from 26% to 28.5% when the inlet gas was heated from 303 K to 443 K. Ozkan $et\ al.^{115}$ on the other hand, reported that the wall and gas temperature should remain as low as possible. In addition, Wang $et\ al.^{118}$ detected an increase in conversion of 0.5–3% when using an external fan to cool the reactor, and they suggested that higher flow rates are preferred, since the latter remove large amounts of heat from the reactor. Brehmer $et\ al.^{117}$ noted a big effect of the wall temperature on the O_3 density and suggested that the recombination of CO and O on the wall would also increase with rising wall temperatures.

The effect of the dielectric material used in the DBD reactor is another topic of debate. By studying the formation of a

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conductive coating, Belov et al. 119 observed that the conductivity of the dielectric might be the most crucial parameter affecting the discharge properties. However, Aerts et al.95 reported no significant effect between quartz or alumina under a wide range of conditions, although it was still concluded that alumina has several advantages in terms of reactor stability. Ozkan et al. 115 found that alumina and quartz perform better than mullite and Pyrex, and that a thicker dielectric leads to a higher conversion and energy efficiency. 115 More sophisticated dielectric materials were created and tested by Li et al. 120,121 and Wang et al. 122 The former investigated the influence of Ca_{0.8}Sr_{0.2}TiO₃ (CST) with 0.5 wt% Li₂Si₂O₅ as the dielectric barrier and achieved an improvement in the conversion by a factor 9 of up to 9, compared to silica glass. Wang et al. 122 investigated the performance of CST ceramics with various amounts of CaO-B2O3-SiO2 (CBS) glass addition (0.5-5 wt%), and found that the addition of 5.0 wt% CBS resulted in an increase in the conversion by a factor of 2.6, while the energy efficiency almost tripled compared to that with 0.5 wt% CBS. However, from the calculation of the plasma power in both studies, it appears that the more sophisticated dielectrics mainly increased the efficiency between the input power and plasma power, and not the effective plasma-conversion energy efficiency. Nevertheless, these dielectrics allowed operation under lower voltages, which could be beneficial for certain processes. In summary, different dielectrics may allow for easier igniting and streamer formation, but not necessarily more energy-efficient plasma-conversion chemistry.

Besides the dielectric material, also the electrode material can be varied. Wang $et~al.^{118}$ studied the effect of changing – or coating – the high voltage electrode and obtained an order of activity as Cu > Au > Rh > Fe ~ Pt ~ Pd. The Cu and Au electrodes yielded a relative increase in the conversion by a factor 1.5 compared to an Fe electrode. Furthermore, the maximum energy efficiency of the Au electrode was almost three times higher than the energy efficiency of the Rh electrode under the same conditions. However, besides the fact that some of these electrodes or coatings are more expensive, compared to the inert stainless steel electrode, they also are susceptible to chemical erosion (i.e. oxidation) and plasma sputtering (as observed for Au).

As mentioned above, some researchers have also added inert gases, such as N2, Ar and He, to ignite the plasma more easily. This also has several effects on the discharge characteristics, conversion, energy efficiency and even by-product formation in the case of N2. The addition of He and Ar leads to an increase in the CO₂ conversion, but the effective conversion decreases, since there is less CO₂ present in the mixture and the increased conversion is not sufficient to counteract this drop in the CO₂ fraction. As a result, the energy efficiency decreases as well. 118,123,124 The addition of N2, on the other hand, shows a completely different behaviour. Snoeckx et al. 125 discovered that the presence of N₂ in the gas mixture of up to 50% barely influences the effective CO₂ conversion and the corresponding energy efficiency. Indeed, N2 enhances the absolute CO2 conversion, due to the dissociation of CO2 upon collision with N2 metastable molecules, and this effect is strong enough to

compensate for the lower CO_2 content in the mixture. However, the presence of N_2 in the mixture leads to the formation of unwanted by-products, *i.e.* N_2O and several NO_x compounds, with concentrations in the range of several 100 ppm, ¹²⁵ which can give rise to severe air pollution problems.

Other modifications being investigated to improve the CO₂ conversion in a DBD reactor are the use of micro plasma reactors, ¹²⁶ a hybrid DBD reactor on the surface of an SOEC¹²⁷ and a pulsed power supply. ¹¹⁵ A micro plasma reactor provides a stronger electric field and a higher concentration of reactive species, while offering better control of the processing parameters. ¹²⁶ Applying DBD plasma on the surface of an SOEC allows the *in situ* exclusion of O₂ during CO₂ splitting, resulting in an increase in the conversion by a factor of 4. ¹²⁷ Finally, the use of a power supply in the pulsed or so-called burst mode, instead of injecting the power in a continuous AC mode, is reported to lead to an increase in conversion and energy efficiency by a factor of 1.5 for a duty cycle of 50%. ¹¹⁵

Besides all the experimental work, great advances have also been made in modelling the plasma chemistry for CO₂ conversion in a DBD. 90,95,124,128-131 The main findings are that the splitting of CO2 is dominated by electron-impact reactions with ground-state molecules and, predominantly, by electron-impact excitation followed by dissociation. Electron-impact ionization is also important, but is compensated by the fact that a large fraction of the formed ions will eventually recombine, resulting in the formation of CO2. Splitting from the vibrationally excited states is found to be of minor importance in a DBD. 95,128 A reduced chemistry model consisting of only 9 species and 17 reactions was presented by Aerts et al.,95 which allowed identifying the main dissociation mechanisms. A 1D fluid model, with roughly the same chemistry, was developed by Ponduri et al. 130 Furthermore, recently, a thorough examination of the cross-sectional data was performed, 94,132 as well as a careful examination of all the rate coefficients and a comparison of the performances of different models. 129

Finally, as mentioned in Section 4.3, a packing can be added to the DBD reactor to enhance the conversion and energy efficiency. In the case of pure CO₂ splitting, the addition of a packing will not influence the formation of products, since no hydrogen source is available. Hence, most of research work focuses on increasing the conversion and energy efficiency by physical effects. Generally, when tested, the CO₂ conversion in packed-bed DBD reactors is always higher than in the corresponding empty reactors, albeit Fig. 20 clearly illustrates that the highest efficiencies found to date are for regular DBD reactors, achieved by varying the different parameters discussed above. Nevertheless, at high conversions, we see that the packed-bed reactors are generally more efficient. As such, in general, adding a packing seems to allow the system to operate at the same energy efficiency, but significantly increases the conversion. Several materials have already been investigated, more specifically, glass wool, 133 glass beads, 134 silica gel, 114 quartz, 114 quartz wool, 133,135 quartz sand, 135 Al₂O₃, 114,133,135,136 CaTiO₃, 114 ZrO₂, 133,137 SiO₂, 133 BaTiO₃, ^{133,134,136} MgO¹³⁵ and CaO. ¹³⁵ The best results have been obtained for ZrO2137 and CaO,135 with conversions in the range of

30-45% and energy efficiencies in the range of 5-10% (see Fig. 20). One of the more recent and interesting works is from Butterworth et al. 136 The authors stated that the testing of different materials should be performed according to a more predefined protocol, since the packing particle size affects the discharge phenomena and the chemistry within the packed bed. More specifically, the efficacy of CO2 conversion is strongly affected by the particle size, whereby small particle sizes (180-300 µm) can increase the CO₂ conversion by up to 70%. However, they also increase the reactor breakdown voltage and they lead to partial discharging, i.e. a drop in the fraction of the reactor where plasma formation occurs. Comparison with the works of other researchers shows that quite often, insufficient electric field strengths are applied for complete reactor discharging to occur. Hence, packing materials for plasma-catalysis should be tested with equivalent reactor operating conditions. It is therefore important to ensure that either: (a) complete discharging occurs in the reactor, or (b) the partial reactor discharging is quantified. 136 A similar message, regarding the need to compare material performances with similar reactor set-ups, was given in the recent work of Michielsen et al. 133

From all these data in the literature, we can conclude that a DBD reactor can provide reasonable conversions of up to 40%, but the energy efficiency is still at least a factor of 3-4 away from the necessary 60% mark. DBDs have the advantage of being very scalable and easy to operate, but their current energy efficiency makes it doubtful that they will be the most suitable technology for pure CO2 splitting.

5.1.2. MW and RF plasmas. The dissociation of CO₂ using MW and RF discharges, was already extensively studied both theoretically and experimentally in the 1970-1980s,²¹ but gained renewed interest with the current global challenges regarding CO₂ emissions. Already back in the 1970-1980s, it was concluded that MW discharges were ideal for obtaining high energy efficiencies for CO2 conversion, due to a combination of their relatively high electron density and low reduced electric field (see Section 4). These conditions favour the excitation of the asymmetric mode vibrational levels of CO2. 138 This efficient dissociation channel is a combination of several reaction steps excitation of the lower vibrational levels by electrons, followed by collision between vibrational levels, gradually populating the higher levels, and dissociation of the excited vibrational levels stimulated by collisions with other molecules - covering a whole spectrum of vibrational levels.87

The main set-ups used for CO2 conversion are the (surfaguide) MW discharge (2.45 GHz and 915 GHz) and RF discharges (13.56 MHz). More details about these set-ups can be found in Section 4. To obtain the best values for the conversion and energy efficiency, several approaches have already been proposed, including changing the applied power, gas flow rate, flow type, reactor geometry, gas temperature, and admixture gases, or by introducing (catalytic) packing materials, as well as by carrying out extensive plasma chemistry modelling to gain a better insight into the underlying mechanisms. Fig. 21 summarizes all the data available from the literature. The highest energy efficiencies reported in the 1970-1980s were up to 80% for subsonic flow conditions, and up to 90% for supersonic

flows. 21,139,140 Recently, several attempts have been made to reproduce these high energy efficiencies. 141-144 To date, maximum energy efficiencies of 50% have been successfully achieved, with both MW and RF discharges. 141-143 The results obtained by Bongers et al. 142 were near the thermal dissociation limit, indicating that the thermal dissociation mechanism is predominant. The authors concluded that the applied values for a reduced electric field, i.e. 70-80 Td, were too high, and future experiments should be directed towards values around 20-50 Td, while keeping the temperature in the discharge as low as possible. It should be noted that all the experiments reporting such high energy efficiencies were performed at reduced pressures, which might not be beneficial for highthroughput industrial implementation, despite the high flow rates of up to 75 SLPM. 142 Indeed, increasing the pressure leads to a significant decrease in energy efficiency. 140 At atmospheric pressure, Spencer et al. 144 obtained a maximum energy efficiency of 20%, which is nevertheless still a factor two better than that obtained for a DBD (see previous section).

Fig. 21 also demonstrates that, in general, surfaguide MWs achieve the highest energy efficiencies (up to 90%). Furthermore, just like for the DBD results, the conversion increases with increasing the SEI, while the energy efficiency decreases above an SEI of 0.1-1 eV per molecule. When plotting the energy efficiency as a function of the conversion, it is clear that the more recent results typically have an energy efficiency of 10-50% in the entire range of conversions up to 95%. Again, like for the DBD results above, it is important to note that some of the data have been recalculated to represent coherent values for the conversions and energy efficiencies. A more detailed influence of the different parameters is discussed below.

For the MW discharges, the same trade-off as described for the DBD between energy efficiency and conversion as a function of SEI is present. 141,142,144-146 Hence, to successfully increase the energy efficiency, one must be able to increase the conversion degree without increasing the specific energy input of the system, which essentially requires using techniques other than increasing the input power. 145 It is also clear that the highest energy efficiencies are obtained for low SEI values (0.1-1 eV per molecule), while the conversion is low at these conditions, hence conflicting requirements are encountered. 143

As already mentioned, pressure has one of the most important influences on the MW discharge and on its performance for CO₂ conversion, and most studies are carried out at reduced pressure. One of the reasons for this is that at low pressure the three-body recombination reaction $CO + O + M \rightarrow CO_2 + M$, which is the most important reaction limiting the effective CO2 conversion in the gas phase, becomes negligible. 146 An optimum operation pressure seems to exist around 150 mbar (at 5 SLPM), but the latter depends on other operating conditions, such as the gas flow rate. 141 Furthermore, the pressure plays an important role in which mode - contracted, diffuse or combined - the plasma is operating.141

Several more complex flow types and geometries have already been studied to optimize the MW discharge performance for CO2 conversion. Supersonic flows in MW discharges Review Article Chem Soc Rev

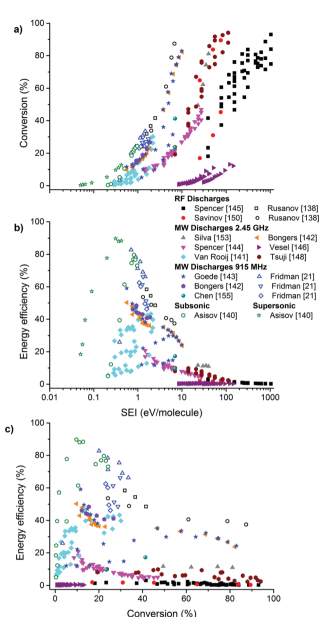


Fig. 21 Experimental data collected from the literature for CO_2 splitting in MW and RF discharges, showing the conversion (a) and energy efficiency (b) as a function of the SEI, as well as the energy efficiency as a function of the conversion (c). The open symbols represent the data from the 1970–1980s. Note, some of the data have been recalculated from the original references to take, among others, dilution effects into account.

have proven to reduce the losses of vibrational levels upon collision with ground-state molecules (*i.e.* the so-called VT relaxation), which occurs mainly at higher temperature, and has allowed achieving an energy efficiency of up to 90%, 140 in agreement with theoretical calculations. 139 Bongers *et al.* 142 observed an energy efficiency of only 15% for a supersonic flow, but by additionally quenching the plasma, an energy efficiency of up to 47% was observed, as predicted for plasmas in the thermal regime (see Section 2, Fig. 2). Similarly, the addition of a vortex gas flow, more specifically a reverse vortex, led to a significant improvement in CO₂ conversion and energy efficiency compared to a forward vortex flow. 142

For MW discharges, the effect of gas temperature is quite complicated. A signature of the desired non-equilibrium conditions in a MW plasma is a low (or moderate) gas temperature (in the order of 1000-2000 K), while vibrational and electron temperatures are higher (i.e. 3000 up to 8000 K and 1-5 eV, respectively). 89,141 For low pressures and reduced electric fields (E/n), gas temperatures around 2000 K have been observed. However, when increasing pressure at an SEI of about 0.5 eV per molecule, the discharge undergoes a transition from a diffuse to a contracted regime, and the temperature rises steeply to 14 000 K.141 When this transition occurs and the vibrational and rotational gas temperature are in thermal equilibrium, low values of energy efficiency are observed. 144 Recent modelling studies suggest that keeping the temperature under control is beneficial for the vibrational excitation, and is a key parameter for more efficient CO₂ conversion. 88-90 Bongers et al. 142 noted that the E/n in their MW plasma set-up was too high to achieve VT non-equilibrium conditions and that the high temperatures in the plasma core could account for the observed high values of E/n. It is thus clear that the gas temperature should be kept as low as possible to reduce vibrational energy losses via VT relaxation.88

Just like for the DBD plasma, some papers have also reported on the addition of other gases to CO2 in the MW plasma, i.e. Ar, 144,147,148 He 148 and N2. 149 For Ar, Spencer et al. 147 observed no effect on CO production, suggesting it does not affect the collisional processes benefiting dissociation, but, depending on the set-up, it might be necessary as an electron source to ignite the discharge. 144 Tsuji et al. 148 observed higher CO₂ conversions when diluted in Ar compared to He, and furthermore, the conversion increased but the energy efficiency decreased when adding more Ar or He. In contrast, just like for the DBD, N₂ can play a very important role in the CO₂ conversion, but the mechanism in the MW plasma is different than in the DBD plasma. Indeed, while in the DBD, the metastable electronically excited N2 molecules give rise to enhanced CO2 conversion (see the previous section),125 while in the MW plasma, the improvement is due to the vibrationally excited N2 molecules.149 The energy difference between the first vibrational level of N2 and CO2 is very small, making the fast resonance transfer of vibrational energy from N2 to CO2 possible. 150 As such, N₂ can help with the vibrational pumping of the asymmetric mode of CO2 and thus can enhance the CO2 conversion. On the other hand, the vibrationally excited N2 molecules can also react with O atoms, leading to the production of NO_x in undesirable concentrations, as was also observed for a DBD (see previous section). 149

Just like for DBDs, big leaps forward have been made in the past few years regarding modelling the plasma chemistry to better understand and improve CO₂ conversion in MW discharges. ^{87–90,94,131,151,152} The added complexity for MW plasmas compared to the plasma chemistry in DBDs stems from the very effective excitation to vibrational states, which can lead to the so-called ladder-climbing effect and eventually very efficient dissociation of the vibrationally excited CO₂ molecule (as discussed in Section 4.1.4). This means that all the vibrational levels up to the dissociation limit need to be taken into account in accurate models for a CO₂ MW plasma, as well as all the

reactions with these different vibrationally excited levels. The models predict that, besides electron-impact dissociation of the vibrationally excited levels, also collisions with neutrals will become important as dissociation mechanisms, as was also suggested by one optical characterization study, 153 and this is the key to achieving maximum energy efficiency of the CO₂ splitting process. 87 It has also been shown that at low E/nvalues, ¹⁵¹ and in post-discharge conditions, ¹⁵² the rates from the vibrational-dissociation mechanism overcome the electronimpact dissociation. To be able to perform multidimensional modelling investigations for these type of discharges, i.e. where the vibrational levels play an important role, a reduced chemistry set has been developed in which, among others, the vibrational levels are lumped into a limited number of groups, to avoid the need to solve equations for all the individual CO₂ vibrational levels.154

Finally, in contrast to the DBD research, no work has been performed on adding a (catalyst) packing in the MW discharge zone, and only a few papers have reported on adding a postdischarge (catalytic) packing. 144,155 Of course, as mentioned in Section 4.3, this is due to the added complexity of adding packing materials to a MW discharge. Spencer et al. 144 added a monolith structure with and without a Rh/TiO2 catalytic coating in the post-discharge zone. For both cases, a slight energy efficiency loss was observed and the Rh/TiO2 was deemed inappropriate due to the possible stimulation of the backward reaction of CO + $O_2 \rightarrow CO_2 + O$. On the other hand, Chen et al. 155 reported that the use of plasma-pretreated-TiO₂supported NiO catalysts in the post-discharge zone led to an energy efficiency of 17%, which was an increase by a factor 2 compared to the plasma-only case. However, this increase was suggested to arise from the dissociation of CO₂ at the catalyst surface with oxygen vacancies through dissociative electron attachment, which is an inherently less efficient dissociation process than the step-wise vibrational excitation (as shown in Section 4.1.4; Fig. 12). We believe that the addition of a catalyst to a MW discharge for pure CO2 splitting will only be beneficial if it is capable of either effectively lowering the E/n value of the discharge due to its physical effects and/or stimulating the dissociation of vibrationally excited CO2 molecules on the surface. Furthermore, the development of MW discharges operating at lower temperatures (≤1000 K) would allow the implementation of thermal catalysts in the discharge zone, rather than use of the current post-discharge packing.

From all these data, it should be clear that MW discharges are more than capable of surpassing the 60% efficiency mark for pure CO2 splitting. However, the best results of this earlier work^{21,139,140} have not yet been reproduced to date, while the best energy efficiency in the more recent work appears to be around the thermodynamic equilibrium value of 45-50%. This observation, together with the reported gas temperatures, makes it questionable whether vibrational excitation plays the major role here. Nonetheless, from modelling insights, 88,89 it is evident that when the set-up can be tailored to achieve the correct strong non-equilibrium conditions, 141 vibrational excitations can lead to energy efficiencies of up to 90%. The main disadvantage

of MW discharges is, however, their current requirement to operate at low pressures, in order to reach this strong nonequilibrium, and thus these high energy efficiencies.

5.1.3. GA plasmas. It is evident that both DBD and MW discharges have their distinctive advantages and disadvantages. DBDs operate at atmospheric pressure, but cannot make use of the most energy-efficient dissociation process by vibrational interactions, while MW discharges can, but the vibrational pathways are only fully exploited when they operate at reduced pressure. The GA set-up tries to combine the best of both worlds, offering the possibility to operate at atmospheric pressure and at the same time trying to reach a strong enough nonequilibrium to stimulate the most efficient dissociation process through vibrational excitation. 100

As explained in Section 4.2.3, there are two main GA reactor geometries used for CO2 conversion. The first one relies on simple two-dimensional (2D) electrode blades. However, this configuration has a few disadvantages: the residence time in the plasma is quite short, flow rates are more limited and, due to its geometry, only a limited fraction of the gas flow is processed by the discharge (e.g. about 20% depending on the actual geometry). 97,156,157 As a result, this limits the theoretical maximum possible conversion to $\sim 20\%$. The GAP configuration, on the other hand, is based on cylindrical electrodes, and the gas follows a vortex flow pattern. The gas in the reverse (inner) vortex flow passes exactly through the arc in the longitudinal direction, which ensures longer residence times in the discharge zone, even at high flow rates. Based on the gas flow calculations, about 40% of the gas flow can be processed by the discharge, 158 doubling the theoretical maximum conversion compared to the classical 2D electrode configuration.

Beside these geometry variations, other work has focused on changing the applied power, gas flow rate, flow type, interelectrode gap, admixture gases and plasma chemistry modelling. Again, all the data available in the literature are plotted in Fig. 22. Most experiments report a maximum energy efficiency of around 40-50%, 97,100,159 with the highest experimentally reported and also calculated energy efficiency being 65%.97,160 Most of the conversion results, however, remain below 15%, for both the regular GA and the GAP set-ups. The only exception is the work of Indarto et al. 161 (35% conversion), where the mixture was diluted with N2, but at the expense of the energy efficiency (<5%). As such, compared to DBDs, which also work at atmospheric pressures, GA plasmas deliver about the same conversion, but the energy efficiency is in general 3-4 times higher. A more detailed discussion on the influence of the different parameters is given below.

It should be no surprise that again a trade-off between the energy efficiency and conversion as a function of SEI is observed. 97,100,159,161,162 In general, the conversion increases and the energy efficiency decreases with increasing the SEI, as can be seen in Fig. 22, and the same optimal SEI range of 0.1-1 eV per molecule, in terms of energy efficiency, is observed as for the MW discharges. For a regular GA, the conversion clearly increases and the energy efficiency decreases when more power is supplied.97 The GAP geometry, on the other hand, can

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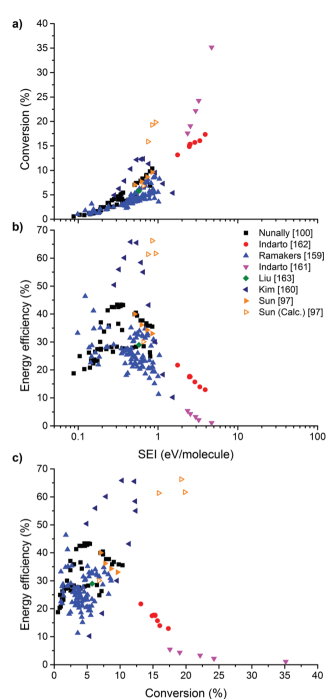


Fig. 22 Experimental data collected from the literature for CO₂ splitting in a GA, showing the conversion (a) and energy efficiency (b) as a function of the SEI, as well as the energy efficiency as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

operate in two regimes depending on the power input, i.e. a low and a high current regime, with the highest energy efficiency but lowest conversion observed for the former and the opposite for the latter. 159 Furthermore, it is reported that the lower the gas flow rate is, the higher the conversion is, both in the low and high current regimes, which is a direct result of the longer residence time of the gas in the discharge. 159 The same, albeit a

slightly less pronounced effect, was observed for the regular GA.161 Kim et al.,160 however, observed an optimum conversion while varying the gas flow rate. This was probably the result of reaching an optimum SEI (0.45 eV per molecule), since this value is in the same order as the SEI range at which Nunnally et al. reported a peak energy efficiency (0.30-0.37 eV per molecule).100

Besides the gas flow rate, an important parameter for the GAP is the vortex flow type, which can be modulated by adjusting the reactor geometry. 100,159,163 Operating in a reversed vortex flow (RVF), compared to a forward vortex flow (FVF), provides an increase in residence time and thermal insulation of the discharge from the reactor walls. As a result, the RVF delivers higher energy efficiencies at higher SEI values, leading to an improved conversion. 100,159 It is also this vortex flow that allows for the higher gas flow rates to be processed compared to a regular GA, and theoretically also to obtain higher maximum conversions because the gas passes through the arc in the longitudinal direction, thereby yielding a longer residence time.

For the regular GA, the interelectrode gap can be varied to improve the CO₂ conversion, and the best result was observed for the smallest interelectrode distance. Indeed, increasing this distance leads to a larger arc volume and a corresponding drop in plasma power and electron density, and consequently also a drop in CO₂ conversion.⁹⁷

Only one research group has reported the use of additive gases, other than CH4, H2O or H2; more specifically, the addition of N2, O2 and air. 161 The experiments revealed that adding N2 had a positive effect on the conversion (up to a factor 2 upon the addition of 95% N2), but due to the dilution, a detrimental effect on the energy efficiency was observed (down by a factor of 10-20). Furthermore, the presence of N₂ leads to the unwanted production of NOx. The addition of O2 and air showed a decrease in the conversion, indicating the possible strong negative effect of the presence of O2 (impurities), presumably due to Le Chatelier's principle.

Due to the more complex behaviour of the gas flow and of the arc movement, and also the relationship between both in a GA, modelling work is more limited in the literature. For the regular GA, a simple plasma kinetic model has been developed, 161 while recently a more detailed 0D chemical kinetic model97 and a 1D quasi-gliding arc mode164 have been presented. The results from these modelling studies show that the electron-impact dissociation of vibrationally excited CO₂ is predominant for an arc temperature of 1200 K and the recombination between CO and O atoms is the main conversion limiting reaction. 97 Reducing the arc temperature to 1000 K can significantly increase the conversion and energy efficiency, because it limits the recombination reaction rate and enhances the importance of the higher vibrational levels in CO₂ dissociation.97 Just like for MW chemistry, lumping of the vibrational levels into a limited number of groups has been successfully performed, opening future perspectives for 2D and 3D GA modelling. 164 A successful first attempt at modelling the more complex GAP with a 0D chemical kinetics model supported by 3D gas flow modelling has also been reported, and has confirmed the assumption, in line with the regular GA, that

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the vibrational levels play an important role for the energyefficient conversion of CO2. 159

In summary, the data in the literature show that GA discharges succeed in exploiting the most energy-efficient CO₂ dissociation channel based on vibrational excitation, while operating at atmospheric pressure. Energy efficiencies of 45% are no exception and even results above the target value of 60% have already been reported. On the other hand, model calculations reveal that the non-equilibrium character of the GA - making full potential of the vibrational excitation/dissociation pathway - could be further exploited to improve the energy efficiency. Just like for MW discharges, operating GA plasmas at lower gas temperatures might be the key for achieving this. In addition, the main limiting factor compared to MW discharges appears to be the conversion, due to the limited fraction of the gas flow that is currently processed by the discharge. Smart reactor design, such as enhancing the processed gas fraction, should lead to the necessary advancements to overcome this limitation in the future.

5.1.4. Other plasma types. Beside the three most common plasma types discussed above, a number of other non-thermal plasmas have been used for CO₂ conversion, including corona discharges, 165-170 glow discharges, 171-174 non-self-sustained discharges, 174 capillary discharges 175 and nanosecond pulsed discharges. 176 For more details and background regarding these different discharges, we refer to the respective references. When available, the relevant data have been extracted from the literature and the results are presented in Fig. 23. In general, the performance of all these different discharges is similar to that observed for a DBD, with maximum conversions of up to 40% and energy efficiencies below 15%. The main exception is the non-self-sustained discharge investigated by Andreev et al., 174 which reached a conversion of 50% with a corresponding energy efficiency of almost 30%. The advantage of this discharge type is its ability to control the mean electron energy by changing the E/n. The higher energy efficiency is reached when operating at values around 20 Td, which favours the vibrational excitation mechanism, as shown in Section 4.1.4 (Fig. 11 and 12). However, just like for the MW plasmas, this discharge operates at reduced pressures (1550 Pa).

5.1.5. Summary. To summarize, in Fig. 24, we plot the energy efficiency as a function of the CO₂ conversion, grouped per discharge type, for all the data discussed above. Furthermore, both the thermal equilibrium limit (see Section 2.1; Fig. 2) and the target energy efficiency of 60% (see the beginning of this section) are displayed. This figure allows us to draw the following conclusions.

First of all, it is clear that, although DBDs are among the most extensively studied for CO2 conversion, and indeed are already successfully applied80 for commercial O3 production and VOC removal, they appear to be unsuitable for the efficient conversion of CO₂. Their energy efficiency remains a factor of 4 too low, even when combined with a packing, in order to justify them as industrially competitive. The same applies to most of the other plasma types listed in Section 5.1.4 above.

Second, the best results for GA plasmas are capable of reaching the set energy efficiency target, namely 60%. Moreover,

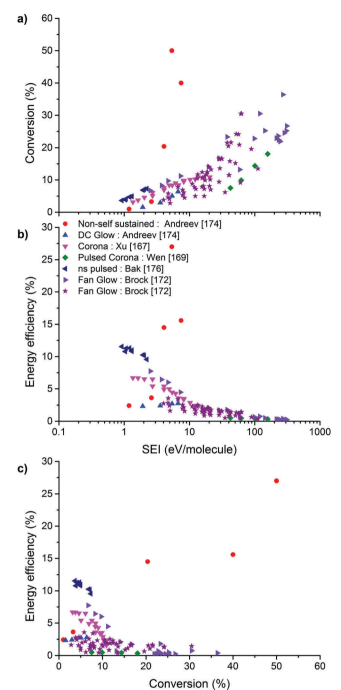


Fig. 23 Experimental data collected from the literature for CO₂ splitting in other plasmas, besides DBD, MW/RF and GA, showing the conversion (a) and energy efficiency (b) as a function of the SEI, as well as the energy efficiency as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account

almost all of the results obtained are far above the thermal equilibrium limit, which is especially interesting, keeping in mind that the GA plasmas operate at atmospheric pressure. This demonstrates the non-equilibrium character of this type of plasma, even at atmospheric pressure, and the benefits of being able to exploit this behaviour through the energy-efficient **Review Article**

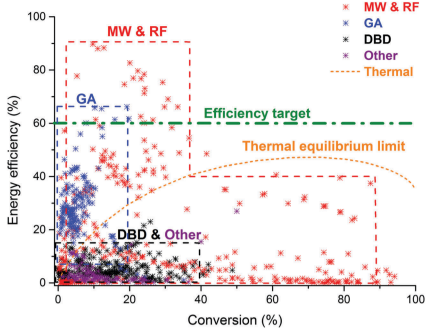


Fig. 24 Comparison of all the data collected from the literature for CO₂ splitting in the different plasma types, showing the energy efficiency as a function of the conversion. The thermal equilibrium limit and the 60% efficiency target are also indicated

dissociation of the CO₂ vibrational levels. Moreover, modelling has revealed that this non-equilibrium character could (and should) be further exploited, to further enhance the energy efficiency. To date, the main challenge is the limited conversion, which remains below 20% because only a limited fraction of the gas passes through the active arc plasma.

Finally, if, for now, we ignore the fact that most MW discharges used for CO₂ conversion operate at reduced pressure, in contrast to the commercially more interesting atmospheric pressure of GA plasmas, it is clear that MW discharges offer a wide variety of possibilities. Even up to conversions of 40%, the energy efficiency target is easily crossed and they clearly operate in a non-equilibrium regime, thus favouring the step-wise vibrational-dissociation mechanism. Conversions in the range of 40-90% are also possible, albeit with maximum energy efficiencies of only up to 40%. Under these conditions, the MW discharges most probably operate in the thermal regime. Nevertheless, this shows the wide variety of both conversions and energy efficiencies achievable with MW discharges for the conversion of pure CO2.

5.2. $CO_2 + CH_4$: dry reforming of methane

Contrary to pure CO₂ splitting, DRM can yield a wide variety of products. This has several implications for the evaluation and comparison of the various studies in the literature, mainly because it affects the definition of the energy efficiency. To determine the true energy efficiency of the process, we would need to take all the formed products into account - both gaseous and liquids - to determine the theoretical reaction enthalpy. Another possibility would be to determine the thermal energy efficiency (see Section 6), based on the higher (or lower)

heating value of the output, i.e. the products, relative to that of the input, i.e. the reactants and power input. However, typically only the selectivity (or yield) towards the syngas components, CO and H2, and light hydrocarbons is given for most studies in the literature. Thus, at the moment these strategies are impossible to pursue when presenting an overview of the available literature results in the sections below. Therefore, in this section, we use the energy cost, in units of eV per converted molecule, to compare the different discharges, together with their total conversion. As an energy efficiency target for syngas production, we take the same 60% value for the general stoichiometric DRM process, which equals an energy cost of 4.27 eV per molecule converted. Indeed, 100% energy efficiency would yield an energy cost equal to the standard reaction enthalpy of 2.56 eV per molecule (as outlined in Section 2.2.1). In our opinion, this is the best representation to compare the results in the literature, considering the fact that in most of the studies, the higher hydrocarbons and the valuable liquid fraction containing oxygenates (e.g. formaldehyde, methanol, ethanol, formic acid, acetic acid) are neglected. However, as explained at the beginning of Section 5, this 60% energy efficiency target is only valid for the comparison towards syngas production. When liquids (such as methanol) are formed through the direct oxidative pathway, which has already been demonstrated using the plasmachemical conversion of CO2 with a co-reactant, the energy efficiency requirements are drastically lowered by a factor of two to three. This is a direct result of circumventing the energy-intensive step of further processing the syngas into the desired liquid products.

Plasma-based DRM has received a lot of attention in recent years. Some overviews can be found in the literature. 21,41,177-179 In a recent review on the liquefaction of methane, 112 DRM was Chem Soc Rev

considered as oxidative plasma-based CH₄ liquefaction. It can be performed in two ways: either as a two-step process, yielding mainly syngas, which can then be further processed into Fischer–Tropsch liquids or methanol, or as a one-step process, aiming for the direct formation of value-added oxygenated products, such as formaldehyde, methanol, ethanol, dimethyl ether and formic acid. These two options are therefore called the indirect and direct oxidative plasma liquefaction approaches, respectively. An overview of the progress in the field of plasma-based DRM is given below, starting again with DBD, MW and GA plasmas. However, in contrast to CO₂ splitting, a vast amount of research has been performed with some other discharges as well for DRM, so these have been given their separate sections below.

First some general trends applicable for all discharges are summarized. We start off with the SEI, because for all discharges studied, it is clear that for DRM - just like for pure CO₂ splitting - the SEI is the major determining factor for the conversion and energy cost, as it combines the effects of power and residence time. Typically, a higher SEI leads to a higher conversion, but also incurs a higher energy cost. The conversions of CH₄ and CO₂ increase almost linearly when a higher voltage or input power is applied, as observed for DBD, 106,180-190 MW, 191 GA, ^{192–194} corona, ^{195–198} spark, ^{199–205} APGD^{206,207} and nanosecond pulsed discharges. 208-212 Generally, this is accompanied by an increase in energy cost. Regarding product selectivity, some different trends are reported as a function of power for the different discharge types. In general, in DBDs, higher H_2 and CO yields are reported, with the ratio of H₂ to CO remaining constant, 184,187,188 as well as a sharp drop in the selectivity of the light hydrocarbons. 186,189,190 Furthermore, some studies observed no significant changes in the selectivities in a DBD, except for an increase in carbon deposition, 184 while other studies report an increase in CO and H₂ selectivity, 186,187,190 or even the opposite trend. 188 For corona discharges, the H2 selectivity seems to decrease, while the CO selectivity exhibits the opposite trend, 195,196 and as a result, the H₂/CO ratio greatly depends on the discharge power. 197 In spark discharges, a more pronounced carbon deposition at higher input power has been reported, 202 as well as an increased selectivity for H2 and CO and a drop in C2H2 selectivity, pointing to an enhancement of the reforming reactions over coupling reactions. 202,203 APGD discharges exhibit a higher CO selectivity and coke deposition at higher input voltages and these conditions favour the production of unsaturated hydrocarbons, such as C2H2.206 On the other hand, for nanosecond pulsed discharges, upon a higher SEI, the mass balance points to a loss of oxygen and carbon, which are converted into water and carbon powder or are deposited, respectively.²⁰⁸ Finally, for MW and GA plasmas, no specific trends have been reported yet for the effect of power on the product selectivities. Thus we can conclude that, albeit with some exceptions, a higher discharge power generally improves the conversion, but does not change the reaction pathways significantly.

A higher total flow rate has the opposite effect to the power, and leads to lower conversions and energy costs for DBD, ^{106,180,183,185,186,188,189,213} GA, ¹⁹⁴ corona, ^{196–198,214} spark, ^{199,202} APGD^{206,207} and nanosecond pulsed discharges. ²⁰⁹

No results have been reported yet for MW plasmas. On the other hand, the flow seems to have almost no effect on the selectivity towards the syngas components CO and H2 and hence also not on the syngas ratio itself, for DBD, 183,185,189,213 GA and APGD. 206,207 At higher flow rates, some researchers have also reported a decrease in H_2 and CO selectivity in DBDs, ^{186,188} while the selectivity towards hydrocarbons increases. 180,183,186,188,213 For corona discharges, a slightly higher H2 and CO selectivity is observed, but the syngas ratio remains almost unchanged. 196-198,214 For spark discharges, it is reported that the H₂ selectivity rises and the CO selectivity drops, at higher flow rates. 199 Finally, for MW and nanosecond pulsed discharges, again no specific data have been reported yet for the effect of flow rate on the product selectivities. Nevertheless, in general we may again conclude that the dissociation and formation mechanisms are, albeit with some exceptions, not significantly influenced by a change in the flow rate, although the selectivities can be altered.

Besides the SEI (or the power and gas flow rate), the CH₄/CO₂ mixing ratio has a tremendous influence on the CH₄ and CO₂ conversions, as well as on the product selectivities. In general, when adding more CH₄ to the mixture, the effective CH₄ conversion rises, while the effective CO₂ conversion drops. Note that we talk about the effective conversions (as defined in Section 4.4.2; eqn (4)) and not about the (absolute) conversion itself, in order to take the effect of the change in gas mixture into account. At the same time, the selectivity towards (light) hydrocarbons increases, as well as the selectivity towards H2, while the CO selectivity decreases, when more CH₄ is present in the mixture. This inevitably leads to an increase in the syngas ratio. Thus, the H₂/CO ratio is highly dependent on the inlet feed, which makes the ratio easily adjustable over a wide range to fit the Fischer-Tropsch or methanol synthesis requirements. This was found for all discharges, i.e. DBD, 182,183,186-190 GA, ^{192–194,215} MW, ¹⁹¹ corona, ^{197,214,216–218} spark, ^{200,202,204,205} APGD^{207,217,219,220} and nanosecond pulsed discharges.^{209,211,221} The formation of carbon black and its deposition on the electrode and reactor walls for mixtures seems to occur with a high CH₄/CO₂ ratio (>1), and this deposition can highly influence the discharge operation. 119 For lower $\mathrm{CH_4/CO_2}$ ratios, this is not observed, neither in DBD, ^{182,189,190} GA, ^{192–194,215} corona, ^{195–197} spark.^{200,205} APGD^{207,217,219,220} nor nanosecond pulsed discharges, 209 as CO2 prevents carbon formation. For APGDs, a higher CH₄/CO₂ ratio is reported to increase the water formation. In contrast, for a DBD¹⁸⁷ and spark discharge, ²⁰² a higher water production is observed at low CH₄/CO₂ feed ratios, presumably due to the occurrence of the reverse water-gas shift reaction.

These trends can be explained by the following reactions: increasing the CH_4/CO_2 ratio leads to a more pronounced H_2 formation and at the same time it reduces the amount of O species in the mixture. Hence, the reaction $C+O\to CO$ is reduced, leading to a lower CO selectivity, and increased carbon deposition. On the other hand, the reaction $H_2+O\to H_2O$ becomes more important, giving rise to the higher H_2O production, but it is not significant enough to balance the higher H_2 production, thus explaining the increased H_2 selectivity. This also means that the deposited C is still an active species in the

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reactions, and it is suggested that the addition of O2 207 and/or H₂O²¹⁷ can prevent carbon deposition. Finally, it is reported that the rate of CO production is always higher than the rate of CO₂ reduction, proving that CO is formed by both the reduction of CO₂ and the oxidation of CH₄. 214

The effect of the CH₄/CO₂ mixing ratio on the total conversion and hence on the energy cost is less straightforward. For nanosecond pulsed discharges, depending on the set-up, different optimum values have been reported for this mixing ratio for achieving the lowest energy cost. Ghorbanzadeh et al. 209,212 reported a considerable increase in the energy cost when methane becomes predominant in the mixture, while Zhang et al. 211 reported the energy cost was cut in half when going from a 1:3 to a 3:1 CH₄/CO₂ ratio. For a DBD, the effect of the mixing ratio on the total conversion also appears to depend on the since both an increase 182,186,222,223 decrease 106,183,188 are widely reported upon the addition of more CH₄. Snoeckx et al., ¹⁸¹ Zheng et al. ²¹³ and Pinhão et al. ²²⁴ all reported an initial increase in total conversion upon adding CH₄ when the CH₄/CO₂ ratio was <1, and a decrease when the ratio became >1. These trends suggest that the CH₄ conversion is strongly affected by the CO2 conversion, especially when the ratio is <1 and when CO_2 is thus the main component in the mixture. For the other plasma types, no specific data have yet been reported in the literature on the effect of the CH₄/CO₂ mixing ratio on the total conversion. Finally, as the energy cost is inversely proportional to the obtained conversion (cf. Section 4.4.3; eqn (7)), the effect of the CH₄/CO₂ mixing ratio on the energy cost is opposite to the effects described above for the total conversion.

Besides these general trends for the effect of the SEI and the CH₄/CO₂ mixing ratio, which are very similar for the various discharge types, some specific trends characteristic for each plasma reactor are reported for DRM. These are summarized in the next sections, along with a summary of all the values for conversion and energy cost reported in the literature.

5.2.1. DBD plasmas. Due to its simple design and ease of use - also in combination with packing materials - again most of the research to date has been performed with DBDs, with the coaxial reactor (see Section 4.2.1; Fig. 13) being the main geometry. To improve the conversion and energy costs, and to tune the product distribution, several approaches have been investigated, including changing the SEI, applying pulsed power, changing the gas flow rate or feed ratio, changing the reaction temperature or pressure, applying multi-electrode configurations or different electrode materials, and mixing with other gases, *i.e.* Ar, He, N_2 , as well as introducing (catalytic) packing materials. Some of these effects have already been described above, but the others are explained in more detail below. Furthermore, extensive modelling work has also been performed to obtain more fundamental insights into the plasma chemistry and in turn to aid in the improvement of future experiments. For all the experiments reported in the literature, the syngas components H₂ and CO were the major reaction products, with smaller amounts of C2 and C3 hydrocarbons also detected, in the order: $C_2H_6 \gg C_3H_8 > C_2H_4 >$ C₂H₂. In even lower quantities, hydrocarbons up to C₅ have been detected, where the selectivities of unsaturated hydrocarbons are again much lower than those of paraffins. The major products detected when the condensate was analysed 183,222,225,226 were C₄-C₁₀ hydrocarbons and oxygenates, including methanol, ethanol, 1-propanol, acetic acid and other e.g. tertiary alcohols, ketones, esters and carboxylic acids.

Fig. 25 summarizes most of the results, in terms of both conversion and energy cost, available in the literature. Due to

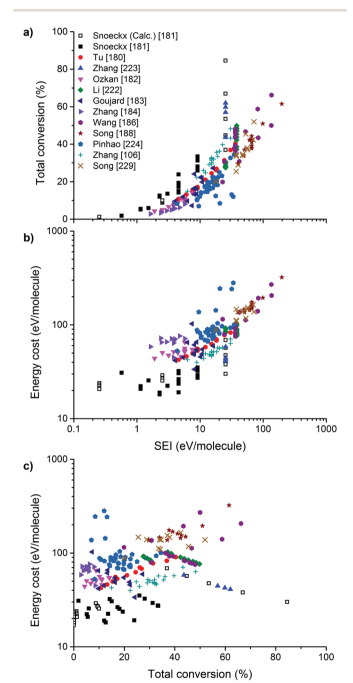


Fig. 25 Experimental data collected from the literature for DRM in a DBD, showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

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Mahammadunnisa [235] a) 100 Zena [213] Tu [180] Tu [240] Gallon [246] 80 Zhang [242] Pham [243] Total conversion (%) Sentek [244] Eliasson [248] 60 Wang [239] Wang [233] Song [188] Krawczyk [245] Zhang [106] Mei [238] 20 Zheng [237] Chung [189] Zheng [241] Zou [250] 1000 Energy cost (eV/molecule) 100 10 1000 0.1 10 100 SEI (eV/molecule) c)

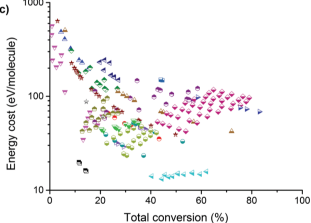


Fig. 26 Experimental data collected from the literature for DRM in a packed-bed DBD, showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

the large amount of data available for packed-bed reactors, those data are discussed and presented further below in Fig. 26. The conversion and energy cost increase again upon a rising SEI. When plotting the energy cost as a function of the conversion, it is clear that most of the results are situated above an energy cost of 20 eV per molecule and below a conversion of 60%, albeit with some exceptions. The highest experimental conversion reported is 66%, 186 and the lowest experimentally

observed energy cost is 18 eV per molecule. 181 This is more than a factor of 4 higher than the set efficiency target of 4.27 eV per molecule (see the beginning of Section 5.2). Model calculations predict that higher conversion and lower energy costs should be achievable by a careful selection of the operating conditions, as will be outlined below. 181 It is important to note that some of the data in this figure have again been recalculated to represent coherent values for the conversions and energy costs. For instance, as also explained for pure $\rm CO_2$ splitting, the mixture is sometimes diluted with He, Ar or $\rm N_2$ to obtain an more easily ignitable and more stable discharge, but this influences the results – an effect that was not always taken into account in the original data. A more detailed influence of the different parameters is discussed below.

The effect of the temperature and pressure on DRM in a DBD has been investigated by several authors. Low pressures seem to favour the reactant conversion, whereas the syngas ratio does not show any pressure dependence. 187 On the other hand, the selectivity of both CO and light hydrocarbons is reported to increase with rising pressure. 190 For a wall temperature range of 353-523 K, Zhou et al. 187 observed slightly higher conversions and product yields upon increasing the temperature, with the syngas ratio being independent of the temperature. Zhang et al. 184 and Goujard et al. 183 investigated the temperature effect in the range of 297-773 K and 298-873 K, respectively. Both the conversions and the hydrocarbon selectivity were reported to increase with temperature, while the H₂ and CO selectivity decreased. However, Zhang et al. 184 found an increase in the syngas ratio, whereas Goujard et al. 183 observed the opposite, noting also that when the temperature reached 773 K, the CO₂ conversion appeared to increase more rapidly than the CH₄ conversion, indicating a shift in the dominant reforming processes around 673 K. 183 These results indicate that while the electron-impact dissociation reactions govern the reactant conversions, the thermochemistry can be used to control the consecutive product reaction pathways.

Several changes with respect to the electrode configuration and materials have been reported as well. Wang et al. 186 found little influence of the discharge gap on the reactions, while Li et al.222 observed higher conversions upon reducing the discharge gap. At the same time, Wang et al. 186 reported a significant influence on the conversions using multi-stage ionization (i.e. multiple DBD and 'afterglow' zones in one reactor by placing several electrodes in series), while only small effects were observed by Li et al.222 Both groups, however, found a significant positive influence on the syngas product selectivities when applying multi-stage ionization. Rico et al. 185 reported better conversions with porous electrodes, while the (catalytic) effect of different inner electrodes only showed very little influence on the conversions, 227,228 but a significant increase in the oxygenate synthesis for nickel and copper, suggesting a catalytic role of the metallic surface. 228 Finally, applying a pulsed power also seemed to have a beneficial effect on the conversion.229

As mentioned above, some researchers added inert gases, such as N₂, Ar and He, to ignite the plasma more easily.

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However, this also has several effects on the discharge characteristics, conversion and energy cost. Ozkan et al. 182 investigated the effect of Ar and He dilution, but did not compare the results with pure CH₄/CO₂ mixtures. The CH₄ conversion was higher for He than for Ar addition, while the opposite was found for CO2 conversion. This could be related to a change from filamentary to glow discharge when switching from Ar to He, which significantly influences the shape of the EEDF and thus the electron-impact reactions. Zhang et al. 106 indicated that, although the conversion increases when adding Ar, the latter is not sufficient to counteract the lower CH₄/CO₂ feed content, since the conversion rates are lower compared to a mixture without Ar. On the other hand, they observed a positive effect towards the syngas component selectivities. Pinhão et al. 224 reported a positive effect on the breakdown voltage with increasing He content, resulting in a rise in the conversion rates with increasing the He content up to 80%. However, a still higher He content showed detrimental effects on both the conversion and selectivity, but again no comparison for the mixture without He was made. In a follow-up modelling study by Janeco et al., 92 a shift in the electron velocity distribution function due to the addition of He was observed.

Kolb *et al.* studied the addition of the molecular gases O_2 and H_2O on the performance of DRM in a DBD. 230,231 They found that O_2 aids in the conversion of CH_4 but also produces CO_2 , 231 which is not beneficial. H_2O , on the other hand, only has a small negative influence on the conversion of CO_2 , but significantly enhances the conversion of CH_4 and, more importantly, the production of valuable oxygenates, such as formal-dehyde and methanol. 230 This study showed that a tri- or more general multi-reforming process, using a mixture of $CO_2/CH_4/H_2O$ or a combination of different reforming processes in series, could be a very interesting next step to pursue using plasma technology.

Due to the historical interest in thermocatalytic DRM and the ease of implementing a (catalytic) packing in the discharge zone of a DBD, as explained in Section 4.3, a lot of research has already been performed towards plasma-catalytic DRM and hybrid reactors. A complete but brief overview is given below, but for more detailed information on the effects and mechanisms behind plasma-catalysis, we refer to the existing excellent literature on this topic. 102,103,232 Most of the work to date has been performed with packed-bed reactors, but some research also exists on fluidized-bed reactors. 233,234 Contrary to pure CO₂ splitting, for DRM, the addition of packing can greatly influence the formation of products, due to the availability of CH₄ as a hydrogen source. Hence, the research work focuses both on increasing the conversion and lowering the energy cost by physical effects, as well as on chemical effects to steer the product distribution towards more value-added chemicals, such as light hydrocarbons and oxygenates. Packed-bed DBD reactors generally yield similar conversions and energy costs as the corresponding empty reactors, as can be seen when comparing Fig. 25 and 26. Nevertheless, the use of a catalytic packing can drastically alter the chemical pathways and thus the selectivity and product distributions, which would be

beneficial for exploiting the direct oxidative pathway of DRM, instead of utilizing the indirect process through syngas production.

A broad spectrum of materials have already been investigated for the plasma-catalytic DRM, of which Ni is by far the most commonly used active phase, such as in Ni/ γ -Al $_2$ O $_3$, $^{106,180,188,213,233,235-240}$ Ni/SiO $_2$, 238,241,242 Ni–Fe/ γ -Al $_2$ O $_3$, 237 Ni–Fe/SiO $_2$, 237,242 Ni–Cu/ γ -Al $_2$ O $_3$, 106 Ni 0 /La $_2$ O $_3$, 183 Ni/MgO, 238 Ni/TiO $_2$, 238 NiFe $_2$ O $_4$, 237 NiFe $_2$ O $_4$ #SiO $_2$, 237 LaNiO₃/SiO₂,²⁴¹ LaNiO₃²⁴¹ and LaNiO₃@SiO₂.²⁴¹ Furthermore, alumina is the most commonly used support, i.e. in Ni/ γ -Al₂O₃, ${}^{106,180,188,213,233,235-240}$ Ni-Fe/ γ -Al₂O₃, 237 Mn/ γ -Al₂O₃, 213 $Cu/\gamma-Al_2O_3$, 106,213,234 $Co/\gamma-Al_2O_3$, 213 $La_2O_3/\gamma-Al_2O_3$, 243 Ag/ γ -Al₂O₃, ²⁴⁴ Pd/ γ -Al₂O₃, ²³⁴, 244 Fe/ γ -Al₂O₃ ²⁴⁵ and Cu-Ni/ γ -Al₂O₃,¹⁰⁶ or even in its pure form. ^{185,188,244,246} Many other catalytic systems are based on zeolites, e.g. 3A, 246 A4, 247 NaX, 248 NaY²⁴⁵ and Na-ZSM-5. ²⁴⁵ Besides, studies have also been conducted using BaTiO₃, ^{185,242} a mixture of BaTiO₃ and NiSiO₂, ²⁴² ceramic foams (92% Al₂O₃, 8% SiO₂) coated with Rh, Ni or NiCa, 249 quartz wool,²⁴⁶ glass beads,¹⁸⁹ a stainless steel mesh,²⁴² starch,²⁵⁰ BZT (BaZr_{0.75}T_{0.25}O₃) and BFN (BaFe_{0.5}Nb_{0.5}O₃). For a regular ACpacked DBD, the best result was obtained for the Zeolite Na-ZSM-5, with a total conversion of 37% and an energy cost of 24 eV per converted molecule (see Fig. 26). 245 Just like for pure CO2 splitting, the addition of a catalyst does not seem to make the process more energy efficient, but it does yield higher conversions at the same energy cost. The best overall results in a packed-bed DBD were obtained for a quasi-pulsed DBD packed with BFN and BZN, with total conversions in the range of 45-60% and an energy cost in the range of 13-16 eV per converted molecule (see Fig. 26), 189 which is lower than that for a DBD without packing, but this might also be due to the pulsed operation.

Besides the experimental work, major insights have been obtained in recent years based on modelling of the DRM process for a DBD. Different kinds of models and computational techniques have been successfully developed, including semi-empirical kinetic models, ^{187,251,252} zero-dimensional chemical kinetic models with both simplified²⁵³ and extensive chemistry sets, ^{131,181,254} a one-dimensional fluid model, ²⁵⁵ a so-called 3D Incompressible Navier–Stokes model combined with a convection–diffusion model, ²⁵⁶ a hybrid artificial neural network-genetic algorithm, ²⁵⁷ a model focusing on a more accurate description of the electron kinetics ⁹² and density functional theory (DFT) studies, to investigate the reaction mechanisms. ^{226,258} Due to the complex chemistry taking place in a DRM, the development of accurate multidimensional models with extensive chemistry is currently restricted by computational limits.

Some key findings of these models are presented here. Snoeckx *et al.*¹⁸¹ performed an extensive modelling study, with detailed plasma chemistry, spanning a wide range of experimentally accessible conditions. The model predicted that increasing the SEI at a constant gas ratio and frequency results in a higher total conversion. However, the increase in conversion is not entirely proportional to the rise in SEI, resulting in somewhat higher energy costs with increasing the SEI. The lowest energy cost was predicted to be 16.9 eV per molecule, but this corresponds to a very low value for the total conversion, *i.e.*

0.015%. On the other hand, the highest total conversion predicted by the model, i.e. 84.2%, corresponded to an energy cost of 30.1 eV per molecule. A larger amount of CO2 was observed to lead to a higher total conversion and lower energy cost. This is attributed to the O atoms formed by the electron-impact dissociation of CO2, which reacts very effectively with the H atoms originating from the electron-impact dissociation of CH₄. As shown in the kinetics analysis of Snoeckx et al., ²⁵⁴ the conversion of CH4 is normally limited by the fast backward reaction, i.e. $CH_3 + H \rightarrow CH_4$, but when more O atoms are available, this reaction is of minor importance compared to the reaction O + H/OH \rightarrow OH/H₂O. Thus, by limiting the backward reaction, the conversion of CH₄ rises dramatically with increasing CO₂ content, leading to a higher total conversion. Another important effect is the total number of micro-discharge filaments that the gas molecules experience when passing through the DBD reactor. It seems that for most cases a larger number of filaments, but with lower energy, yields higher values for the conversion and lower energy costs, compared to a smaller number of filaments, but with more energy deposited per filament. 181 In some other models by Janeco et al. 92 and Goujard et al., 253 the effect of adding He to the CH₄/CO₂ mixture was investigated. Both groups reported that He, as well as all other species that may result from the electron collisions in the gas mixture, can significantly change the electron velocity distribution function, the electron reaction rates and the energy losses, which can ultimately affect the energy cost when taken into account in the model.

Summarizing all the data in the literature, we can conclude that a DBD reactor can provide reasonable conversions of up to 60% and 80% for empty (i.e. non-packed) and packed-bed DBDs, respectively. However, the energy cost lies in the range of 20-100 eV per molecule, which corresponds to an energy efficiency of 12.8-2.6%, assuming that only syngas is formed. This is at least a factor 5 away from the necessary 4.27 eV per molecule benchmark for the energy cost (or the 60% benchmark for energy efficiency), as defined at the beginning of this section, with no positive outlook on the horizon. Indeed, even extensive modelling studies have predicted that the lowest achievable energy cost, obtained after careful selection of the operating conditions, would be 16.9 eV per molecule (see above). 181 Hence, despite a simple reactor design for scalability, the ease of implementing a wide variety of packings and its industrial success for ozone generation (and possibly VOC abatement), at this point, the high energy cost - and hence low energy efficiency - makes it doubtful that DBDs will be the most suitable technology for DRM into syngas. Nevertheless, due to its ease of use, research with DBDs still can yield valuable insights and knowledge, which may be transferable to other more efficient discharges. On the other hand, if suitable catalysts could be found for the production of valuable oxygenates with high yields through direct (plasma-catalytic) oxidative liquefaction, 112 the energy efficiency target would be significantly lower (by a factor of 2-3 in the case of methanol), so this could change the analysis drastically. However, many challenges for this pathway remain, such as finding suitable catalysts, and advances in energy efficiency are also still required.

5.2.2. MW plasmas. Compared to the extensive work available on pure CO2 splitting in MW plasmas, almost no work exists for DRM using a MW plasma. This is in great contrast to the extensive work performed for DRM in DBD reactors (see above). We are aware of only one study for a pulsed MW plasma, 259 and one for a continuous MW plasma, where a comparison is made with the introduction of a Ni catalyst. 191 The pulsed set-up achieved CH4 and CO₂ conversions of 71% and 69%, respectively, at an energy cost of 6.5 eV per molecule for a CH₄/CO₂ ratio of 1.5. Syngas was again the major product, but C₂H₂ and C₂H₄ were also detected. ²⁵⁹ Furthermore, no coking was observed. The data are presented together with the results for the GA plasma in Fig. 27.

Cho et al. 191 obtained conversions in continuous MW plasma in the same range as in the pulsed MW plasma of Zhang et al., 259 but at much high powers, up to 1.5 kW compared to 120 W, and at low flow rates (100 mL min⁻¹ ν s. 200 mL min⁻¹), so their energy costs were unacceptably high, i.e. between 93 and 343 eV per molecule. Adding a Ni catalyst to the discharge was reported to lead to a higher conversion by 10% for CH₄ and by 15% for CO₂. Furthermore, the H₂/CO ratio changed drastically from 1.7 for the plasma-only case to 1.2 when combined with the catalyst. As additional products, only C2H4 was detected, with increasing selectivity when more CH4 was added to the feed. 191

In Section 3.5, we also discussed catalysis based on MWheating. Fidalgo et al.76 observed the formation of microplasmas in their so-called microwave receptor/catalyst set-up. However, the MW source was simply used in that case to heat the gas, with the creation of microplasmas at the catalyst surface only a side effect of the strong electric fields at the sharp edges.

In general, the few studies reported so far in literature for DRM by MW plasmas show high conversions, as well as a high selectivity towards CO and H₂. Furthermore, based on the pure CO₂ splitting results presented in the previous section, we might expect that a large treatment capacity is in principle possible, as well as high energy efficiencies. Nevertheless, experimental evidence to date remains extremely scarce. One of the major unanswered questions is related to the role played by the vibrational levels of CH4 in the CO2 dissociation, and more specifically in the ladder-climbing effect, which was explained in Section 4.1.4 (cf. Fig. 12). In particular, it is not clear whether they will affect (i.e. stimulate or inhibit) this process or not, as well as how the dissociation of CH₄ itself will be influenced.

5.2.3. GA plasmas. Most of the research for DRM with GA plasmas is based upon 2D bladed electrodes, 192,194,215,260-263 while only some studies have reported the use of a rotating GA discharge, 193,264-266 and even then, mostly for biogas instead of a pure CH₄/CO₂ mixture. 264-266 All the data available in the literature are plotted in Fig. 27. Conversions in the range of 30-50% for both CH₄ and CO₂ have been reached, with energy costs as low as 1-2 eV per converted molecule. The best result achieved a total conversion of 39%, with an energy cost of only 1 eV per molecule. 193 The main products formed seem to be the syngas components, unsaturated C2 hydrocarbons (i.e. C₂H₂ and C₂H₄) and solid carbon. Additionally, the

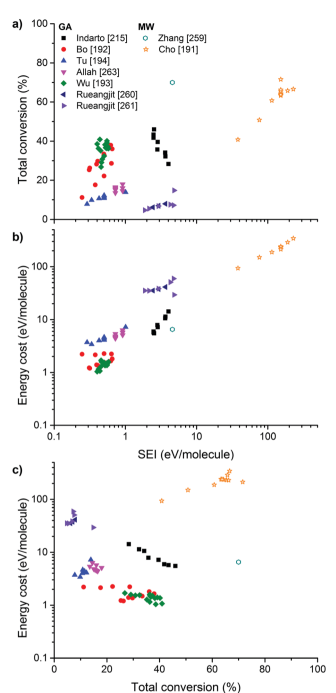


Fig. 27 Experimental data collected from the literature for DRM in MW (open symbols) and GA plasma (closed symbols), showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account

formation of different carbon materials, including spherical carbon nanoparticles, multi-wall carbon nanotubes and amorphous carbon, have also been reported. 194 Hence, compared to a DBD, the distribution of C2 hydrocarbons seems to be shifted from C₂H₆ to C₂H₂ and C₂H₄, and there is no formation of C₃ hydrocarbons reported. 194 This shift is commonly observed for discharges where the gas temperature is higher. Furthermore, compared to a DBD, which also operates at atmospheric pressure, GA plasmas deliver about the same conversion, but the energy cost is, in general, 20 times lower. This follows directly from Fig. 27, because the energy input is also an order of magnitude lower. A more detailed discussion on the influence of the different parameters is given below.

For GA discharges, the addition of inert gases has not yet been studied in the literature for DRM, but some studies have reported mixtures with reactive gases. Rueangjit et al. 260,261 investigated the dry reforming of simulated natural gas with a high CO₂ content, represented by CH₄: C₂H₆: C₃H₈: CO₂ in a molar ratio of 70:5:5:20. However, only a comparison with mixtures without CO2 was made, and it was then concluded that CO2 considerably enhances the conversion of the hydrocarbons in the feed.²⁶⁰ In addition, very high syngas ratios, i.e. around 10-20, could be obtained by adding these small amounts of hydrocarbons in the feed, due to their lower C-H bond energy. 260,261

Furthermore, some research has also been performed on the combination of dry reforming with O2 and steam using GA plasmas, to mimic biogas feeds as an input source. 261,262,264-266 However, in this case, CO2 is not only converted but is also being produced from the reaction of the CH4 dissociation products with O atoms, 261,264,265 and in the case of H2O addition, even a net CO₂ formation can be observed. 262,266 It is suggested that when adding O2, the main process becomes the partial oxidation of methane, while the main involvement of CO₂ is in the reverse water-gas shift reaction.²⁶⁴ The main positive effect of adding O2 is minimization of the carbon deposit.261 When adding steam, it seems again possible to obtain high syngas ratios, i.e. around 7.262,266

For DRM, the addition of a catalytic bed after the GA discharge zone has been investigated in two studies.^{262,263} In both studies, a NiO/Al₂O₃ catalyst was used. It was found that the CO₂ and CH₄ conversion increased by 24% and 16%, respectively. The selectivity of the syngas components, on the other hand, was found to drop slightly and the unsaturated hydrocarbons, C₂H₂ and C₂H₄, were formed with selectivities ranging from 16% to 19%. Lower NiO loading and smaller particle sizes appeared to be beneficial. Finally, it was suggested that there is scope for further development and optimization using a fluidized bed for maximizing the plasmacatalyst interactions. 263

It is clear that the obtained energy costs for DRM using a GA plasma are much lower than those obtained for DBDs, while at the same time they allow achieving much higher feed processing capacities. Furthermore, the energy costs achieved already, i.e. around 1 eV per molecule, are already a factor of 4 better than the required target for syngas production. No data is available yet for the DRM of the GAP set-up, but we could expect even further improvements, as already demonstrated for pure CO₂ splitting (see Section 5.1.3 above). Finally, the use of a catalytic spouted bed might be able to influence the product distributions, as already reported for pure CH₄ reforming. 105

5.2.4. Corona discharges. As described in Section 4.2.4, corona discharges can operate in both a negative and positive mode, and both modes can be used to perform DRM, although

positive coronas seem to exhibit slightly higher conversions, while the H₂/CO ratio is slightly higher for negative coronas.¹⁹⁷ This effect can be attributed to the different characteristics and generation mechanisms between both operation modes. Nevertheless, most research on DRM is performed with positive coronas. Fig. 28 summarizes the available data from the

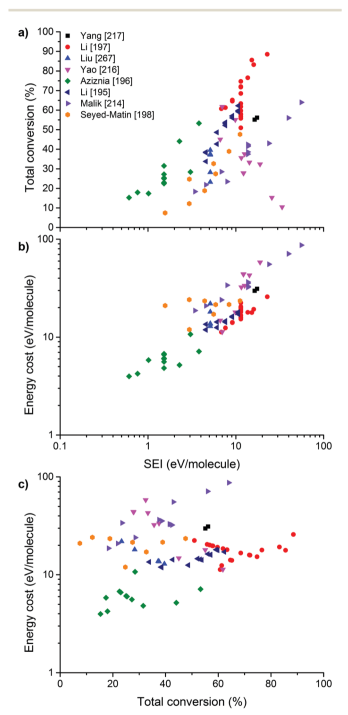


Fig. 28 Experimental data collected from the literature for DRM in a corona discharge, showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

literature. Both the conversion and energy cost increase with the SEI. Conversion values in the entire range up to 90% are achieved, with energy costs around 20 eV per molecule. The best result comprises a total conversion of 44% with an energy cost of 5.2 eV per molecule. 196 The main products formed are the syngas components, as well as C₂H₂, with some minor production of C₂H₆ and C₂H₄. ²¹⁶

A pulsed corona was investigated by Yao et al.,216 who reported that the conversion and CO selectivity increase with rising frequency, while the hydrocarbon selectivities showed the opposite trend.

Some experiments were also performed by placing catalysts in the corona discharge. Liu et al. 267 observed a significant conversion of CH₄ and CO₂ over a NaOH-treated Y zeolite placed in the discharge zone. The selectivity shifted towards higher hydrocarbons, e.g. propane, n-butane, isobutane, 1-butene and n-pentane. Aziznia et al. 196 investigated the effect of Ni/Al₂O₃ in the discharge zone. Upon the addition of Al₂O₃, the conversions decreased slightly; the CO selectivity also decreased, but the H2 selectively increased remarkably. When adding up to 20 wt% Ni, the CH₄ and CO₂ conversions increased from 18% and 25% to 23% and 36%, respectively. Furthermore, the selectivity towards CO was favoured over other carbon-containing compounds, and as a result, the H₂/CO ratio decreased, i.e. from 0.7 to 0.6 for Ni 20 wt%. It was suggested that CO2 molecules are readily chemisorbed and dissociated into CO on the surface of Ni-based catalysts, and in a high electric field the Ni sites become active and affect the conversion and selectivity. Finally, Li et al. 195 used both Ni/Al2O3 and the HZSM-5 zeolite ($SiO_2/Al_2O_3 = 38$) as catalysts. No significant effect on the conversions was observed, but the catalysts did affect the selectivities, whereby a higher CO and lower H₂ selectivity were observed for Ni/Al₂O₃, while higher H₂ and lower CO selectivities were measured for the HZSM-5 zeolite as the catalyst.

It is clear that corona discharges exhibit several similarities with DBDs, with an energy cost in the range of 4-100 eV per molecule and conversions of up to 60% (with exceptions even up to 80%). Thus, much work would still be required to lower the energy cost by a factor 2-3, which would be needed before corona plasmas could become a competitive alternative. In addition, it is difficult to achieve a high treatment capacity due to a corona's localized breakdown. 177 This leads us to believe that, at this point, corona discharges have the same negative outlook as DBDs.

5.2.5. Spark discharges. Spark discharges are also being explored for DRM, but the developments are rather new and limited to date. 199-205,268-272 The available data are presented in Fig. 29, showing a pronounced increase in total conversion with the SEI. Furthermore, the energy cost is between 3 and 10 eV per molecule, with the best results achieving a total conversion of 85% at an energy cost of barely 3.2 eV per molecule. 199 In addition to H2 and CO, C2H2 is also formed with selectivities of up to 40% and minor quantities of C₂H₄ and C₂H₆. ²⁰²

A spark discharge with different interelectrode distances and varying pressure and frequency was investigated by Zhu et al.²⁰³ With the interelectrode distance rising from 3 to 9 mm, a higher conversion was observed, followed by a slight decrease

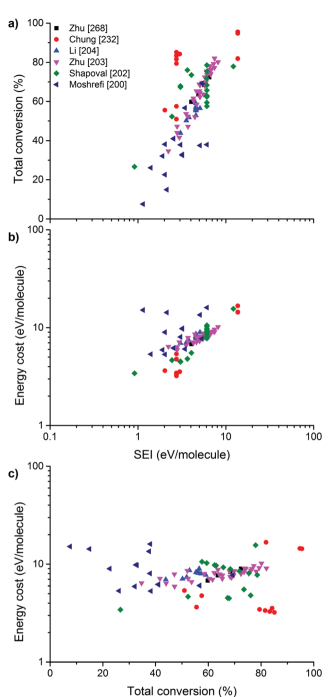


Fig. 29 Experimental data collected from the literature for DRM in a spark discharge, showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

for even larger gap widths, while the selectivities were only slight affected. At increased pressures, i.e. up to 2.5 bar, higher conversions were reported for the same SEI. 203,268 The selectivities again appeared unaffected. When increasing the frequency from 5 to 80 kHz at the same SEI, an increase in the conversions of CH₄ and CO₂ from 67% and 58% to 74% and 63%, respectively, was observed. This might be explained by the relatively

higher density of active species at higher frequencies. The product selectivities, on the other hand, showed only a minor dependence on the frequency.

Just like aforementioned for the GA discharge, for spark discharges, a lot of research has already been performed for the addition of O2 to the mixture to simulate biogas reforming. 204,269-272 The same observations were made regarding a shift from DRM to partial oxidation reactions. 204,269-271 with the suggestion that the water-gas shift reaction is the major or even the only effective contributor to the CO₂ conversion.^{270,271}

Finally, some experiments were also performed for the implementation of catalysts in spark discharges. Chung et al. 199 used BaZr_{0.05}Ti_{0.95}O₃ (BZT), with a perovskite structure and ferroelectric properties, as a packing material inside the spark discharge zone. Compared to the empty reactor, the CH₄ and CO₂ conversions increased from 53% and 49% to 84% and 77%, respectively. This indicated that the combination induced synergistic effects to reduce the specific energy cost. Two types of packing were tested: coarse (C-BZT) and fine (F-BZT) particles. C-BZT showed a better performance than C-BZT, possibly because the void space was larger in the former, leading to a higher electron density.

In summary, spark discharges allow a high conversion, i.e. up to 85%, while demonstrating low energy costs, i.e. 3-10 eV per molecule. At the same time, they can achieve high selectivities towards syngas, which is interesting for the indirect oxidative liquefaction. The direct formation of oxygenates has, however, not been observed up to this point.

5.2.6. Atmospheric pressure glow discharges (APGDs). The APGD shows quite some similarities with DBD and corona discharges. However, to date, the application of this discharge type for DRM is rather limited. 206,207,217,219,220 Nevertheless, it has some distinctive properties, which make it more suitable than its two companions, such as its high electron density and proper plasma temperature for vibrational excitation. The highest obtained conversions of CH₄ and CO₂ are 99% and 90%, while the main products are H2 and CO, with a ratio that can easily be modulated with the CH₄/CO₂ ratio.²¹⁹ Fig. 30 summarizes the data available in the literature, with the best result achieving a total conversion of 89% at an energy cost of only 1.2 eV per molecule. 219 Both the conversion and energy cost show a clear increasing trend with the SEI.

Long et al. 220 added N2 to the CH4/CO2 mixture to generate a stable discharge, but increasing the N2 flow rate led to a decrease in the CH₄ and CO₂ conversions from 46% and 34% to 37% and 22%, respectively, while the H₂ and CO selectivity only changed slightly.

Furthermore, Long et al. 220 combined the APGD plasma jet with a post-discharge catalyst bed. A γ-Al₂O₃ carrier and 12 wt% $Ni/\gamma-Al_2O_3$ catalyst were used. The performance for the $\gamma-Al_2O_3$ was similar to the plasma-only results, while the Ni/γ-Al₂O₃ catalyst significantly enhanced the process. The conversion of CH₄ and CO₂ increased by 14% and 6%, respectively, while the yields of H₂ and CO increased by 18% and 11%, respectively. The single pass conversion was, however, too low and N2 was needed to maintain a stable discharge.220

In summary, the APGD seems to be promising for DRM, based on its proper plasma temperature for vibrational

excitation and its high electron density, as it has a higher conversion ability compared to DBD and corona discharges, i.e. conversions up to 90% are reported for energy costs as low as 1 eV per molecule. Nevertheless, more research will be needed to fully demonstrate the capabilities. One of the major challenges remains a further enlargement of the process to treat large volumes.

5.2.7. Nanosecond pulsed discharges. Nanosecond pulsed discharges are another relatively new alternative used for DRM, for which relatively high energy efficiencies, or low energy costs, have been reported, compared to other discharge types. All the data available in the literature are plotted in Fig. 31. Total conversions in the 40-60% range have been achieved, with

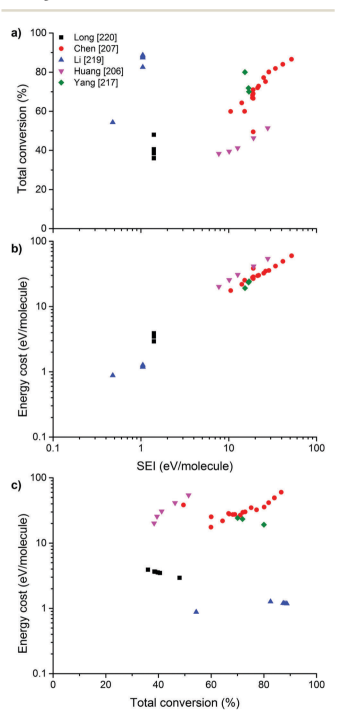


Fig. 30 Experimental data collected from the literature for DRM in an APGD, showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

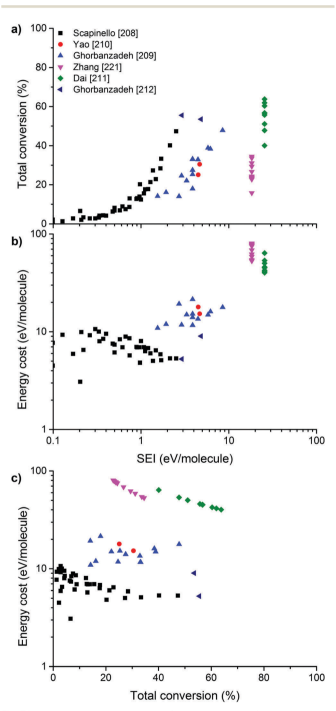


Fig. 31 Experimental data collected from the literature for DRM in nanosecond pulsed discharges, showing the conversion (a) and energy cost (b) as a function of the SEI, as well as the energy cost as a function of the conversion (c). Note that some of the data have been recalculated from the original references to take, among others, dilution effects into account.

energy costs around 3-10 eV per converted molecule. 208,212 The best results comprise an energy cost of only 5.3 eV per molecule for a total conversion of 56%, 212 or 47%. The most abundant hydrocarbons formed are C₂H₂ (and C₂H₄). This is different from the results for DBD where C₂H₆ is mostly formed. Hence, the product distribution, with a prevalence for unsaturated hydrocarbons, in nanosecond pulsed discharges seems to be more similar to GA or spark discharges. 208 Work in this area has been performed on pulsed coronas with a pulse width of both 100 ns²⁷³ and 330 ns,^{211,221} and on pulsed glow-arc discharges with a typical pulse width in the range of 20-70 ns. 208-210,212

In this type of discharge, the amount of power dissipated in the plasma is determined by the pulse voltage and pulse frequency. 209-212 A higher pulse voltage or repetition frequency leads to a higher power, which hence increases the SEI and conversions. The influence of the pulse voltage seems to be stronger than that of the pulse frequency. 211 Regarding product selectivity, increasing the pulse frequency gives rise to a higher C₂H₄ selectivity and a decrease in C₂H₂ selectivity. ^{210,211}

Zhang et al.221 investigated the effects of adding various catalysts to the discharge and observed significant effects. Pure γ-Al₂O₃ significantly increases the CH₄ conversion, while both the CO₂ conversion and C₂ selectivity decrease. La₂O₃/γ-Al₂O₃ catalysts, in contrast, give lower CH₄ conversions and higher C₂ selectivities, i.e. more than 60%. Pd/γ-Al₂O₃ and Pd-La₂O₃/ γ -Al₂O₃ catalysts show the same results as pure γ -Al₂O₃ and La₂O₃/γ-Al₂O₃, respectively, but the C₂ product distribution shifts from C₂H₂ to C₂H₄.²²¹

In summary, these nanosecond pulsed discharges allow a high conversion, i.e. of up to 50-60%, while demonstrating relatively low energy costs, i.e. 3-10 eV per molecule. In previous sections, pulsed power DBD and corona discharges were mentioned, operating in the microsecond pulse regime, but it is clear that the latter did not attain the high conversions and energy efficiencies of nanosecond pulsed discharges. This leads us to believe that the nanosecond timescale is essential to create the necessary strong non-equilibrium, 209 as is also the case in the filaments of a DBD.

5.2.8. Summary. It is clear that DRM has already been explored more extensively than pure CO₂ splitting, and several promising results have already been obtained for various types of plasmas. On the other hand, the DRM process is clearly more complex than pure CO₂ splitting. The CH₄/CO₂ ratio plays an extremely important role, not only in the conversion rates and energy cost, but even more importantly in the product distributions and syngas ratio. A balance appears to be required, since at high CH₄/CO₂ ratios carbon deposition can cause detrimental effects, while at low ratios the H atoms are lost to H₂O, and next to H₂ formation. Several discharges also exhibit high selectivities towards syngas, which is beneficial for the indirect oxidative pathway, while others might be more interesting in the long run due to their suitability for a direct oxidative pathway, in combination with catalysts. 112

To summarize, in Fig. 32 we plot the energy cost as a function of the total conversion, grouped per discharge type, for all the data discussed above. Furthermore, both the thermal equilibrium limit (see Section 2.2.1; Fig. 3) and the target energy cost of 4.27 eV per molecule for the production of syngas (see the beginning of Section 5.2; indicated as "Efficiency target") are displayed. Note that the y-axis is reversed, to allow a better comparison with Fig. 24 for pure CO₂ splitting, where we plotted the energy efficiency. This figure allows us to draw the following conclusions.

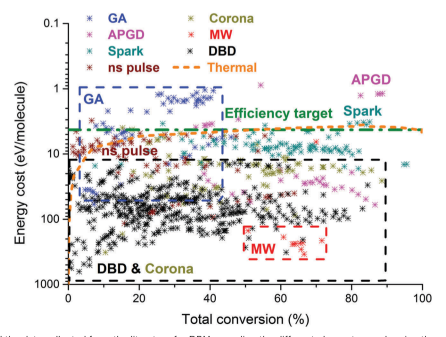


Fig. 32 Comparison of all the data collected from the literature for DRM regarding the different plasma types, showing the energy cost as a function of the conversion. The thermal equilibrium limit and the target energy cost of 4.27 eV per molecule for the production of syngas (corresponding to a 60% efficiency target) are also indicated. Note that the y-axis has been reversed for easy comparison with Fig. 24.

First of all, it is clear that for DRM - just like for CO₂ splitting - although DBDs are by far the most extensively studied, they do not allow the energy-efficient conversion of CO₂ and CH₄ into syngas. Their energy cost currently remains a factor of 5 too high, even when combined with a packing, and we do not expect this to be sufficiently improved in the near future for them to become industrially competitive. On the other hand, if suitable catalysts could be found that would allow the direct high yield production of valuable oxygenates, this would change the analysis drastically, due to the more favourable energy requirements for the one-step process, as explained above. The same conclusion applies to most studies on corona discharges and the several studies on APGDs, with some exceptions as mentioned below.

Second, there is a significant lack of data with MW plasmas on DRM compared to pure CO₂. Research in this area is highly recommended to evaluate whether the potential of MW discharges for DRM is as high as for pure CO2 splitting.

Third, the best results for GA plasmas are easily capable of surpassing the energy efficiency (or energy cost) target set for DRM. Even for conversions up to 40%, the energy efficiency target is easily crossed and they clearly operate in a strong non-equilibrium regime, as evidenced when compared to the thermal equilibrium limit. Moreover, further improvements can still be expected from this discharge type, as the full capabilities of the novel type of GAP, which has many advantages compared to classical GA discharges (see Sections 4.2.3 and 5.1.3 above), have not vet been explored for DRM.

Fourth, although the use of spark discharges and nanosecond pulsed discharges is relatively new and studies are still scarce, energy costs in the range of 10 eV per molecule are reported, and even as low as 3 eV per molecule, thus already reaching the efficiency target, even with the limited amount of research performed to date. For the nanosecond pulsed discharges, conversions are currently limited to 40-50% at these energy costs, while for spark discharges, conversions up to 85-95% have already been achieved. This clearly shows the potential for this type of discharge, and more research should be performed to further exploit their possibilities.

Finally, as mentioned above, for APGDs, some results obtained in the literature seem to be very promising. Indeed, as appears from Fig. 32, the best overall results were clearly achieved for APGDs, which could reach a high conversion, such as 88% for the spark discharge, in combination with low energy costs, such as 1.2 eV per molecule for the GA discharges. Thus, at this stage, APGDs together with GA discharges seem to be the most powerful, providing significant conversions at high energy efficiencies for DRM.

5.3. $CO_2 + H_2O$: artificial photosynthesis

As is clear from the sections above, to date a lot of research is based on both pure CO₂ splitting and the dry reforming of methane. There has also been some research on pure H₂O splitting with plasmas, i.e. for the production of hydrogen. 21,274-278 However, research on the simultaneous conversion of CO2 and H2O - or so-called artificial photosynthesis - into syngas or oxygenated

products by plasma is rather limited. The different plasma set-ups used to date for combined CO2/H2O conversion are: a regular DBD,^{279,280} ferroelectric packed-bed DBD,²⁷⁹ DBD packed with Ni/γAl₂O₃, ²⁸¹ MW discharge, ^{282–285} GA, ^{100,162} surface discharge²⁸⁶ and a negative DC corona discharge.²⁸⁷

5.3.1. DBD plasmas. Futamura *et al.*²⁷⁹ investigated two different plasma reactor types: a ferroelectric packed-bed reactor (using BaTiO₃ pellets) and a silent discharge reactor, a.k.a. a DBD. For both reactors, the CO₂/H₂O mixture was diluted to 0.5-2.5% in N₂. For the DBD reactor, a CO₂ conversion of only 0.5% was obtained, with product yields of 0.7% for H₂, 0.5% for CO and 0% for O2, but no oxygenated compounds were mentioned. The packed-bed reactor was much more successful and a CO2 conversion of 12.3% was reached, with product yields of 12.4% for H₂, 11.8% for CO and 2.8% for O₂, but again, nothing was mentioned concerning the formation of oxygenates.

Mahammadunnisa et al. 281 investigated the effect of a DBD reactor packed with a Ni/γ-Al₂O₃ catalyst, in both its unreduced and partially reduced forms, on combined CO₂/H₂O conversion. For the measurements without a catalyst, a CO2 conversion of 12-25% was obtained, depending on the SEI. The higher the power or the longer the residence time, the higher the conversion was. The reported selectivities were 18-14% for H2 and 97-99% for CO, leading to a syngas ratio of 0.55-0.18. When adding the catalysts, the conversion and syngas ratio increased, to 18-28% and 0.95-0.45, respectively, for the unreduced catalyst (NiO/ γ -Al₂O₃), and to 24-36% and 0.66-0.35, respectively, for the partially reduced catalyst (Ni/γ-Al₂O₃). In this case, the added catalyst leads to a combination of physical and chemical effects, since, beside the enhanced conversion, the NiO catalyst is responsible for a further reduction of the produced CO to CH₄ (and CH₃OH), as well as the other compounds that were detected, i.e. C2H2, propadiene and carbon nanofibres of the partially reduced catalyst. Furthermore, it was concluded that higher flow rates lead to a higher H₂/CO ratio, and thus to more economical syngas production.

Snoeckx et al.280 recently performed a combined experimental and computational study for CO2/H2O conversion in a DBD. CO₂ and water vapour were used as the feed gases, with varying the H₂O content in the mixture between 0% and 8% for three different SEI values. It was demonstrated that adding a few % of water to a CO₂ plasma leads to a steep drop in the CO₂ conversion, and then both the CO₂ and H₂O conversion keep on decreasing slightly when adding more water. The main products formed were CO, H2 and O2, as well as H2O2 - up to 2% for high SEI values and water contents. The combination of the experiments with a computational chemical kinetics study allowed the researchers to analyse the chemical kinetics and to construct and investigate the different chemical pathways to clarify the experimental results. In general, it was concluded that for a DBD, the main reactive species created are OH, CO, O and H, of which the OH radicals will quickly recombine with CO into CO₂, thereby limiting the CO₂ conversion upon the addition of water. At the same time, the O and H atoms will undergo subsequent reactions to form H₂O again, thus explaining why the H₂O conversion is also limited. Furthermore, the fast

reaction between O/OH and H atoms also explains why no oxygenated products were formed, because it occurs much faster than the possible pathways that might otherwise lead to oxygenates.²⁸⁰

5.3.2. MW plasmas. Ihara *et al.*^{282,283} were the first to investigate the combined CO_2 and H_2O conversion. In their first study²⁸² a 1:1 mixture of CO_2/H_2O was investigated for a MW plasma. They detected low yields of oxalic acid and H_2O_2 in the cold trap condensate by reversed-phase chromatography using UV and conductivity detectors. The H_2O_2 production was very dependent on the discharge power, and a maximum yield of 0.024% was obtained. Furthermore, they assumed that H_2 and O_2 are generated, although these products were not measured. Interestingly, they also found the deposition of a transparent solid crystal film on the reactor walls, corresponding to oxalic acid.

In their follow-up study, 283 they used the same MW set-up, but alternative detection techniques, i.e. steam chromatography and gas chromatography-mass spectrometry (GC-MS). The CO₂/H₂O gas mixture was varied from a 1:4 to a 1:1 ratio, but the full conditions for the presented results were not mentioned. This time, methanol formation, instead of H₂O₂ and oxalic acid, was reported, albeit in very low concentrations < 0.01%, both in the effluent stream and in a transparent solid crystal film that was deposited on the reactor walls. Therefore, the authors concluded that two pathways for methanol formation could be considered: the direct formation from CO2 and H2O in the plasma and the reformation of deposited polymeric material on the walls during the plasma reaction with H2O. Most importantly, they observed that the methanol yield increased by a factor of 3.5 when increasing the pressure from 240 to 400 Pa, and so they stated that the system pressure is one of the most important parameters to consider.

Chen et al. 284 applied a surface-wave MW discharge for the simultaneous dissociation of CO_2 and $\mathrm{H}_2\mathrm{O}$. The formation of syngas and O_2 was observed, but no hydrocarbons or oxygenates were detected. The influences of the gas mixing ratio, the SEI and the feed flow rate on the H_2 and CO production were studied. It was found that syngas with a ratio close to 1 can be produced when the $\mathrm{CO}_2/\mathrm{H}_2\mathrm{O}$ ratio in the gas mixture is 1:1.

In a follow-up study, 285 the authors combined their MW setup with NiO/TiO2 catalysts. In this work, they reported an increase in CO₂ conversion from 23% to 31%, along with a lower energy cost from 30.2 to 22.4 eV per molecule, when adding 10% H₂O for the plasma-only conditions. At the same time, a lower gas temperature was observed, which might be due to the higher heat capacity of water and the induced endothermic reactions. In turn, this lower temperature might be responsible for the higher conversion, since a lower gas temperature in MW plasmas is beneficial for energy-efficient CO₂ conversion, due to there being lower vibrational losses by VT relaxation (see Section 5.1.2 above) and a reduced backward reaction rate, i.e. the recombination of CO back into CO₂. 88,89 When adding a NiO/TiO₂ catalyst treated with an Ar plasma, the CO₂ conversion further increased to 48%, with an energy cost of 14.5 eV per molecule. Still though, no oxygenated products were detected. It was suggested that CO2 is adsorbed at an oxygen vacancy on the catalyst surface, reducing the threshold for the

dissociative electron attachment process to 2 eV, creating CO, an adsorbed O atom at the vacancy and an electron. Subsequently, the adsorbed O atoms interact with OH to form O_2 and H as well as with gas phase O atoms to recombine to O_2 . As such, this seems to be some evidence that the reactions and species limiting the CO_2 and H_2O conversion, as described in the chemical reaction pathways for a pure DBD plasma²⁸⁰ (see previous section), can indeed, as suggested, be hijacked by the implementation of a catalyst.

5.3.3. GA plasmas. As mentioned in Section 5.1.3 on pure CO2 splitting, Indarto et al. 162 and Nunnally et al. 100 applied a GA to investigate the conversion and energy efficiency of CO₂. The former investigated the addition of water vapour in the range of 5% to 31% for a classical GA. A decrease in CO₂ conversion (from 7.1% to 3.0%) and an increase in energy cost (from 89 to 189 eV per molecule) was observed over this range compared to 13.4% and 53 eV per molecule for pure CO₂. One of the suggested reasons for this was the increased instability of the GA with the increasing water vapour concentration. Nunnally et al. 100 investigated the effect of adding 1% water vapour for a GAP, and showed an increase in the energy cost from 9.5 to 14.8 eV per molecule. While Indarto et al. 162 attributed this to arc instabilities, no instabilities were observed for the GAP. Therefore, it was concluded that this higher energy cost is due to vibrational energy losses through VT relaxation. This process is relatively slow for CO₂ molecules, ^{21,100} leading to the high energy efficiency of pure CO₂ splitting in a GA and MW plasma. However, for H₂O this VT relaxation is much faster, ^{21,100} hence it is believed that water absorbs part of the vibrational excitation energy of CO₂ and subsequently loses this energy quickly through VT relaxation, leading to the observed drop in energy efficiency, or vice versa, the rise in energy cost.21,100

5.3.4. Other plasma types. Another discharge type, namely surface discharge, was applied for both a 1:1 mixture of CO_2/H_2 and CO_2/H_2O in a comparative study by Hayashi *et al.*²⁸⁶ The CO_2 conversions achieved were 15% and 5%, respectively. In both cases, the major products were the same, *i.e.* CO, CH_4 , dimethyl ether (DME) and formic acid, and in the case of H_2 as co-reactant, also water formation.

Guo et al.²⁸⁷ reported the use of a negative DC corona discharge for the reaction between CO₂ and H₂O, varying the water vapour content between 10% and 43%, and the pressure between 1 and 4 bar. They observed a drop in CO₂ conversion with increasing gas flow rates (hence decreasing SEI) and increasing water content. The main products formed were ethanol and methanol, in roughly a 3:1 ratio, with a total molar yield of up to 4.7% and CO₂ conversion reaching a maximum of 16% at 1 bar, 50 mL min⁻¹ and 23% water vapour. Other compounds detected were H₂ and CO. Increasing the pressure had a beneficial effect on the methanol and ethanol yields. For example, when going from 1 to 4 bar, the ethanol yield increased from 3.2% to 11.9%. Unfortunately, some of the reported results seem to be contradictory/inconsistent, making it difficult to interpret the obtained results.

5.3.5. Summary. In general, from all the data presented in the available literature, we may conclude that, for plasma-only cases, the addition of even small amounts of water (1-2%) leads

to a significantly lower CO₂ conversion. This declining trend continues upon the addition of even more water, albeit less severe. As a result, the addition of water also leads to higher energy costs. For a classical DBD reactor, the energy cost is already quite limited, thus making it unsuitable for the combined CO₂/H₂O conversion. As shown in Sections 5.1 and 5.2 above, MW and GA plasmas are far more efficient. Moreover, they operate at somewhat higher temperatures, i.e. in the order of 1000 K or more, which enables the addition of more H₂O vapour. Nevertheless, it has been suggested that H2O might quench the vibrational levels of CO₂, 100,162 thus reducing the most energy-efficient conversion process, and therefore raising the energy cost. Both GA studies reported in the literature, i.e. those by Nunnally et al. 100 and Indarto et al., 162 seem to confirm this hypothesis, while the MW results from Chen et al. 285 seem to contradict it. However, the MW set-up of Chen et al. might be less prone to this quenching mechanism, since it operates at low pressures (30-60 Torr). Furthermore, the presence of water in the low pressure MW case also seems to lead to a cooling effect, resulting in a higher energy efficiency. Nevertheless, the low pressure operation is still less ideal for industrial implementation. Fig. 33 summarizes the energy cost per converted CO2 molecule as a function of the conversion, for those discharges where data were available.

The main products formed are again H2 and CO, like in the case of DRM (see Section 5.2), as well as O2, but some papers also report the production of hydrogen peroxide (H₂O₂), ^{280,282} oxalic acid (C₂H₂O₄), ²⁸² formic acid (CH₂O₂), ²⁸⁶ methane (CH₄), ^{281,286} dimethyl ether (C₂H₆O, DME), ²⁸⁶ methanol (CH₃OH), ^{281,283,287} ethanol (C_2H_5OH) , 287 acetylene (C_2H_2) , 281 propadiene $(C_3H_4)^{281}$ and even carbon nanofibres (CNF). 281 Unfortunately, most data on the formation of these products are only qualitative and mainly incomplete, making it impossible to deduce a general trend on product yields or selectivities. Although research on

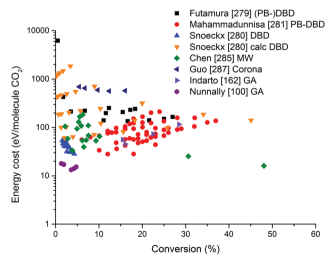


Fig. 33 Comparison of all the data collected from the literature for the artificial photosynthesis ($CO_2 + H_2O$) in the different plasma types, showing the energy cost per converted CO2 molecule as a function of the conversion. Note, some of the data have been recalculated from the original references to take, among others, dilution effects into account.

this process is more limited than for DRM - and hence the road ahead is longer - this clearly indicates that, just like for DRM above, the plasmachemical conversion of CO2 with H2O as a co-reactant allows for the development of the highly desirable direct oxidative pathway.

It is clear from the results obtained in the literature that CO₂ and H2O seem to be unsuitable to create oxygenated hydrocarbons in a one-step process by means of a pure non-thermal plasma, i.e. in the absence of a catalyst, with the exception of the negative DC corona discharge. There are too many steps involved in generating these oxygenates, such as CH₃OH, in an efficient way, and all of them involve H atoms, which will have the tendency to quickly recombine with OH into H₂O or with O₂ into HO2, which also reacts further with O into OH, and hence H₂O. The problem at hand is thus that the interactions of H atoms with oxygen species (either OH, O3, O2 or O atoms) are too fast and their tendency to form H₂O is too strong.

On the other hand, CO₂/H₂O plasmas can deliver an easily controllable H₂/CO ratio with a rich hydrogen content – even up to 8.6 - when sufficient amounts of water can be added to the CO₂ plasma, as was demonstrated in the computational study by Snoeckx et al.280 Hence, they might be suitable to create value-added chemicals, such as oxygenates, in a two-step process, which is good news. However, the interaction between H₂O and CO₂ dissociation products, i.e. the recombination between OH and CO into CO2, and the recombination of H and OH into H2O, limit the CO2 and H2O conversion, and thus the formation of useful products. Besides syngas, the direct production of sufficient amounts of hydrogen peroxide, which can be used as a disinfectant or for biomedical purposes, seems possible. However, the formation rate of H_2O_2 is also partially limited by the destruction reaction of OH + H₂O₂ towards H₂O and HO2. Therefore, rapid removal of the formed product (i.e. H₂O₂), e.g. by means of a membrane, would be an important aspect for further improving this process. Based on these findings, and the work by Mahammadunnisa et al.,281 it is evident that a CO2/H2O plasma should be combined with a catalyst in order to produce value-added chemicals. 102,107

As such, it appears that, due to the inherent nature of this mixture and the plasma set-ups, the future for the combined CO₂ and H₂O conversion by pure plasma technology does not look so bright at the moment. Nevertheless, we believe that a possible promising way forward is by its combination with specific tailored catalysts to produce value-added chemicals.

Such catalysts should be able to selectively let the plasmagenerated CO and H₂ react into oxygenates, such as methanol, and subsequently allow separation of the methanol from the mixture. As mentioned in the next section, Eliasson et al.²⁸⁸ applied a CuO/ZnO/Al₂O₃ catalyst in a CO₂/H₂ discharge, which resulted in an increase in methanol yield and selectivity by more than a factor of 10. Several other reported catalysts used already for CO2 conversion with H2 in traditional thermal catalysis might also be interesting to investigate for their suitability in plasma-catalysis in a CO2/H2O mixture, such as Ni-zeolite catalysts, for which methanation is reported, 289 a Rh₁₀/Se catalyst, yielding an ethanol selectivity of up to 83%, 11

and a Ni–Ga catalyst for the conversion into methanol.²⁹⁰ Moreover, a lot of research into catalytic CO₂ hydrogenation is showing promising results for CuO/ZnO/ZrO₂ and for Cu/ZnO-based catalysts promoted with Pd and Ga, as well as Pd/ZnO and Pd/SiO₂ with the addition of Ga.⁵¹ In general, multicomponent systems (Cu/ZnO/ZrO₂/Al₂O₃/SiO₂) have reported good performances for the formation of methanol starting from CO/CO₂/H₂ mixtures,¹¹ making them potentially also very interesting for plasma-catalysis of a CO₂/H₂O mixture (as well as for DRM), because the CO/CO₂/H₂ mixture is anyway generated *in situ* during plasma-based conversion.

Based on the chemical kinetics pathway analysis presented by Snoeckx *et al.*²⁸⁰ (see Section 5.3.1 above), two pathways might be interesting and realistic to achieve: (i) promoting the recombination of OH radicals to H_2O_2 or (ii) promoting the reduction of CO to methanol. In both cases, the thermodynamic aspects at the nanoscale will become very important, especially since plasma-catalysis is a far-from-equilibrium process.¹⁰⁷ The critical point will be the arrival and binding (*e.g.* physi- or chemisorption) of the reactants to the catalyst surface. To be successful, this process should be faster than the recombination rate of OH with H into H_2O . Of course, these suggestions are only speculations, and further research will be needed to investigate these possibilities in practice.

We need to give a final critical note concerning safety. We need to be cautious about the explosive mixture that can be formed during this process - which could of course also occur for some of the other novel technologies discussed in Section 3, when O2 and H2 are not produced separately - due to the presence of O2, together with CO, H2 and an ignition source in a plasma set-up. At the research level, this will probably never be a problem due to the low volumes and conversions. However, when going to a pilot or industrial scale, i.e. with larger volumes and conversions, the risk will increase significantly. Consequently, both the capital and operating costs will increase drastically to ensure safe operations. One way to circumvent this problem is by diluting this mixture with an inert gas, such as argon or helium. In this case, however, an additional separation (for the products) and recuperation (for the inert gas) step will need to be included, which will also increase the cost. Furthermore, part of the input energy will be lost due to the electron-impact excitation and ionization of these gases. Therefore, this option will reduce the energy efficiency and increase the operating cost, but it will ensure safe operation.

5.4. CO₂ + H₂: hydrogenation of CO₂

Research on the plasma-based hydrogenation of CO_2 – using H_2 as a co-reactant – is as limited as the combined conversion of CO_2 and H_2O . Historically, this is in part due to the high cost of hydrogen. Like for multiple other novel technologies discussed in Section 3, the sustainable and economically viable production of H_2 is indispensable for pathways relying on the use of H_2 as a co-reactant. Although interest into the plasma-based hydrogenation of CO_2 emerged around the same time as the use of H_2O as a co-reactant, most of the work performed is actually very recent. The different plasma set-ups used to date:

are a DBD, ^{288,291} packed-bed DBD, ^{288,292} MW discharge, ^{285,293,294} RF discharge²⁹⁵ and a surface discharge. ²⁸⁶

5.4.1. DBD plasmas. Eliasson et al. 288 investigated the hydrogenation of CO2 to methanol in a DBD, both with and without the presence of a CuO/ZnO/Al2O3 catalyst. The experiments were performed at a gas pressure of 8 bar and a H₂/CO ratio of 3:1. For the plasma-only case, the major products found were CO and H₂O, which is not surprising keeping the results of CO₂/H₂O in mind (see the previous section). Other components detected were CH₄ and methanol, with a selectivity of only 3-4% and 0.4-0.5%, respectively. By adding the catalyst, the methanol yield increased about 10 times and the selectivity was 10-20 times higher. Furthermore, by optimizing the system to use low power and high pressure, it became possible to further enhance the methanol selectivity over the CH4 selectivity. The results indicate that the discharge shifts the region of maximum catalyst activity from 220 °C to 100 °C. Nevertheless, the electric power used is considered to be prohibitive for methanol production on an industrial scale, due to the low yield, i.e. $\sim 1\%$.

The methanation of CO2 in a DBD packed with Ni/zeolite pellets was investigated by Jwa et al.289 Conventional and plasma-assisted catalytic hydrogenation were compared with a varying nickel loading for a temperature range of 180-360 °C for a stoichiometric 4:1 ratio of H₂/CO₂. For the conventional catalytic hydrogenation case, a conversion of 96% was observed at 360 °C, while for the plasma-assisted hydrogenation, the same conversion was already reached at 260 °C. It was assumed that the reactive species generated in the plasma reactor could speed up the rate determining step of the catalytic hydrogenation. The hydrogenation of CO₂ involves the dissociation of CO_2 to CO and O on the active site of the Ni/zeolite ($CO_{2.ads} \rightarrow$ CO_{ads} + O_{ads}). The rate determining step in the CO₂ conversion into CH₄ would be the same as for CO conversion (CO_{ads} → C_{ads} + O_{ads}). Subsequently, the dissociated species react with hydrogen to produce CH₄. This increased methanation rate is believed to arise from an increase in the surface concentration of carbon. The plasma can help to dissociate the adsorbed molecules, and hence surface carbon can be produced by both thermal activation and plasma activation, ultimately resulting in a higher methanation rate. Without Ni (bare zeolite), the conversion of CO and CO2 was found to be less than 1%, implying that the zeolite support together with the plasma cannot efficiently convert CO2 into CH4.289

Nizio *et al.*²⁹² investigated the effect of a packed-bed DBD with Ni-Ce_xZr_{1-x}O₂ catalysts on the hydrogenation of CO₂ for a stoichiometric H₂/CO₂ ratio of 4:1. At 90 °C without a catalyst, the CO₂ conversion was around 5%, without any methanation taking place. For the same conditions, 78% CO₂ conversion and methanation with a selectivity of 99% and an energy cost < 3 eV per molecule CH₄ was achieved when adding the catalyst. This demonstrated the activity of plasma-catalytic systems at low temperature ($T_{\rm g}$ < 260 °C) and the possibility of enhancing both the conversion and selectivity of the process under study, by combining plasma with catalysts. Furthermore, the plasma-catalytic system showed almost no difference in activity for the

different ceria-zirconia supports, while in the absence of plasma this was the case. The latter clearly indicates that it is indeed the plasma that initiates the methanation process by dissociating adsorbed CO₂ molecules - a reaction that does not take place catalytically below 250 °C.

Recently, Zeng et al. 291 investigated the performance of CO2 hydrogenation in a DBD at atmospheric pressure and low temperature, both with and without a catalyst, i.e. Cu/γ-Al₂O₃, $Mn/\gamma-Al_2O_3$ and Cu-Mn/ $\gamma-Al_2O_3$. Some plasma-only results were also presented. The CO₂ conversion was found to be 7.5%, and the main products were CO, H₂O and CH₄. The CO and CH₄ selectivity were 46% and 8%, respectively. No methanol production was reported, although this might have been due to the inability to detect this compound by the GC set-up used. Adding the catalysts enhanced the process: the CO₂ conversion increased to 8-10%, while the CH₄ selectivity remained around 6.9-8.6%, but the CO selectivity and yield were enhanced to 76-80% and 6.4-7.9%, respectively.

Finally, de Bie et al. 296 performed an extensive computational study on the hydrogenation of CO2 in a DBD, using a onedimensional (1D) fluid model. The H₂/CO₂ mixing ratio was varied in the entire range from 1:9 to 9:1. The most abundant products predicted by the model were CO, H₂O and CH₄, and to a lower extent also formaldehyde, C₂H₆, O₂ and methanol. The CO₂ conversion was found to be rather low (2-7% in all the gas mixing ratios), especially when compared to typical values found for DRM (3-20%) at comparable conditions. This was thought to be the result of the lack of high enough concentrations of CH2 and CH3 radicals, which aid in the CO2 conversion. Moreover, a very similar chemical behaviour as described above for the CO₂/H₂O mixture²⁸⁰ was reported. Indeed, the CO₂ conversion was limited due to the formation of CHO (CO + H + M \rightarrow CHO + M), which reacted back to CO_2 (CHO + O \rightarrow CO₂ + H). Furthermore, a lot of subsequent reactions were needed to form the desired hydrocarbons or oxygenates, such as CH4 and methanol, making their overall production rates negligible. Hence, it was concluded that a CO₂/H₂ mixture in a DBD-only set-up is not suitable for the production of value-added chemicals.

5.4.2. MW and RF plasmas. Maya²⁹³ employed a MW discharge to explore the possibility of obtaining formic acid from a H₂/CO₂ mixture. The pressure in the reactor was, however, very low, i.e. about 1-2 Torr. The main products observed were CO and water, with some secondary products when the H₂/CO₂ ratio exceeded 1:1, including acetylene, methane, methanol, ethylene, formaldehyde and formic acid.

A so-called RF impulse discharge was used by Kano et al. 295 to study the combined H₂/CO₂ conversion into CH₄ and methanol at low gas pressures (1-10 Torr). For a H₂/CO₂ mixing ratio of 4:1, the main products detected were CO and H₂O, as well as CH₄ and to a lesser extent, i.e. one order of magnitude lower, methanol. Formaldehyde and formic acid were not observed. When the repetition frequency was lowered from 60 to 10 kHz, the formation of products decreased as well, while the CH₄ production almost disappeared. The maximum CO₂ conversion and CH₄ selectivity obtained were 26% and 21%, respectively. The most efficient production of CH₄ took place for a mixing

ratio of 6:1 H₂/CO₂, which was larger than the stoichiometric ratio (4:1).

De la Fuente et al. 294 recently used a surface-wave MW plasma reactor without a catalyst to study the effect of the gas flow rate, H2/CO2 mixing ratio and SEI on the performance for CO₂ hydrogenation. The main products found were CO and H₂O as well as 200 ppm C₂H₄ and 10-20 ppm methanol, but remarkably no CH₄ was detected. The best performance was obtained for a H₂/CO₂ mixture of 3:1, obtaining a CO₂ conversion of 82% and an energy cost of 28 eV per converted CO2 molecule, which are the best values reported in the literature for plasma-only operation. The CO₂ conversion was even higher than for pure CO₂ splitting, which was 65%, with an energy cost of 35 eV per molecule. It was predicted by means of a zero-dimensional (0D) reactor model that the dominant intermediate species are H and O, which was also found by OES measurements. The higher conversion of CO2 when adding H2 was thought to be the result of the slightly lower ionization energy of H atoms, resulting in higher electron densities. Furthermore, as these atoms cannot be excited vibrationally and/or rotationally, it might lead to higher electron temperatures.²⁹⁴

Chen et al.285 added a Ni/TiO2 catalyst to a surface-wave MW plasma, to not only investigate a CO2/H2O mixture (see Section 5.3.2 above), but also a H_2/CO_2 mixture in a ratio of 1:9. For the plasma-only case, a reduction in the CO2 conversion from 23% to 14% occurred. When adding the catalyst, the conversion was enhanced by a factor of 2, up to 28%, which was, however, still lower than the value for the pure CO₂ case (41%), and also no methanol or CH₄ formation was observed.

5.4.3. Other plasma types. As also mentioned in Section 5.3 above, Hayashi et al. 286 investigated both the effects of H2O and H₂ as an additive gas for the plasma-based conversion of CO₂ using surface discharge. A 1:1 mixing ratio was used at atmospheric pressure. The products observed were again CO and H₂O as the main components and additionally CH₄, DME and formic acid. For the highest SEI conditions, the maximum CO₂ conversion was approximately 15%. Although higher than for the CO₂/H₂O case, these conversions and yields are considered to be too low.

5.4.4. Summary. It is clear from the above results that the conversions in CO₂ hydrogenation are about a factor of 2-3 lower (and the energy costs the same factor higher) than for DRM and pure CO₂ splitting. Therefore, we may conclude from the limited data available in the literature that the hydrogenation of CO2 for plasma-only cases is not successful. Fig. 34 summarizes the energy cost per converted CO₂ molecule as a function of the conversion for those discharges where data were available.

From all the data presented in the available literature, the main products formed are clearly CO and H2O. Some secondary products are also reported, but always in much smaller, indeed almost negligible, amounts, with the most important ones being CH₄^{286,288,291,293,295,296} and methanol, ^{288,293–296} although some papers also report the production of formaldehyde (CH2O), 293,296 formic acid (CH2O2),286,293 dimethyl ether (C2H6O, DME),286 acetylene(C₂H₂),²⁹³ ethylene (C₂H₄)^{293,294} and ethane (C₂H₆).²⁹⁶

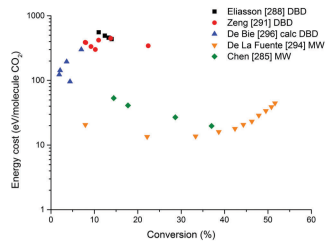


Fig. 34 Comparison of all the data collected from the literature for CO₂ hydrogenation in the different plasma types, showing the energy cost per converted CO₂ molecule as a function of the conversion. Note, some of the data have been recalculated from the original references to take, among others, dilution effects into account.

In spite of these low amounts, the fact that these products can be formed is an indication that the plasmachemical conversion process has the potential for the development of the highly desirable direct oxidative pathway if suitable catalysts could be found.

Thus, CO₂ conversion using H₂ as an additive, without a catalytic packing, shows a high resemblance to the use of H₂O, which is of course not surprising, since the same reactive intermediate species and hence reactions are responsible for the plasma chemistry taking place. In accordance with that observation, in a pure plasma set-up, the combination of CO2 and H₂ also seems to be unsuitable to create methane and/or methanol (or other oxygenated hydrocarbons) in a one-step process. The same number of steps - still too many - are involved in generating these value-added hydrocarbons or oxygenates in an efficient way. All these steps involve H atoms, which will have the same tendency to quickly recombine into OH and subsequently H2O.

Nevertheless, from the limited studies available, it appears that the combination with a catalytic packing is a viable option to produce value-added chemicals in an efficient way. We would like to make the same suggestions as in the previous section, namely that the catalyst should be able to selectively let the plasma-generated C (or CO) and H2 react into methane (or oxygenates). Multicomponent systems (Cu/ZnO/ZrO₂/Al₂O₃/ SiO₂) that show good performance for CO/CO₂/H₂ mixtures¹¹ are potentially very interesting, since this mixture is anyway generated in situ during the plasma-based conversion of CO₂/H₂. Additionally, based on the work of Jwa et al.²⁸⁹ and Nizio et al., 292 the most effective and efficient pathway seems to be a low-power Ni-based plasma-catalytic set-up, in which the plasma takes care of the rate determining step, namely dissociating the adsorbed CO into C, allowing for the catalyst to reduce the C with H to CH₄.

Benchmarking of plasma-based CO₂ conversion against the other (traditional and novel) technologies

Finally, in this section, we compare plasma technology with the traditional catalytic approach and with the other novel approaches for CO2 conversion, based on all aspects explained in the previous sections. Furthermore, we identify the advantages and limitations of plasma-based CO₂ conversion with respect to the other technologies, which will help us to define future priorities for the development of plasma-based CO2 conversion systems.

6.1. Process versatility

A first comparison is made based on the versatility of the different CO2 conversion technologies. Table 1 shows the different technologies and the four different CO2 conversion approaches that were discussed. Traditional thermal catalysis, as discussed in Section 2, can be successfully applied for DRM and the hydrogenation of CO2, but seems unsuccessful at this point for pure CO₂ splitting, while data for the combined conversion of CO₂ and H₂O are unavailable. The novel technologies, discussed in

Table 1 Overview of traditional thermal catalysis and the different emerging technologies, indicating their suitability for the four different CO₂ conversion processes discussed in the previous section. The colour coding gives an additional visual sense for how efficiently this process could be applied when it is achievable, as discussed in more detail in the text: inefficient (red), to be proven (orange) or efficient (green)

	CO ₂ splitting	CO ₂ /CH ₄	CO ₂ /H ₂	CO ₂ /H ₂ O
Catalysis	Х	X	Х	NA
Electrochemical	X	NA	NA	Х
Solar thermochemical	X	NA	NA	X ^a
Photochemical	NA	NA	NA	Х
Biochemical	NA	NA	NA	Xp
Plasmachemical	Х	Х	Xc	Xc

^a CO₂ and H₂O are not converted at the same time, see discussion in Section 3.2. ^b H₂O is a vital nutrient for the growth of the algae, see discussion in Section 3.4. CWhen used in combination with catalysts (plasma-catalysis), see Sections 5.3 and 5.4.

Section 3, are all applicable to converting CO₂ together with H₂O. Furthermore, the electrochemical and solar-thermochemical-based technologies are also capable of pure CO₂ splitting. The combinations with CH₄ and H₂ are not generally reported in the literature. Plasmachemical conversion, on the other hand, currently is the only technology reporting successful application for all four processes, i.e. pure CO2 splitting, DRM, the hydrogenation of CO2 and the combined conversion of CO2 and H2O. As discussed in detail in Section 5, plasma-based pure CO₂ splitting and DRM are obviously the two most successful processes today, with some types of plasmas already surpassing the posed 60% energy efficiency target for syngas production and achieving conversions of up to 90-95%, even without the combination with catalysis. Furthermore, the production of liquids in a one-step process, through a direct oxidative pathway (in combination with catalysts), has already proven successful, with further research in full progress. In this case, avoiding the intermediate syngas step, and thus circumventing the need for an additional Fischer-Tropsch or methanol synthesis and subsequently the methanol/ethanol to olefins synthesis, would significantly reduce the energy efficiency target and make it more competitive (i.e. by at least a factor of 2–3 in the case of methanol). Studies regarding the hydrogenation of CO2 using plasmas in combination with catalysis are very promising as well, but more

research is still required. Although technically possible, the combined conversion of CO2 and H2O - the so-called artificial photosynthesis - with plasma technology seems the least promising from the data available to date. Nonetheless, here as well the combination with catalysis could prove beneficial, if a suitable catalyst could be found, as discussed in Section 5.3.5 above, but to date, research in this area is very scarce.

This high process versatility makes the plasmachemical conversion of CO₂ very interesting as a CCU technique. It allows the process to be used at a wide variety of locations, independent of the available product feed, and even adjustable to a variable product feed at a specific location by the easily adaptable process. This process versatility gives the plasmachemical conversion a substantial benefit over the other technologies.

6.2. Advantages/disadvantages

It is clear that, although all the technologies, regardless of their maturity, have specific or even unique advantages, there is always a flip side to the coin. Table 2 provides a visual overview of the different technologies discussed in Section 3, as well as for plasmachemical conversion, together with a list of some of their key distinctive advantages and disadvantages.

Table 2 Overview of traditional thermal catalysis and the different emerging technologies, indicating their distinctive key advantages and disadvantages. The colour coding gives an additional visual sense for the impact of the feature, as discussed in more detail in the text: negative (red), undesirable/neutral (orange) or positive (green)

	Use of rare earth metals	Renewable energy	Turnkey process	Conversion and yield	Separation step needed	Oxygenated products (e.g. alcohols, acids)	Investment cost	Operating cost	Overall flexibility
Traditional catalysis	Yes	\ <u>-</u>	No	High	Yes	Yes	Low	High	Low
Catalysis by MW-heating		Indirect						Low	Low
Electro- chemical	Yes	Indirect	Nob	High	Yes ^c	Yes	Low	Low	Medium
Solar thermo- chemical	Yes	Direct	NA	High	No	No	High	Low	Low
Photo- chemical	Yes	Direct ^a	Yes	Low	Yes	Yes	Low	Low	Low
Biochemical	No	Direct ^a	No	Medium	Yes ^d	Yes	High /low	High	Low
Plasma- chemical	No	Indirect	Yes	High	Yes ^e	Yes	Low	Low	High

^a Bio- and photochemical processes can also rely on indirect renewable energy when they are coupled with artificial lighting, see Sections 3.3 and 3.4. b Electrochemical cells are turnkey, but generally the cells need to operate at elevated temperatures and the cells are sensitive to on/off fluctuations, see Section 3.1. ^c The need for post-reaction separation for the electrochemical conversion highly depends on the process and cell type used, see Section 3.1. d Biochemical CO2 conversion requires very energy-intensive post-reaction separation and processing steps, see Section 3.4. The need for post-reaction separation for plasma technology highly depends on the process, see Section 5.

From an economical and sustainable point of view, the use of rare earth metals is one of the key disadvantages of most technologies at this point, except for biochemical and plasmachemical conversion. As mentioned before, the grand challenge for the other emerging technologies to be successful (as well as for traditional catalysis) is switching to inexpensive earthabundant metals. It cannot be stressed enough that this is a critical make or break point, but which is not an issue for the plasmachemical conversion.

The discussion regarding the reliance on renewable energy in a direct or indirect manner (see Sections 1, 3 and 4) is more nuanced. The ability of solar thermochemical, photochemical and biochemical conversion to directly utilize renewable energy, i.e. solar radiation, can be considered to give them an edge over the other novel technologies, i.e., catalysis by MW-heating, electrochemical and plasmachemical conversion, which rely on electricity, and hence indirect renewable energy. The former have the advantage of skipping an energy conversion step, which would otherwise always leads to further energy losses, hence the energy efficiency of the latter is intrinsically determined by the efficiency of the renewable electricity generation. On the other hand, this also means that the MW-heating, electrochemical and plasmachemical processes can rely on other renewable energy sources as well, e.g. wind, hydro, wave and tidal power, increasing their versatility and employability, since they can be installed and operated independent of the availability of solar light. This significantly increases the application flexibility of MW-heating and electrochemical as well as plasmachemical conversion. Note that bio- and photochemical technologies may also rely on indirect renewable energy, when used in combination with artificial lighting.

The wide-scale adoption of these renewable energy sources, and their intermittent character, poses a challenge for the efficient storage and easy transport of the electricity produced. Not only is there a need for peak shaving, but more importantly a need for grid stabilization, which requires technologies to follow the irregular and at times intermittent supply of renewable electricity in a flexible manner. Technologies that are able to harness this energy and convert it into carbon neutral fuels could play an important role for the future energy infrastructure industry. Due to the intermittent character of renewable energies, a flexible storage process, which can easily be switched on and off to follow the supply, i.e. a so-called turnkey process, would be most ideal. Photochemical and plasmachemical conversion are the only two processes that truly meet this condition. They can be turned on with the flick of a switch, since no pre-heating or long stabilization times (<30 minutes) are required, as is the case for traditional catalysis. Likewise, they can be turned off with the flick of a switch, since no sensitive cool-down times are required and there is no risk of damaging the reactors with repeated on-off cycles. The problem with electrochemical cells is that, not only do they operate at elevated temperatures, but they are not designed- indeed, they suffer - for repeatedly being turned on and off, while biochemical processes need to be looked after on a continuous basis, e.g. regarding the nutrients, light, temperature, mixing.

From an industrial point of view, high conversions and yields are required, and most technologies are already capable of delivering on these requirements, with the exception of the photo- and biochemical conversion. A closely related key parameter is the solar-to-fuel conversion efficiency, which is discussed separately in the next section (Section 6.3).

Furthermore, when assuming that all or most of the feed is converted, the question arises as to whether an additional postreaction separation step is required for the products. Solar thermochemical and electrochemical conversion (depending on the process and cell type) are the only two technologies capable of generating separated product streams, i.e. separated CO and O₂ streams in the case of pure CO₂ splitting. This is an important advantage, since the separation of CO and O2 is rather energy intensive at this point, although in the future, membranes might bring a convenient solution to this problem. Currently, this is probably the major disadvantage of plasma technology, as the products are all in one feed, i.e. for pure CO₂ splitting, a mixture of CO and O2 is produced, and for DRM a mixture of (mainly) CO and H2, with - highly depending on the discharge type – some (minor) side-products, like hydrocarbons and oxygenates. Of course, a mixture of CO and H2 should not be a big problem, if the mixture will be immediately processed further through Fischer-Tropsch or methanol synthesis, since the plasmachemical conversion is able to deliver any desired syngas ratio, mainly depending on the gas mixing ratio in the feed. Nevertheless, research on the *in situ* trapping of species⁴⁰ or a combination with other technologies²⁹⁷ is highly valuable, and requires more attention.

At the same time, this problem can also become less significant when focusing on the direct conversion of CO₂ into value-added liquid oxygenated products by means of plasmacatalysis, ^{102,103} including alcohols, aldehydes and acids, which are easier to separate. In the long run, as already mentioned before, this direct oxidative pathway to produce more value-added products is highly preferred rather than the current indirect method, which involves first producing CO and H₂, which are then further processed. At the moment, except for solar thermochemical conversion, all the technologies are already able to produce more value-added chemicals in a direct fashion, although the yields are still extremely low and most research is still in its early stages.

The investment and operating costs are in general considered to be low for most technologies. The main exceptions are solar thermochemical conversion, which has a high investment cost for concentrating the solar energy, and biochemical conversion, where both investment and operating costs can be high, depending on the bioreactor type. Furthermore, the plasmachemical conversion is a highly modular technology, which is not dependent on an economy of scale and thus allows for local on-demand production capabilities.

Finally, one of the key-features for CCU techniques is their overall flexibility. This is actually a combination of many of the features discussed above. The plasmachemical conversion has a tremendous advantage here, due to its feed flexibility (CO₂, CO₂/CH₄, CO₂/H₂ and CO₂/H₂O), its energy source flexibility

(solar, wind, hydro, nuclear power), its operation flexibility (instant on/off, power scalability) and its flexibility of scale. None of the other technologies possess this unique combination of features required for successful worldwide implementation as a CCU technique.

In conclusion, plasma technology fares very well in this comparison, with its main disadvantages being: (i) its current need for post-reaction separation processes, (ii) the fact that the energy efficiency is dependent on the reactor type and (iii) the need for combination with catalysis to steer the product yield and selectivity.

6.3. Solar-to-fuel efficiency

In our opinion, the most interesting measure to compare the different technologies is based on how well solar energy is converted into chemical energy, known as the solar-to-fuel conversion efficiency ($\eta_{\text{solar-to-fuel}}$). First, we calculate this value for the various plasma-based technologies, based on the data presented above. Subsequently, we compare these solar-to-fuel efficiencies with the values for the other emerging technologies.

Where possible (see the discussion in Section 6.4 below), we calculated the solar-to-fuel conversion efficiency for all the pure CO₂ splitting data, based upon:

$$\eta_{\text{solar-to-fuel}} (\%) = \frac{E_{\text{out}}}{E_{\text{in}}} = \frac{\Delta H_{\text{c,fuel}} \left(\text{kJ mol}^{-1} \right)}{\text{SEI} \left(\text{kJ mol}^{-1} \right)} \times \eta_{\text{PV}} (\%)$$
(9)

where $\Delta H_{c,fuel}$ is the standard enthalpy of combustion of the fuel, based on the high heating value (HHV) of all the products so basically this is the energy output. Furthermore, the SEI is the specific energy input based on the plasma power, which is thus the energy input. Finally, η_{PV} is the photovoltaic efficiency for electricity production, which we need to be able to compare the technologies relying on direct and indirect solar energy. Of course, as discussed in Section 6.2 above, one of the key benefits of plasma technology is its energy source flexibility. Hence, when (renewable) electricity from other sources is used, this term can be removed from the equation (just like for the electrochemical conversion). As mentioned above, it is added here to allow an easy comparison of all the different novel technologies. However, in the overall comparison, it is important to keep this note in mind.

For the simple case of CO_2 splitting, this means $\Delta H_{c,fuel}$ equals:

$$\Delta H_{c,\text{fuel}} \text{ (kJ mol}^{-1}\text{)} = (\chi_{\text{CO}_2} \times S_{\text{C,CO}} \times 283 \text{ (kJ mol}^{-1}\text{)})$$

$$+ (\chi_{\text{CO}_2} \times S_{\text{O,O}_2} \times 0 \text{ (kJ mol}^{-1}\text{)})$$

$$\Rightarrow \Delta H_{c,\text{fuel}} \text{ (kJ mol}^{-1}\text{)}$$

$$= \frac{\dot{n}_{\text{CO,out}}}{\dot{n}_{\text{CO}_2,\text{inlet}}} \times 283 \text{ (kJ mol}^{-1}\text{)}$$

$$= Y_{\text{C,CO}} \times 283 \text{ (kJ mol}^{-1}\text{)}$$

where γ stands for the conversion, S and Y are the selectivity and yield, respectively, which are here expressed with respect to

C (for CO) and to O (for O_2), and \dot{n} stands for the molar flow rate.

Thus, since the C-based selectivity for CO in the case of pure CO₂ splitting in most cases is 100%, the equation for the solarto-fuel efficiency equals here the definition for energy efficiency (eqn (6); defined in Section 4.4.3) - as well as the thermal energy efficiency (based on the HHV) - multiplied by the photovoltaic efficiency:

$$\begin{split} \eta_{\rm solar-to-fuel}(\%) &= \frac{\Delta H_{\rm c,fuel} \; \left(\rm kJ \; mol^{-1}\right)}{\rm SEI} \; (\rm kJ \; mol^{-1})} \times \eta_{\rm PV}(\%) \\ &= \frac{\chi_{\rm CO_2} \times 283 \; \left(\rm kJ \; mol^{-1}\right)}{\rm SEI} \; (\rm kJ \; mol^{-1})} \times \eta_{\rm PV}(\%) \\ &= \eta \; (\%) \times \eta_{\rm PV} \; (\%) = \eta_{\rm Thermal, HHV}(\%) \times \eta_{\rm PV}(\%) \end{split}$$

For the DRM data, the equation becomes a bit more delicate. First of all we need to take the HHV of the converted CH4 in the feed into account as part of the denominator, since this counts as an energy input.

$$\begin{split} \eta_{\text{solar-to-fuel}}(\%) &= \frac{E_{\text{out}}}{E_{\text{in}}} \\ &= \frac{\Delta H_{\text{c,fuel}} \left(\text{kJ mol}^{-1}\right)}{\text{SEI} \left(\text{kJ mol}^{-1}\right) + \Delta H_{\text{c,conv CH}_4} \left(\text{kJ mol}^{-1}\right)} \\ &\times \eta_{\text{PV}}(\%) \end{split}$$

Second, as mentioned in the sections above, a wide variety of products is formed, which all need to be taken into account. Hence, for DRM, the solar-to-fuel efficiency would only equal the definition for energy efficiency (egn (6)) multiplied by the photovoltaic efficiency, when the stoichiometric reaction as defined in Section 2.2.1 occurs, i.e. $CO_2 + CH_4 \rightarrow 2CO + 2H_2$.

Note that since the SEI is based on the plasma power, this efficiency does not take the energy losses into account, which occur at the level of the power supply and power coupling with the plasma. The data regarding these energy losses are generally not reported, since those may vary greatly, depending on the plasma type and power supply used, but they are independent from the plasma process under study. Furthermore, a lot of (successful) research progress is still being made in minimizing these electrical losses when going from outlet power to plasma power.

It should be noted that the solar-to-fuel conversion efficiency could only be calculated based upon the available data in the literature. If certain reaction products were not measured, their contribution to the enthalpy of combustion could not be taken into account. Due to the scarcity of information regarding CO₂/H₂O and CO₂/H₂ mixtures, we only consider CO₂ splitting and DRM in the following analysis.

For the following discussions, we consider a PV efficiency value of 25% based on the efficiencies of the current commercial PV systems at the cell level.111 It is important to keep in mind that advancements in PV efficiency have a direct positive

influence on the solar-to-fuel efficiency of plasma technology. For example, PV efficiencies of up to 45% have already been reported on a lab scale, ²⁹⁸ a value which would almost double the solar-tofuel efficiency of plasma technology as reported below.

It is reported in the literature that - on a purely economic basis - for the various novel technologies to be cost competitive with existing chemical and fuel processes, a solar-to-fuel conversion efficiency of 20% is likely to be needed for the production of syngas. 65,110 Nonetheless, as already indicated above, all of these emerging technologies have additional advantages compared to the traditional catalytic technologies currently used. Furthermore, when the intermediate syngas step can be circumvented, i.e. due to the production of liquids through a direct oxidative pathway, this solar-to-fuel conversion efficiency target decreases drastically, e.g. for methanol, a solar-to-methanol efficiency of 7.1% would already be considered feasible. 110 This is the greatest advantage of the plasmachemical conversion (in combination with catalysis), namely its already proven capability of producing a wide variety of liquid chemicals in a direct (although currently unselective) manner.

Table 3 presents the best data obtained from the literature on solar-to-fuel efficiencies, together with the obtained conversions, for the different plasma types discussed in Section 5. It should be no surprise that the plasma set-ups that showed the best performance also have the highest solar-to-fuel efficiency. For pure CO₂ splitting, MW and GA plasmas are the two most

promising discharge types, with maximum achieved solar-tofuel efficiencies to date of 22.5% and 16.4%, respectively. For DRM, APGDs and GA plasmas have obtained the highest solarto-fuel efficiencies up to now of 23.0% and 22.1%, respectively. However, most other discharges, except for DBDs, already reach efficiencies between 11% and 15%. This means that the chemicals and fuels produced by the plasmachemical conversion of CO2 into syngas could already be cost competitive, depending on the critical notes made above on product separation costs and power supply efficiencies.

Finally, it is important to realize that this analysis is only a screenshot of the current production of syngas. When a successful shift is made towards the direct oxidative production of liquids, the performance of the different plasma types could change drastically. Indeed, certain plasma types (i.e. DBDs) allow for the easier implementation of catalytic materials, and the latter will play an important role in the selective production of these value-added compounds through the direct oxidative pathway. Furthermore, by circumventing the energy-intensive conversion of syngas to the desired liquids through the Fischer-Tropsch or methanol and subsequent synthesis processes, the required solar-to-fuel conversion efficiency needed to be competitive decreases by a factor of two to three.

Finally, in Table 4, we compare the reported solar-to-fuel conversion efficiencies obtained to date, as well as the theoretical maximum values, of plasmachemical conversion with

Table 3 Overview of the best solar-to-fuel efficiencies, along with the obtained conversions, calculated from the data in the literature for the different plasma types discussed in Sections 1.1 and 1.2, and presented in Fig. 24 and 32. For some plasma types, two or three 'best values' are listed, as some conditions lead to the best conversion, while others lead to the best efficiency; cf. the trade-off between both, as discussed in previous sections. Processes reaching efficiencies below 10% are considered inefficient (red), while those between 10-15% are considered promising (orange), above 15% very promising (green) and values above 20% might already be cost competitive (green, underline, bold). Note that this analysis applies to the production of the syngas components CO and H2; when considering the direct oxidative pathway to produce liquid fuels, lower values will already be cost competitive (see text)

	CO ₂ s	plitting	Dry reforming of methane		
	Xco ₂	$\eta_{solar-to-fuel}{}^a$	χ_{CH_4}/χ_{CO_2}	$oldsymbol{\eta_{solar-to-fuel}}^b$	
DBD	25.8% ¹¹⁵	5.8%	7% / 3% ¹⁸¹	4.2%	
DBD	42.0% ¹³⁵	1.7%	88% / 78% ²⁴¹	1.8%	
	9.7% ¹⁴⁰	<u>22.5%</u>			
MW (1980s)	35.9% ¹³⁸	17.1%			
	87.4% ²¹	9.4%			
	9.7% ¹⁴²	12.6%			
MW (2010s)	29.5% ¹⁴¹	10.8%			
	82.9% ¹⁴³	6.1%	71% / 69% ²⁵⁹	11.6%	
	12.2% ¹⁶⁰	16.4%	13% / 9% ¹⁹²	<u>22.1%</u>	
GA			41% / 36% ¹⁹²	15.5%	
			45% / 34% ²¹⁵	11.1%	
APGD	50.0% ¹⁷⁴	6.75%	94% / 77% ²¹⁹	<u>23.0%</u>	
Ns-pulsed	7.1% ¹⁷⁶	2.5%	61% / 50% ²¹²	14.5%	
Corona	6.1% ¹⁶⁷	1.6%	23% / 36% ¹⁹⁶	12.2%	
Spark			87% / 83% ¹⁹⁹	14.7%	

^a Based on eqn (9) above, using a PV efficiency of 25%. ^b Based on eqn (12) above, using a PV efficiency of 25%.

the other novel technologies. As mentioned in Section 3.1, as water electrolysis powered by renewable energy is already a more mature technology, we have added it here to the comparison.

Water electrolysis yields efficiencies in the range of 16-19% for a PV efficiency of 25% and an electricity-to-hydrogen efficiency of 65-75%.⁵⁸ Despite this success, electrochemical CO2 splitting, on the other hand, faces large efficiency losses due to overpotentials, and the theoretical efficiency is expected to be below 15%. For the solar thermochemical approach, theoretical efficiencies of 16-19% or 35-50% are assumed, 58,61,65 depending on the heat recovery (see details in the cited references). Nevertheless, efficiencies above 10% are still pending experimental demonstration with robust and scalable solar reactors. 59,61,65,110 For the photochemical conversion, the efficiency is theoretically limited to a maximum below 5% or 17%, 68 depending on the band gap energy of the photocatalysts. However, the actual solar-to-fuel conversion efficiencies obtained to date are generally much lower (<2%), 58,68 with some exceptions when coupled with an electrolytic cell (10.9%).68 For the biochemical conversion, a maximum efficiency for the conversion of light to stored chemical energy of about 4.5% has been calculated.³⁵ However, this value is rarely achieved. Only in exceptional cases will dry matter yield exceed 1% or 2%.³⁵

Finally, it is clear from Table 4 that the highest value obtained to date on a lab scale for the plasmachemical conversion of CO2, i.e. 23% (see Table 3 above), already appears to be competitive with the current most mature technology (i.e. water electrolysis) to transform renewable energy into chemicals and fuels through the intermediate production of the syngas components CO and H2. Furthermore, the theoretical maximum energy efficiency is 90-95%, when multiplied by the power supply efficiency and the PV efficiency. This is based on the theoretical - and experimentally proven - most energy-efficient CO₂ dissociation process through vibrational excitation (see Section 4.1.4). Hence, the construction, use and matching of

Table 4 Overview of the currently obtained and theoretical maximum solar-to-fuel conversion efficiencies to give the syngas components CO and H₂ for the different emerging technologies described in Section 3, as well as by plasma technology

Solar-to-fuel efficiency	Currently obtained	Theoretical maximum
Water electrolysis	16-19% ^a	65–75% ⁵⁸ (x PV efficiency)
Electrochemical conversion Solar thermochemical conversion	NA $0.4-0.8\%^{61,65}$ $1.7-3.5\%^{59}$ $7.1\%^{110}$	<15% ⁵⁸ 16-19% ^{61,65} 35-50% ⁵⁸
Photochemical conversion	NA $< 2\%^{58}$ $0.01-10.9\%^{68}$	$<5\%^{58}$ $\sim 17\%^{68}$
Biochemical conversion	NA	4.5% ³⁵ <i>x</i> harvesting efficiency
Plasmachemical conversion	23% ^{a,b}	90–95% x power supply efficiency (x PV efficiency)

^a Based on a PV efficiency of 25%. ^b Without taking the power supply efficiency into account, due to lack of this data in the literature.

efficient power supplies for generating the plasma is highly critical. Nonetheless, we should not forget the critical notes made above on product separation. Additional - and indeed substantial - research is needed to provide an answer to these issues, just like in the case with all the other emerging technologies and their specific concerns. Due to its emerging character, interdisciplinary research towards plasma technology, especially in combination with other fields, such as catalysis will, undoubtedly, lead to further advancement and breakthroughs in this field.

6.4. Research recommendations

To conclude, we would like to provide some research recommendations for plasma technology, based on all the collected data from the literature, to further advance the field of plasmabased CO₂ conversion. The first recommendation is related to the need for a more standardized framework in reporting data, to allow easier comparison of data within and outside the field of plasma technology. The most important criterion here is to be able to compare the conversions and energy cost/efficiency. Regarding the conversions, this requires a clear reporting on the presence of diluting agents (i.e. N2, He, Ar) when used, since these agents can significantly influence the plasma process, as discussed in Section 5. Furthermore, Pinhão et al. 299 elaborately described the effect of gas expansion during plasma processing, which is to date often neglected, but which can have a tremendous influence on data accuracy, for example on the conversions obtained through GC measurements. 92,95,125 Concerning the energy cost/efficiency, first of all there should be no room for interpretation about whether the reported power is the applied input power or the measured plasma power and through which electrical techniques, such as Lissajous plots, this power was obtained. Second, one should aim to identify all the important products and their selectivities, to be able to determine the energy efficiency based on the enthalpy of the reaction, as well as the so-called fuel production efficiencies. Third, data on power supply efficiency and its optimization are urgently required to report accurate solar-to-fuel conversion efficiencies. In general, as also discussed by Butterworth et al. 136 for the testing of materials for plasma-catalysis, general frameworks on performing experiments and reporting data are becoming essential for the transparent further development of plasma research, both within the plasma community and within the larger CO2 conversion community.

Finally, we also have some more personal recommendations. From the plots in the sections above, showing the energy cost/efficiency as a function of the SEI, it becomes more than clear that most applied SEIs are far too high, especially for the combinations with catalysis. An SEI range of 0.1-5 eV per reactant molecule (taking possible dilutions into account) should be the target to achieve energy-efficient conversions, as also recommended by Fridman.²¹ Furthermore, since plasma processes are very susceptible to the effect of impurities in the feed, more research and insights are needed towards the effect of real-life gas compositions on both the physical side (e.g. discharge stability) and on the chemical side (e.g. product distributions, harmful by-products).

7. Conclusions and outlook

The conversion of CO₂ – preferably into value-added products – is considered one of the great challenges of our century. In the past decade, substantial progress with several novel technologies has been made. The last section of this review intended to provide the reader with a state-of-the art and critical assessment of an – up till now – rather underexposed emerging technology: the plasmachemical conversion of CO₂. To achieve this, we provided an introduction to the basic concepts of plasma, and we demonstrated its viability as a gas conversion technology, finally putting it in the broader context of its peers.

From the advances in the area of CO_2 conversion discussed in this review, one outcome stands out without doubt. The question is not 'if' one of these novel technologies will be industrially competitive, but rather 'when' and 'which one(s) will play the leading role'. To shine a brighter light on – the final burning question – whether plasma technology could be the answer – or at least be part of the equation – we briefly summarize here the arising opportunities and challenges for plasma technology in the field of CO_2 conversion.

Plasmas possess some important advantages over certain other (novel) technologies: (i) they can operate at room temperature using any source of (renewable) electricity, (ii) they have a large flexibility in terms of the feeds that need to be processed, (iii) they provide an extremely flexible 'turnkey' process, which allows for the efficient storage of energy, peak shaving and grid stabilization, (iv) the reactors have low investment and operating costs, (v) they have a simple scalability both in size and applicability, and (vi) last but not least, the technology does not rely on rare earth materials – making it rather unique at this point. This unprecedented combination of features gives plasmachemical conversion a very high overall flexibility, making it an extremely useful and valuable technology for CCU.

The flip side of the coin is that the reliance of plasma technology on indirect solar energy in the form of electricity is – at the same time – a limiting factor for the solar-to-fuel conversion efficiency, especially compared to technologies that can directly harvest solar energy. Nonetheless, this also allows plasma technology to rely on other renewable energy sources, and more importantly, it adds to its overall – location and process – flexibility, since it is not dependent on the availability of sunlight. More urgent, however, is the issue of product separation. From all the data available in the literature, it is clear that converting CO_2 – with or without an additional hydrogen source – always yields a mixture of products after reaction. This implies the need for an – often expensive – post-processing separation step.

From the benchmarking discussion, it is clear that plasma technology can definitely play an important role in the field of CO₂ conversion, and it is not beyond our grasp to think about its eventual commercial implementation, be it on a large or

small scale. Nevertheless, as always, only a few candidates seem suitable for the specific task at hand, as we outline here.

Two main CO₂ conversion strategies were discussed: (i) pure CO₂ splitting and (ii) CO₂ conversion in combination with a co-reactant serving as the hydrogen source, i.e. CH4, H2 and H₂O, yielding processes named the dry reforming of methane (DRM), the hydrogenation of CO₂ and artificial photosynthesis, respectively. At this time, the conversion of CO₂ through the latter two processes is - although possible - inefficient in a plasma-only set-up, not to mention industrially undesirable, due to the formation of possibly dangerous mixtures to be handled. In the future, the combination of plasma with other technologies, such as catalysis or electrolysis, could solve these issues. On the contrary, for both pure CO₂ splitting and DRM, the proposed energy efficiency target of 60% has already been surpassed by several types of plasma reactors, resulting in a solar-to-fuel conversion efficiency for the production of syngas above - the required - 20% mark.

In the short run, pure CO_2 splitting using the most energy-efficient set-ups, *i.e.* MW and GA discharges, appears to have a high potential. In this case, pure CO and O_2 could be produced, after a separation step, or a pure CO stream when *in situ* trapping of the O_2 can be successfully implemented, *e.g.* by combination with other technologies or through the addition of scavengers.

In the long run, it seems evident that CO₂ conversion in combination with a hydrogen containing co-reactant has the highest potential for the efficient production of value-added products, such as alcohols, aldehydes, esters and acids. In particular, the direct oxidative pathway, in which the intermediate syngas step is circumvented, has great potential. For now, however, a lot of research in this area is still needed to increase the selectivity towards these valuable bulk chemicals over the currently produced syngas. This will most probably need to be achieved in combination with catalysts. However, even today, APGD and GA discharges already provide an energy-efficient alternative to produce syngas in any desired ratio through DRM.

By all means, plasma technology is not just an overlooked unicorn - although its combination of features does make it quite unique for CCU - but, as mentioned several times throughout this work, and just like for all other emerging technologies in this field, several important challenges remain. General challenges comprise the need for further fundamental research, concerning: (i) the plasmachemical processes taking place in warm discharges, such as MW and GA, which stimulate vibrational excitation; whereby the latter, representing the most energy-efficient dissociation pathway, should be further exploited by enhancing the non-equilibrium character of these plasmas to further improve the energy efficiency, (ii) the combination of plasma with catalysis and the possibility of taking advantage of the synergetic effects and the selective production of value-added compounds, (iii) in situ O/O2 trapping by using scavengers, or combinations with other technologies, such as catalysis or SOEC, to further enhance the conversions and yields towards the desired value-added compounds, (iv) the demonstration of successful scale-up for other discharges than DBDs and

(v) the production and tuning of efficient power supplies for generating and sustaining the plasma.

Moreover, some important discharge-specific research challenges can also be identified. For MW plasmas, the ultimate goal should be to achieve the same energy efficiency at atmospheric pressure as currently obtained at reduced pressure for CO₂ splitting, while more research towards DRM is essential. For GA plasmas, the limited amount of gas that passes through the active arc region, and thus that can be converted by the plasma, is currently limiting the overall CO₂ conversion, and more research in this direction is needed, e.g. by improving the reactor design. Furthermore, some other plasma reactors show good performance, but the available data and insights are still limited, thus requiring more research to explore their full potential. Of course, inherent to every review, the above analysis represents the current state-of-the-art. Plasma-technology-based CO₂ conversion is relatively new and it is clearly a fast advancing field with - just like for any emerging technology - ample room for improvement and new (interdisciplinary) developments.

It appears that the ultimate plasma for CO_2 conversion would be a system with a high electron density (N_e) and a low electron temperature (T_e) – resulting from a low reduced electric field (E/n) – which are essential to stimulate and achieve a high fraction of vibrational excitation. At the same time, this system should operate at a low gas temperature (T_g) to minimize the VT relaxation losses and backward recombination reactions. Finally, to be industrially attractive, this should be achieved at atmospheric pressure.

Future research towards the direct oxidative production of oxygenates, as well as multi-reforming processes with a combination of gases (e.g. CO₂, CH₄ and H₂O or O₂) or several reforming processes in series, using plasma technology appears very promising and should be pursued. Furthermore, more interdisciplinary research into the combination of plasmas with other novel technologies is highly desirable. For catalysis, the combination is already proving alluring, and it is our opinion that combinations with electrolysis, although even more challenging, could also lead to synergetic effects. As is the case for many important issues brought forward in the current age, with a high probability the future will rely on a true amalgam of technologies and solutions, each for its specific tasks, rather than the 'one solution (or technology) fits all' mentality of the old age.

From the current analysis, it is evident that pure CO_2 splitting could reach industrial implementation at a faster pace. Nevertheless, CO_2 conversion using a hydrogen source as a co-reactant should certainly be further pursued, since – when successful – in the long run it offers us the possibility of producing a wide variety of value-added chemicals and fuels, starting from the same building block and allowing the flexibility to tune the output depending on the market's needs. On that account, we believe plasmas could be – at least part of – the bright light that shines on our horizon.

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