ABSTRACT: Plasma technology is gaining increasing interest for CO₂ conversion. The gas temperature in (and after) the plasma reactor largely affects the performance. Therefore, we examine the effect of cooling/quenching, during and after the plasma, on the CO₂ conversion and energy efficiency, for typical "warm" plasmas, by means of chemical kinetics modeling. For plasmas at low specific energy input (SEI ∼ 0.5 eV/molecule), it is best to quench at the plasma end, while for high-SEI plasmas (SEI ∼ 4 eV/molecule), quenching at maximum conversion is better. For low-SEI plasmas, quenching can even increase the conversion beyond the dissociation in the plasma, known as superideal quenching. To better understand the effects of quenching at different plasma conditions, we study the dissociation and recombination rates, as well as the vibrational distribution functions (VDFs) of CO₂, CO, and O₂. When a high vibrational–translational (VT) nonequilibrium exists at the moment of quenching, the dissociation and recombination reaction rates both increase. Depending on the conversion degree at the moment of quenching, this can lead to a net increase or decrease of CO₂ conversion. In general, however, and certainly for equilibrium plasmas at high temperature, quenching after the plasma helps prevent recombination reactions and clearly enhances the final CO₂ conversion. We also investigate the effect of different quenching cooling rates on the CO₂ conversion and energy efficiency. Finally, we compare plasma-based conversion to purely thermal conversion. For warm plasmas with typical temperatures of 3000–4000 K, the conversion is roughly thermal.

1. INTRODUCTION
To mitigate the devastating effects of climate change on society, extensive research is carried out on the conversion of greenhouse gases responsible for global warming. Plasma, created by applying electricity to a gas so that a cocktail of reactive species is formed (electrons, ions, radicals, excited species), is of particular interest, since it can convert (excess) electrical energy from intermittent renewable energy sources into chemical energy, it is very flexible, and it can be scaled easily. Many different types of plasma reactors are investigated for CO₂ conversion, such as dielectric barrier discharges (DBDs), microwave (MW) plasmas, atmospheric pressure glow discharges (APGDs), and nanosecond-pulsed plasmas. MW, GA, APGD, and nanosecond-pulsed plasmas yield a better performance than, e.g., DBD plasmas. This is generally attributed to the lower reduced electric field (E/N) of 50–100 Td compared to DBD plasmas (200 Td and above), which leads to a lower electron temperature (1–2 eV), compared to the latter (3 eV and higher).

At lower electron temperatures, more energy goes into vibrational excitation, while electrons at higher energies mainly cause electronic excitation, ionization, and direct electron impact dissociation. The latter explains the lower energy efficiency of DBD plasmas because direct electron impact dissociation requires at least 7 eV energy, which is considerably greater than the bond energy of CO₂ (5.5 eV), so the extra energy is wasted.

The electron energy that goes into the lower CO₂ vibrational levels in MW, GA, and APGD plasmas can either relax to heat in so-called vibrational–translational (VT) relaxation or undergo the so-called vibrational–vibrational (VV) relaxation, in which CO₂ molecules can acquire more vibrational energy. This process is called vibrational ladder climbing and proceeds until the vibrational energy of the CO₂ molecules reaches the dissociation limit. Over the last few years, research on CO₂ plasmas was largely focused on increasing the energy efficiency of CO₂ conversion. A lot of attention was paid to increasing the VT nonequilibrium. The aim was to enhance the vibrational temperature without (significantly) raising the gas temperature. However, in most experiments conducted with MW, APGD, and GA discharges, the plasma appears to be in (or close to) thermal equilibrium, with the gas temperature (nearly) equal to the vibrational temperature.
These high gas temperatures (i.e., >3000 K) can even be reached at relatively low specific energy inputs (SEIs) because the power input is localized in a small gas volume, leading to a much higher local SEI. This phenomenon is especially pronounced in APGD, GA, and MW plasmas. In APGD and GA plasmas, the contracted nature is inevitable, since the discharge is characterized by a plasma column that connects to two electrodes. In MW plasmas, the discharge is known to radially contract with increasing pressure, starting from around 100 mbar,39 the discharge is known to radially contract with increasing column that connects two electrodes.2,15,38 In MW plasmas, the contracted nature is inevitable, since the discharge is characterized by a plasma phenomenon is especially pronounced in APGD, GA, and MW plasmas. In APGD and GA plasmas, the contracted nature is especially pronounced in APGD, GA, and MW plasmas. In APGD and GA plasmas, the contracted nature is especially pronounced in APGD, GA, and MW plasmas.

The high temperatures lead to significant thermal dissociation at the expense of vibrational-induced dissociation. The vibrational distribution function (VDF) exhibits a Boltzmann distribution, with no overpopulation of the higher vibrational levels, and the dissociation mainly occurs from the lower vibrational levels of CO2. While high energy efficiencies can be reached at these temperatures, the overall conversion starts to become constrained by the recombination reactions, mainly of CO with O2 molecules, which become more important at higher temperatures.31,38,39

A possible pathway to limit these recombination reactions at high gas temperatures is by cooling the gas, thereby quenching the converted reaction products. Experimentally, this can be accomplished by a sharp temperature gradient between the plasma core and the surrounding gas. In addition, the gas can be cooled by enhanced radial heat transport, achieved in turbulent flow regimes, or by supersonically accelerating the gas. For high-SEI plasmas, however, in which more than 0.19 eV/molecule goes to heat, this acceleration has to take place in the afterglow to prevent thermal choking of the flow.

Next to limiting the recombination reactions, and thus freezing of the reaction products (so-called ideal quenching), it is also possible to increase the CO2 conversion upon quenching. This is called superideal quenching and happens when a VT nonequilibrium, created or enhanced by the sudden drop in gas temperature, promotes VV relaxation to the higher vibrational levels and further enhances the dissociation reactions. More specifically, this can happen by the reactions of vibrationally excited CO2 molecules with any molecule (M) or with an O atom (i.e., CO2 + M → CO + O or CO2 + O → CO + O2).

Although it has been experimentally and theoretically demonstrated that high energy efficiencies for CO2 conversion can be reached in warm plasmas at temperatures of 3000 K and above, and it is often stated that quenching is promising to reach higher energy efficiencies, more insight is needed to demonstrate its full potential. Therefore, we examine here the effect of quenching on the CO2 conversion and energy efficiency for different plasma conditions. Furthermore, we compare the plasma conversion with purely thermal conversion, for the same power input, to elucidate the difference in performance and to determine the actual role of thermal dissociation in plasma-based CO2 conversion.

2. MODEL DESCRIPTION

2.1. Plasma Model. We apply a zero-dimensional (0D) chemical kinetics model, using ZDPPlasKin. This model solves the continuity equation for the different plasma species

\[
\frac{\text{d}n_j}{\text{d}t} = \sum_j R_j \{a_j^R - a_j^I\} = \sum_j (k_j \prod_l n_l) \{a_j^R - a_j^I\}
\]

in which index \(j\) refers to reaction \(j\) and index \(I\) refers to the different reactants of reaction \(j\). \(a_j^R\) and \(a_j^I\) are the right- and left-hand side stoichiometric coefficients of species \(s\), respectively, taking part in reaction \(j\), \(k_j\) is the reaction rate coefficient, and \(R_j = k_j \prod_l n_l\) is the reaction rate, with \(\prod n_l\) being the product of densities of species present on the left side of reaction \(j\).

Within the ZDPPlasKin framework, a chemical kinetics solver is coupled to the Boltzmann solver BOLSIG+ to calculate the electron energy distribution function (EEDF) using a set of cross sections, corresponding to the reactions listed in Table S1 of the Supporting Information (SI), including superelastic collisions. The EEDF is regularly updated during the simulations, upon changes in the chemical composition, reduced electric field, or gas temperature.

2.2. Chemistry Set. The species in our model are listed in Table 1. The chemistry set is a slightly reduced form of other

<table>
<thead>
<tr>
<th>Table 1. Species Described in the Model</th>
</tr>
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<tbody>
<tr>
<td>Neutral Ground-State Species</td>
</tr>
<tr>
<td>CO2, CO, O2, O, C</td>
</tr>
<tr>
<td>Charged Species</td>
</tr>
<tr>
<td>CO2′, CO′, O2′, CO+, O+, O2−, CO2−, e−</td>
</tr>
</tbody>
</table>

excited states | associated energy [eV] | state^\* |
<table>
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<th></th>
<th></th>
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<tbody>
<tr>
<td>O2[ν−33]</td>
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</tr>
<tr>
<td>CO[ν−50]</td>
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<tr>
<td>CO2[ν−21]</td>
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<td>CO2[e2]</td>
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<td>(030) + (110)</td>
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<tr>
<td>CO2[e3]</td>
<td>0.339</td>
<td>(040) + (120) + (200)</td>
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<tr>
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<td>(∑u) + (∑Π) + (∑Λ)</td>
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<tr>
<td>CO[π]</td>
<td>13.5</td>
<td>(d1Δ) + (eΣg^-)</td>
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</tbody>
</table>

^CO electronic state designation from Grofulovic et al.; O2 and CO electronic states notation from Itikawa.

sets used in the past. The effect of the chemistry reduction on the model results was examined in ref 31. This reduced chemistry set has been used in more recent works, including an uncertainty analysis in which it was argued that more species had a negative impact on the model uncertainty, while they virtually had no influence on the model results.

Given the importance of vibrational ladder climbing in studying the role of thermal (equilibrium) vs nonequilibrium CO2 dissociation, we include all (21) asymmetric mode vibrational levels, up to the dissociation limit, as well as the four (degenerate) lowest symmetric mode vibrational levels, which include both the bending and symmetric stretch modes.
The choice to only include these few symmetric mode levels is justified, since for \( E/N > 20 \text{ Td} \), and thus at the conditions under study here, primarily the asymmetric mode is excited\(^{(2,5)} \) and experiences a very fast \( \text{VV} \) relaxation.\(^{(28,51)} \)

For conditions under study here, primarily the asymmetric mode is temperature. For example, if all CO\(_2\) is dissociated into CO and O\(_2\), 50 and 33 levels are taken into account, respectively. For O\(_2\), these levels again reach the dissociation limit. For CO, there are 63 levels up to the dissociation limit, \( \text{changes in the gas temperature or upon changing the chemical composition (e.g., 1 CO}\(_2\) molecule is converted into CO and O} \) below).

The full set of reactions in our model can be found in the SI (Tables S1–S5) and includes electron impact, ion–ion, and neutral–neutral reactions and \( \text{VT} \) and \( \text{VV} \) relaxation between different molecules.

All reactions in this chemistry set are characterized by an individual uncertainty. While these uncertainties will have an impact on specific calculation results, such as the CO\(_2\) conversion, the trends predicted are typically not affected by these uncertainties.\(^{(49)} \) Note that this quoted study only applied to gas temperatures up to 2000 K, and the effect of the uncertainties on the neutral reactions can lead to higher deviations at the higher temperatures considered here. We therefore focus more on the overall trends rather than on the absolute values of, e.g., CO\(_2\) conversion and energy efficiency.

Most of the rate coefficients and cross sections are generally known for the ground-state or low-vibrational-level molecules. Scaling laws are used to scale these rate coefficients toward higher vibrational levels. To determine the various rate coefficients, we mostly follow the same approach as in previous works,\(^{(28,29,31,45,48–51,54)} \) with an additional correction term on the rates, scaled with the Fridman Macheter \( \sigma \) model. This has been explained in more detail in the SI.

**2.3. Modeling Approach.** We follow the time evolution of a volume element as it moves through a plasma reactor. The model starts with pure CO\(_2\) mimicking the gas at the inlet of the reactor, and a Boltzmann vibrational distribution, corresponding to 300 K, i.e., the gas temperature at the inlet. At \( t = 0 \), the plasma power is applied to the gas (or thermal power when we use the model for pure thermal conversion; see below).

At any time during the simulation, the ideal gas law is applied. We assume that pressure remains constant, so the volume element considered can expand or contract upon changes in the gas temperature or upon changing the chemical composition (e.g., 1 CO\(_2\) molecule is converted into CO and O, thus increasing the number of species). We take this effect into account by introducing a gas expansion factor, \( \beta(t) \), which equals 1 at the start of the simulation and drops when the gas expands. The factor is defined as

\[
\beta(t) = \frac{M(t = 0)T_g(t = 0)}{M(t)T_g(t)}
\]

where \( M \) is the total number of molecules and \( T_g \) is the gas temperature. For example, if all CO\(_2\) is dissociated into CO and \((1/2)\)O\(_2\), the initial number of molecules \( M(t = 0) \) rises to \((3/2)M(t = 0)\), and thus \( \beta = (2/3) \), at constant gas temperature because the volume expands to keep the pressure constant. Similarly, the expansion factor drops upon rising gas temperature. The electron density is not included in this expansion factor, as the ionization degree is too small.

The plasma conditions are defined in the model by setting a certain ionization degree (\( d \)). The electron density (\( n_e \)) then equals \( d \times N \) (with \( N \) being the gas number density). At the same time, a constant (DC) reduced electric field \( E/N \) (with \( E \) being the electric field) is set. The deposited power density (in eV/(cm\(^3\) s)) applied to the plasma is then calculated using Joule’s law

\[
P_{\text{dep,p}} = \sigma E^2 = \sigma N \left( \frac{E}{N} \right)^2
\]

where \( \sigma \) is the conductivity (in C/(cm V s)), given by \( \sigma = e n \mu_e \), the electron mobility \( \mu_e \) is obtained from the Boltzmann solver, and \( e \) is the elementary charge.

To compare the plasma performance with pure thermal simulations, we apply a heat source term with the same power deposition profile \( P_{\text{dep},\text{pl}} \), as in the plasma case

\[
\frac{N_{\text{th}}P_{\text{dep},\text{th}}}{\beta_{\text{th}}} = \frac{N_{\text{pl}}P_{\text{dep},\text{pl}}}{\beta_{\text{pl}}}
\]

with \( N_{\text{th}} \) and \( \beta_{\text{th}} \) being the total gas number density and expansion factor of the thermal simulation, respectively, and \( N_{\text{pl}} \) and \( \beta_{\text{pl}} \) being for the plasma simulation, respectively.

At time \( t = 0 \), the plasma is ignited by applying the power as described above. The plasma is sustained until the applied power reaches the value corresponding to a certain predefined SEI (in eV/molecule), defined as

\[
\text{SEI} = \frac{\int_0^t P_{\text{dep},\text{pl}}(t) dt}{N(t = 0)}
\]

Note that the dependency between the SEI and the residence time is specific to the conditions under study. In reality, a similar SEI can be reached with different deposited power profiles, leading to a different deposited power density. When the power density is lower, the residence time will increase. However, this situation will correspond to lower ionization degrees or reduced electric fields, and we decided to keep these fixed in our study, as characteristic plasma parameters. Also, in a 0D model, the plasma parameters are assumed to be homogeneous. In reality, localized power deposition can also lead to longer residence times for the same total SEI.\(^{(20)} \)

We aim to study the effect of quenching at conditions of maximum CO\(_2\) conversion and energy efficiency. This is typically achieved in MW plasmas at intermediate pressure.\(^{(19,21)} \) We therefore assume a fixed power of 100 mbar. The CO\(_2\) conversion is calculated as

\[
X(t) \text{ [%]} = \left[ 1 - \frac{N_{\text{CO}_2}(t)}{\beta(t)N_{\text{CO}_2}(t = 0)} \right] \times 100\%
\]

where \( N_{\text{CO}_2} \) is the total number density of all CO\(_2\) molecules (i.e., the sum of the ground-state and all vibrationally and electronically excited CO\(_2\) molecules) at a given time \( t \).

The energy efficiency is calculated as

\[
\eta \text{ [%]} = \frac{X \Delta H^s}{\text{SEI}}
\]

where \( \Delta H^s = 2.93 \text{ eV/molecule} \) is the reaction enthalpy of the reaction \( \text{CO}_2 \rightarrow \text{CO} + (1/2)\text{O}_2 \) at the inlet temperature of 300 K.

At the plasma conditions under study, not all of the applied power is used for CO\(_2\) conversion, but a significant fraction is
released as heat. In our model, the gas temperature is calculated self-consistently according to
\[ N \frac{\gamma c_p}{\gamma - 1} \frac{dT}{dt} = P_{el} + \sum_j R_j \Delta H_j - P_{ext} \]
(8)
where \( \gamma = c_p/c_v \) is the ratio of specific heats that is regularly updated depending on the concentration of the different species, \( P_{el} \) is the gas heating power density due to elastic electron-neutral collisions, \( \Delta H_j \) is the heat released (or consumed) by each reaction \( j \), and \( P_{ext} \) is the power loss density due to exchanges with the surroundings.

The heat exchange with the surroundings is particularly important to tune the (non) equilibrium effects. The external cooling term is defined as
\[ P_{ext} = \frac{8\lambda(T_e)}{R^2}(T_g - T_w) \]
(9)
where \( \lambda \) is the gas thermal conductivity, \( T_w \) is the wall temperature (assumed as 300 K), and the reactor radius \( R \) is set to 7 mm.56,57 The gas thermal conductivity is calculated by
\[ \lambda(T_g) = (0.071T_g^0.5 - 2.33) \times 10^{-3} \text{ W/(m K)}. \]

To study the effect of quenching on CO2 dissociation, we added a tuning constant \( c \) to increase or decrease the cooling power (or thermal conductivity), which determines the gas temperature. Without quenching, \( c \) remains the same during the entire simulation. The gas can be quenched by dropping the gas temperature to 300 K (i.e., instantaneous quenching) or by increasing the cooling constant (i.e., \( c_1 \) before quenching and \( c_2 \) after quenching).

In reality, the cooling by thermal conductivity can be increased by gas flow turbulence.16,44 A rise in thermal conductivity by a factor 100 (which in our model corresponds to \( c = 100 \)) is quite feasible for CO2 at room temperature. Indeed, while in our model, the thermal conductivity at 300 K is 0.019 W/(m K), the effective thermal conductivity due to turbulence at 300 K was estimated to be 1–3 W/(m K)17 and hence a factor \( (= 100) \) higher. The highest cooling constant in our model is \( c = 1000 \), which probably may only be realized upon mixing with a cold gas. However, the latter could also result in changes in the gas composition, but these effects are beyond the scope of our study. Indeed, our study is more conceptual.

3. RESULTS AND DISCUSSION

All results are presented for a pressure of 100 mbar, which typically yields high energy efficiencies.4 The ionization degree \( \langle\tilde{d}\rangle \) is set to \( 10^{-6} \), which can be considered as a lower value for diffuse microwave plasmas56 but could also be applicable for glow discharge plasmas.59 The reduced electric field \( E/N \) is set to 50 Td, as this is a value that ensures high vibrational excitation of the asymmetric mode levels of CO2. We also present the results for an ionization degree \( \langle\tilde{d}\rangle \) of \( 10^{-4} \) and \( E/N \) of 100 Td, to illustrate the behavior in a wider range of conditions.

3.1. Effect of the Cooling Constant on the CO2 Conversion, Gas, and Vibrational Temperature During Plasma

Figure 1 shows the time evolution of CO2 conversion (left axis) and gas and vibrational temperature (right axis) for a plasma with SEI = 5 eV/molecule at weak (a) and strong (b) cooling and hence representing a VT equilibrium (or warm) plasma and a nonequilibrium (or cold) plasma, respectively.
values, since VT relaxation is much less important at low gas temperatures. At an SEI of 1 eV/molecule, the CO$_2$ conversion reaches 9.5%, which is more than double of what was reached at the same SEI in Figure 1a. However, when the SEI is higher than 2 eV/molecule, the CO$_2$ conversion rises much more slowly than in Figure 1a. At 5 eV/molecule, a CO$_2$ conversion of 28.3% is reached, which is much lower than in Figure 1a. Thus, at SEI values below or equal to 1 eV/molecule, cooling is detrimental, pointing toward the important role of thermal dissociation. This will be further elaborated in Section 3.7.

3.2. CO$_2$ Conversion and Gas Temperature in the Afterglow of Warm Plasmas. In Figure 2, we plot the time evolution of CO$_2$ conversion and gas temperature in the afterglow of a plasma with high or low SEI. For an SEI of 0.5 eV/molecule (Figure 3a), the optimal quenching moment is at the plasma end. The conversion increases rapidly after quenching, due to the VT nonequilibrium, enhanced by the fast drop in gas temperature. This is called “superideal quenching”. Quenching at maximum conversion also yields a higher CO$_2$ conversion, but since the vibrational temperature drops quickly after switching off the plasma,$^{54}$ the VT nonequilibrium, and thus the increase in conversion, is much smaller than when quenching at the plasma end (cf. yellow and red curves in Figure 3a).

For an SEI of 4 eV/molecule, the optimal quenching moment is at maximum conversion. Since a chemical equilibrium is not yet reached at the plasma end, the conversion further increases after the plasma (see also Figure 2). For both quenching cases, the CO$_2$ conversion remains more or less frozen (except for a small drop immediately after quenching). The VT nonequilibrium created by the drop in gas temperature is not enough to produce a rise in conversion (i.e., superideal quenching). The different trends in CO$_2$ conversion upon quenching, for the two SEI values, will be discussed more in depth in the following sections.

3.3. Instantaneous Quenching. Instantaneous quenching, i.e., quickly reducing the gas temperature to 300 K, can prevent the recombination. Indeed, at 300 K, thermal dissociation and recombination are not important, and the converted reaction products are “frozen”.

We describe here the effect of instantaneous quenching, (i) at the plasma end and (ii) in the afterglow, when the CO$_2$ conversion reaches its maximum without any quenching (see Figure 2).

3.3.1. Quenching after Warm Plasmas. In Figure 3, we plot the time evolution of CO$_2$ conversion (top panels) and gas temperature (bottom panels) for a plasma with SEIs of 0.5 eV/molecule (left panels) and 4 eV/molecule (right panels) at weak cooling (c = $10^{-3}$). Note that a VT equilibrium exists at the plasma end with an SEI of 4 eV/molecule but not at 0.5 eV/molecule (cf. Figure 1a).

Quenching clearly leads to a higher final CO$_2$ conversion in all cases, but the ideal quenching moment is different for the plasma with high or low SEI.

For an SEI of 0.5 eV/molecule (Figure 3a,c), the optimal quenching moment is at the plasma end. The conversion increases rapidly after quenching, due to the VT nonequilibrium, enhanced by the fast drop in gas temperature. This is called “superideal quenching”. Quenching at maximum conversion also yields a higher CO$_2$ conversion, but since the vibrational temperature drops quickly after switching off the plasma,$^{54}$ the VT nonequilibrium, and thus the increase in conversion, is much smaller than when quenching at the plasma end (cf. yellow and red curves in Figure 3a).

For an SEI of 4 eV/molecule, the optimal quenching moment is at maximum conversion. Since a chemical equilibrium is not yet reached at the plasma end, the conversion further increases after the plasma (see also Figure 2). For both quenching cases, the CO$_2$ conversion remains more or less frozen (except for a small drop immediately after quenching). The VT nonequilibrium created by the drop in gas temperature is not enough to produce a rise in conversion (i.e., superideal quenching). The different trends in CO$_2$ conversion upon quenching, for the two SEI values, will be discussed more in depth in the following sections.

Next to the conversion, the energy efficiency of quenching. Figure 4 summarizes both the conversion and energy efficiency as a function of SEI without quenching (blue bars), in the case of quenching at the plasma end (red bars), and at maximum conversion (yellow bars).

For all SEI values, quenching at maximum conversion enhances the final conversion and energy efficiency, compared
to no quenching. The rise is the most pronounced for higher SEI values, where the conversion is the most affected by a gradual drop in gas temperature (Figure 2). Our model predicts a maximum final conversion of 90%, for quenching at maximum conversion, at an SEI of 5 eV/molecule and a maximum energy efficiency of 58%, again for quenching at maximum conversion, but at an SEI of 4 eV/molecule.

Quenching at the plasma end only enhances the final conversion and energy efficiency for SEI below 1 eV/molecule and above 2 eV/molecule. For SEI below 1 eV/molecule, the VT nonequilibrium at the plasma end, further enhanced upon quenching, yields superideal quenching; see also Figure 3a. For SEI values of 1 and 2 eV/molecule, the gas temperature at the plasma end is high enough to allow thermal CO₂ dissociation, but there is no significant drop in conversion in the afterglow due to gradual cooling (cf. Figure 2a) because the CO₂ conversion is limited and thus the recombination reactions are not so important as for higher SEI values. Hence, quenching does not play such a big role in reducing the eventual drop in CO₂ conversion. For higher SEI values, the drop in CO₂ conversion is very pronounced (see Figure 2a), thus explaining why quenching at the plasma end is again beneficial for SEI values above 2 eV/molecule.

3.3.2. Quenching after Cold Plasmas. When strong cooling ($c = 10^{-3}$) is applied, the VT nonequilibrium does not disappear for higher SEIs (see Figure 1b). Figure 5 shows the effect of different quenching times on the evolution of CO₂ conversion, for a cold plasma with an SEI of 4 eV/molecule.

In both cases, the CO₂ conversion slightly rises after quenching (i.e., superideal quenching; see the inset in Figure 3a), but it is followed by a larger drop compared to without quenching. Hence, the final CO₂ conversion is lower for both quenching options than without quenching.

Comparing this situation to Figure 3a, it seems that a VT nonequilibrium can enhance the CO₂ conversion, but for high SEI values, the conversion after plasma is higher, and more recombination can take place, resulting in a lower final CO₂ conversion. Only at low SEI values, nonequilibrium plasmas seem to benefit from quenching, yielding a clearly higher final CO₂ conversion (see Figure 3a).

3.4. Effect of Instantaneous Quenching on the Dissociation and Recombination Rates. To explain the different behavior of instantaneous quenching at low and high SEIs, we now discuss its effect on the dissociation and recombination rates, for a plasma with both low and high SEIs (0.5 and 4 eV/molecule), characterized by VT nonequilibrium and equilibrium at the plasma end, respectively. Figures 6 and 7 show the total contribution of the three most important dissociation reactions (reaction X4 in Table S1 of the SI and reactions N1 and N2 in Table S4) and recombination reactions (N4 and N5 in Table S4 of the SI) to the CO₂ conversion, plotted as positive and negative conversions. For both cases, the quenching moment that provides the maximum
CO₂ conversion (i.e., quenching at the plasma end for an SEI of 0.5 eV/molecule and quenching at maximum conversion in the afterglow for an SEI of 4 eV/molecule, respectively) will be compared with the nonquenched case. Figures S1 and S2 from the SI show the corresponding time evolutions of these rates after quenching, while Figures S3 and S4 show the evolution of the O radical density after quenching.

Figure 6 illustrates that the dissociation rates increase upon quenching, as well as the recombination rate by reaction N5. However, due to the low conversion degree (3−4%), little CO and O₂ are present, and this reaction is not important. The main reason for the higher final CO₂ conversion is the lower contribution from the recombination reaction N4, due to the faster drop in the O radical density, as shown in Figure S3 from the SI.

Figure 7 illustrates that dissociation reactions N1 and N2, as well as recombination reactions N4 and N5, are all reduced. The reduction is the most drastic for N2 and N5. Note that for these two reactions, the conversion value is higher than 100% in the case without quenching, since on average, CO₂ dissociates and recombines multiple times, but the overall net conversion is of course not above 100%. Due to the stronger reduction in the recombination rate, the final CO₂ conversion increases, as also observed in Figure 3b (yellow vs blue curve). However, Figure S2 in the SI shows that upon quenching, both recombination reactions see a fast but short increase with respect to the nonquenched case. This can be correlated with the short decrease in CO₂ conversion, which is
noticed when quenching (see the yellow vs blue curve in Figure 3b).

In the SI, we show the time evolution of the dissociation and recombination rates (Figure S5), and the contribution of these reactions to the overall CO2 conversion (Figure S6), for a nonequilibrium cold plasma with an SEI of 4 eV/molecule (i.e., strong cooling, \( c = 10^{-3} \)). The case without quenching (left) is shown as a reference. The horizontal dashed lines indicate the final CO2 conversion. In contrast to the nonequilibrium plasma of Figure 6, electron impact dissociation (X4) has a negligible contribution to the final CO2 conversion.

3.5. Effect of Instantaneous Quenching on the VDFs of CO2, CO, and O2. The dissociation reactions N1 and N2 and the recombination reaction N5 all have a high activation energy (see Table S4 in the SI). At low gas temperatures, i.e., after quenching, these reactions can only take place if the molecules have sufficient vibrational energy to overcome the activation energy. Therefore, Figures 8 and 9 illustrate the VDFs of CO2, CO, and O2 at different times after quenching for a VT nonequilibrium warm plasma with an SEI of 0.5 eV/molecule that is quenched at the plasma end (cf. Figures 6 and S1, and the red curve in Figure 3a) and an equilibrium warm plasma with an SEI of 4 eV/molecule that is quenched at maximum conversion (cf. Figures 7 and S2, and the yellow curve in Figure 3b), respectively.

In Figure 8, right before quenching (i.e., at the plasma end), the higher vibrational levels of all three molecules already have a large overpopulation (see the red curve); note that this time point is not indicated in the upper panel because of the logarithmic x-axis. This is consistent with the VT nonequilibrium for SEI below 1 eV/molecule in Figure 1a. Upon quenching, the drop in gas temperature results in a faster VV relaxation and a slower VT relaxation, leading to a higher population of higher vibrational levels of CO2, enabling the dissociation reactions N1 and N2. On the other hand, the strong vibrational overpopulation of CO and O2 also enables the recombination reaction N5 (see Figures 6 and S1). Due to the low conversion degree, CO2 is the dominant molecule in the mixture, and the rise in N1 and N2, fueled by the overpopulation of the higher vibrational levels of CO2, is dominant over the rise in recombination reaction N5. However, the VDF of CO2 depopulates much faster than those of CO and O2, explaining why the fast increase in conversion is followed by a slow drop (point 2, purple curve in Figure 8, and Figure S1 in the SI). After 1 ms (time-point 3, green curve in Figure 8), the higher vibrational levels of CO and O2 exhibit no overpopulation anymore, and the
recombination reaction N5 is no longer important (Figure S1 in the SI). The resulting drop in CO₂ conversion is a result of reaction N4, which has a low activation energy, and does not need vibrational energy to take place.

In Figure 9, the VDFs of the three molecules follow the Boltzmann distribution, since VT equilibrium was established at the plasma end (Figure 1a), and quenching occurs some time after the plasma. Due to the high gas temperature, the distribution of the higher vibrational levels is elevated but not overpopulated compared to a Boltzmann distribution. Right after quenching, the Boltzmann distribution drops due to the drop in gas temperature, but the VDFs remain close to the Boltzmann distributions before quenching (see the yellow curves; time-point 1 in Figure 9). A VT nonequilibrium is therefore created by quenching. Similar to Figure 8, the higher levels of CO₂ are somewhat overpopulated due to the increased VV relaxation. While the vibrational temperature, after quenching, of about 3050 K is very similar to the vibrational temperature of Figure 8 (i.e., 3500 K), the overpopulation of the higher and intermediate levels is much lower. Indeed, due to the much lower population of these levels in Figure 9, VV relaxation requires a longer time to significantly populate the higher vibrational levels, since the energy has to come from the lowest levels. This gives more time for VT relaxation to depopulate the VDF. Also, the conversion degree is much higher, so there is less CO₂ to dissociate, and the recombination reactions are more prominent (Figure S2 of the SI), counteracting the eventual rise in CO₂ dissociation, yielding a net drop in conversion (upper panel of Figure 9). Similar to Figure 8, the VDFs of CO and O₂ remain populated for a longer time, so recombination reaction N5 is active for a longer time (time-point 2, purple curves, and Figure S2 in the SI). Due to the higher concentrations of CO and O₂, this results in a higher recombination and a bigger drop in CO₂ conversion with respect to Figure 8.

For completeness, Figure S7 in the SI illustrates the VDFs of CO₂, CO, and O₂ for a VT nonequilibrium cold plasma at an SEI of 4 eV/molecule (i.e., strong cooling; ε = 10⁴), which is quenched at the plasma end (cf. red curve in Figure 5). 3.6. Effect of Different Quenching Cooling Rates. In previous sections, we discussed the effect of instantaneous quenching. This is the ideal case, but in reality, the gas cooling will not be instantaneous. Therefore, we discuss here the effect of different cooling rates (i.e., different cooling constants ε₂ in eq 9) on CO₂ conversion for warm plasmas, i.e., weak initial cooling (ε₁ = 10⁻³).

Figure 10 illustrates the time evolution of CO₂ conversion and gas temperature in the afterglow, for different cooling rates, for a warm nonequilibrium plasma with an SEI of 0.5 eV/molecule that is quenched at the plasma end (cf. Figure 3a).

Without quenching (i.e., weak cooling in the afterglow; ε₂ = ε₁ = 10⁻³), the gas temperature continues to rise in the afterglow (up to about 0.3 s) due to VT relaxation and recombination reactions. Upon increasing ε₂, the gas temperature drops faster (cf. Figure 10b). Hence, recombination becomes less important, resulting in a higher final CO₂ conversion (cf. Figure 10a). For a cooling constant ε₂ ≥ 10⁴, the gas temperature reduces to 300 K within 10 µs (or less) and superideal quenching becomes apparent (cf. Figure 10). Without quenching (ε₂ = ε₁), VT relaxation quickly depopulates the highest vibrational levels in the afterglow. Figure S8 in the SI shows that the VDF of CO₂ is in equilibrium within 100 µs after the plasma end. If the cooling rate is high enough (ε₂ ≥ 10⁴), the gas temperature drops faster than the VDF can relax. This limits the drop in, or even enhances, the overpopulation of the higher vibrational levels of CO₂ thus increasing the dissociation rates, as the CO₂ vibrational energy can overcome the high activation energy of the dissociation reactions N1 and N2.

Figure 11 depicts the time evolution of CO₂ conversion and gas temperature in the afterglow, for different cooling constants, for a VT equilibrium warm plasma with an SEI of 4 eV/molecule that is quenched at maximum conversion (cf. Figure 3b).
In this case, the gas temperature becomes high enough to allow for thermal dissociation of CO$_2$, even after the plasma end. However, due to slow gas cooling, the recombination reactions reduce the CO$_2$ conversion significantly (cf. Figure 11a,b). Upon higher cooling rates, the final CO$_2$ conversion increases. Note that the maximum conversion is not reached with maximum cooling rates. Indeed, when the cooling constant increases from 100 to 1000, the conversion slightly decreases (see the inset in Figure 11a). This is due to the VV relaxation that populates the higher vibrational levels of CO and O$_2$ at lower gas temperatures (Figure 9), so that they can overcome the high activation energy of the recombination reaction N5. The effect is however minor, and in general, we can conclude that for warm plasma conditions (i.e., weak initial cooling), higher cooling rates yield a higher final CO$_2$ conversion, either due to superideal quenching, further enhancing the CO$_2$ conversion (Figure 10), or simply by freezing the CO$_2$ conversion because the recombination reactions become negligible at low gas temperatures (Figure 11).

3.7. Comparison of Plasma-Based and Thermal CO$_2$ Conversion at Different SEI Values and Gas Temperatures and Effect of Instantaneous Quenching. Figure 12 illustrates the CO$_2$ conversion and energy efficiency as a function of SEI for a warm plasma ($c_1 = 10^{-3}$), i.e., warm plasma. The cooling constants $c_2$ in the afterglow are indicated in the legend.

For SEI values $\leq 1$ eV/molecule, the maximum gas temperature is too low for thermal dissociation of CO$_2$. The thermal process therefore shows negligible conversion, while the plasma process exhibits a conversion of about 0.8–5.6%, corresponding to an energy efficiency of 9.5–16.5%, mostly attributed to vibrational-induced dissociation but also electron impact dissociation.

For SEI $\geq 2$ eV/molecule, the conversion and energy efficiency of the thermal process rise, and they are similar to the plasma process, both with and without quenching. In addition, the maximum gas temperatures are also very comparable, due to the high VT relaxation that results in a VT equilibrium (see Figure 1). Upon quenching, a similar behavior takes place because in both cases a VT nonequilibrium is created, since the VDF is frozen in its Boltzmann distribution from right before quenching (see Figure 9). Similarly as in the plasma quenching case, the conversion degree slightly drops due to the higher concentration of CO and O$_2$, promoting a higher recombination (see Figures 3b and 9). However, at high SEI, the conversion and energy efficiency of the quenched thermal case perform slightly better than that of the plasma case. This is because the power goes directly to the neutral species, while in the plasma case, inefficient processes like electron impact dissociation still play a role (see Figures 6 and 7). Without quenching, the final CO$_2$ conversion

Figure 10. Time evolution of CO$_2$ conversion (a) and gas temperature (b) for different cooling rates in the afterglow for a VT nonequilibrium plasma with an SEI of 0.5 eV/molecule that is quenched at the plasma end. The plasma is initially subjected to weak cooling ($c_1 = 10^{-3}$), i.e., warm plasma. The cooling constants $c_2$ in the afterglow are indicated in the legend.

Figure 11. Time evolution of CO$_2$ conversion (a) and gas temperature (b), for different cooling rates, for a VT equilibrium plasma with an SEI of 4 eV/molecule that is quenched at maximum CO$_2$ conversion. The plasma is initially subjected to weak cooling ($c_1 = 10^{-3}$), i.e., warm plasma. The cooling constants $c_2$ in the afterglow are indicated in the legend.
remains constant at 27–28% due to the slow cooling, which promotes recombination reactions.

In Figure 13, we plot the CO₂ conversion and energy efficiency (left axis) as a function of cooling constant (ε), i.e., mimicking warm and cold plasmas at different temperatures (see curves and right axis), for plasma-based and thermal conversion, with an SEI of 4 eV/molecule, without quenching, and for instantaneous quenching at maximum conversion (i.e., the gas temperature drops immediately to 300 K).

At weak cooling (ε = 10⁻³), corresponding to a maximum gas temperature of 4150–4200 K, quenching reaches the highest conversions of 79 and 81%, corresponding to energy efficiencies of 58 and 59%, for the plasma and purely thermal case, respectively. At these high temperatures, the conversion is mostly thermal. The difference between the quenching and no quenching is also high, since the slow drop in gas temperature without quenching is detrimental to the conversion (see also Figure 2). With increasing cooling constant ε, the maximum gas temperature decreases, leading to a lower thermal conversion. Also, the difference with and without quenching decreases, as the higher afterglow cooling also leaves less time for recombination. Note that the higher conversion is already significant for moderate cooling (ε = 0.01 vs 10⁻³), corresponding to low reduction in the maximum gas temperature, as the recombination in the afterglow typically occurs over a long time (order of 1 s; see Figure 2). When the cooling ε reaches 1, the maximum gas temperature is around 2500 K, which is too low for (substantial) thermal dissociation. Therefore, the thermal CO₂ conversion and energy efficiency become negligible. The plasma-based CO₂ conversion and energy efficiency also drop. Indeed, while the gas temperature is too low for substantial thermal dissociation, it is still high enough for considerable VT relaxation, which prevents an efficient vibrational-induced dissociation. This VT relaxation becomes less important upon even stronger cooling, leading to higher vibrational-induced dissociation and a subsequent rise in the final CO₂ conversion and energy efficiency, for ε ≥ 100. These conditions correspond to cold VT nonequilibrium plasmas, where quenching does not enhance the CO₂ conversion or energy efficiency (cf. Figure 5).

It must be realized that the exact values of conversion and energy efficiency are subject to uncertainties, as mentioned in Section 2.2, and should thus be considered with caution, but the trends predicted by the model, for the effect of quenching and the comparison between plasma-based and thermal conversion, should be reliable. Indeed, our modeling results are in reasonable agreement with the model of Kotov and Koelman, predicting maximum energy efficiencies of about 40% at 4 eV/molecule in MW plasmas at comparable pressure and also stressing the dominant role of thermal conversion. den Harder et al.19 and Bongers et al.21 reported experimental energy efficiencies up to 48% for gas temperatures between 3500 and 4000 K at intermediate pressure. In addition, experiments revealed that quenching in the afterglow can enhance the energy efficiency to values around 50%.21 However, exact comparison with these experimental results is difficult due to possible deviations in pressure, ionization degree, and reduced electric field and because of limitations of the 0D model,61 like not describing transport phenomena and assuming spatial homogeneity. These transport phenomena might play an eminent role in facilitating the quenching process.19

3.8. Quenching at Higher Ionization Degree. In the previous sections, we considered an ionization degree of 10⁻⁶. In contracted MW plasmas, the ionization degree can be much higher, up to 10⁻².40 Therefore, we show here the evolution of CO₂ conversion and energy efficiency as a function of SEI for an ionization degree of 10⁻². The reduced electric field is kept at 50 Td.

Figure 13. Energy efficiency and CO₂ conversion (left axis) as well as maximum gas temperature (right axis) as a function of cooling constant ε for the plasma and purely thermal process, with an SEI of 4 eV/molecule, without quenching and with instantaneous quenching at maximum conversion.

Figure 12. CO₂ conversion ((a) left axis) and energy efficiency ((b) left axis) as well as maximum gas temperature (right axis) as a function of cooling constant (ε) for a warm plasma (i.e., weak cooling; ε = 10⁻³) and pure thermal conversion, with and without quenching at maximum conversion.
In Figure 14, we plot the conversion and energy efficiency (left axis) as a function of SEI, for the plasma and thermal process, at low cooling \((c = 10^{-3})\). The gas is instantaneously quenched at maximum conversion, and the maximum gas temperature is shown for both cases (right axis). When comparing Figure 14 with Figure 12, we can see that the conversion and energy efficiency are slightly higher at this higher ionization degree for SEI values \(\leq 2\) eV/molecule. For SEI values \(\leq 1\) eV/molecule, the temperature is too low for thermal conversion. Even at 2 eV/molecule, the plasma-based conversion and energy efficiency are higher than for thermal conversion, both with and without quenching, since a higher ionization degree results in more electron impact reactions, including more electron impact vibrational excitation, and consequently also in more vibrational-induced dissociation.\(^{29}\) Similar to the lower ionization degree of \(10^{-6}\), the highest conversions and energy efficiencies are again found for the higher SEI values (see also Figure 12). Note also that the maximum gas temperature for these higher SEIs is larger than for a lower ionization degree (see Figure 12). This is also observed in experiments.

For an SEI of 3–5 eV/molecule, the conversions and energy efficiencies are very similar for plasma and thermal conversion, both with and without quenching. However, the maximum gas temperature is much lower for the plasma process than for the thermal process. This is because more energy goes to the vibrational levels at a higher ionization degree, resulting in more vibrational-induced dissociation. Despite the difference in maximum gas temperature, and the fact that more of the plasma-based CO\(_2\) conversion takes place through vibrational-induced dissociation, the final CO\(_2\) conversion and hence energy efficiency are similar for the plasma and thermal process. The energy efficiencies at these higher SEI values are very similar to the ones at lower ionization degree (see Figure 12).

### 3.9. Quenching at Higher Reduced Electric Field

All previous results were obtained for a reduced electric field of 50 Td. As this value may be quite low for practical MW plasmas, we present here the evolution of CO\(_2\) conversion and energy efficiency as a function of SEI, for a reduced electric field of 100 Td, again for a low cooling \((c = 10^{-3})\), and an ionization degree of \(10^{-6}\), and for both the plasma and thermal processes, with and without quenching (instantaneous quenching at maximum conversion). The maximum gas temperature is shown for both cases on the right axis.

Figure 15 shows a similar evolution with increasing SEI as at lower E/N (see Figure 12). At low SEI values (SEI = 0.25–1 eV/molecule), the plasma exhibits a low conversion and energy efficiency, and there is no thermal dissociation. At SEI = 2 eV/molecule, the plasma process is less efficient than the thermal process, both with and without quenching. The same applies to higher SEI values upon quenching. This is because the higher reduced electric field makes electron impact dissociation more prominent, but the latter is less energetically favorable, explaining the lower energy efficiency than for pure thermal conversion. At higher SEI values (SEI = 3–5 eV/molecule), the plasma exhibits a low conversion and energy efficiency, and there is no thermal dissociation.
molecule), the plasma and thermal process without quenching become equally efficient because the slow drop in gas temperature reduces the CO$_2$ conversion to a value of 27–28%.

4. CONCLUSIONS

We demonstrated the potential of quenching for plasma-based CO$_2$ conversion, by means of chemical kinetics modeling. We consider the so-called warm plasmas at a pressure of 100 mbar, an ionization degree of $10^{-6}$, and a reduced electric field of 50 Td, but our model predicts that the trends are the same at higher ionization degree ($10^{-4}$) and reduced electric field (100 Td).

We calculated the gas and vibrational temperatures, and the CO$_2$ conversion, as a function of time in the plasma, at weak and strong cooling, mimicking warm and cold plasma conditions, respectively. At weak cooling, high conversion degrees were reached for large SEI values (i.e., up to 96% at SEI = 5 eV/molecule), due to the high gas temperature (∼4000 K), leading to VT equilibrium and thus thermal conversion. However, without additional cooling in the afterglow, the conversion drops dramatically due to the backward (recombination) reactions, leading to a low overall CO$_2$ conversion of 27–28%.

We examined the effect of quenching for warm and cold plasma conditions, and we explained the results based on the dissociation and recombination reactions and the VDFs of the CO$_2$, CO, and O$_2$ molecules. For warm plasma conditions at low SEI, yielding VT nonequilibrium, our model predicts that it is best to quench at the plasma end. In this case, quenching can even further enhance the dissociation by the reactions CO$_2$ + M → CO + O + M and CO$_2$ + O → CO + O$_2$ due to the overpopulation of the CO$_2$ vibrational levels, which helps overcome the activation energy of these reactions. This process is called superideal quenching. For cold plasma conditions at high SEI, which also result in VT nonequilibrium, quenching also enhances the dissociation, but it promotes the recombination reactions even more, leading to a lower final CO$_2$ conversion. For warm plasma conditions at high SEI, yielding VT equilibrium, the high gas temperature enhances the CO$_2$ conversion even after the plasma, so it is more favorable to quench at maximum conversion. Upon fast quenching, both dissociation and recombination reactions stop, so the CO$_2$ conversion is frozen, leading to a much higher final CO$_2$ conversion and overall energy efficiency than without quenching (even up to a factor 3 enhancement).

We also evaluated the effect of different cooling rates in the afterglow, demonstrating overall the importance of fast cooling, although at high SEI values, we also have to account for overpopulation of the higher vibrational levels of CO, which can overcome the activation energy of the recombination reaction, therefore leading to more recombination, resulting in a slight drop in the overall conversion.

Finally, we compared the performance of plasma-based CO$_2$ conversion with purely thermal conversion for different SEI values and for warm (i.e., low cooling) and cold (i.e., high cooling) plasmas, as well as the effect of quenching, on both the CO$_2$ conversion and energy efficiency. For warm plasmas at SEI ≥ 2 eV/molecule, yielding gas temperatures of 3000–4000 K, the plasma-based conversion is mostly thermal, and quenching can enhance the conversion and energy efficiency by up to a factor 3. For low-SEI plasmas at lower temperatures, when thermal conversion is negligible, the overall conversion and energy efficiency are quite low, and in addition, quenching becomes less efficient. For cold plasmas in VT nonequilibrium at high SEI, quenching can even decrease the final CO$_2$ conversion.

In summary, our model provides interesting insights into how quenching can improve the CO$_2$ conversion and energy efficiency for various plasma conditions and cooling conditions. We showed that the highest conversion and energy efficiency can be reached for thermal conversion at high gas temperatures in the plasma, followed by fast quenching. Such quenching can be realized when expanding the flow after the plasma, as demonstrated experimentally, but it may also be accomplished by mixing with a cold gas. In addition, the quenching might be combined with heat recovery for preheating the gas entering the plasma reactor, so that less plasma power is wasted for gas heating and all of the power could be used for the conversion process.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.0c04257.

Reactions included in the models as well as their rate coefficients or cross sections; scaling laws; effect of instantaneous quenching on the evolution of the dissociation and recombination rates for a plasma subjected to low cooling; effect of instantaneous quenching on the O atom density; effect of instantaneous quenching on the dissociation and recombination rates for a plasma subjected to strong cooling; effect of instantaneous quenching on the VDFs of CO, CO, and O$_2$ for a plasma subjected to strong cooling; relaxation time of the CO$_2$ VDF at low SEI (PDF)

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Notes

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