Transition metal carbides as novel materials for CO₂ capture, storage, and activation†

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The capture and activation of the greenhouse gas carbon dioxide (CO₂) is a prerequisite to its catalytic reforming or breakdown.

Here we report, by means of density functional theory calculations including dispersive forces, that transition metal carbides (TMCs) can be utilized to uptake and activate CO₂ on their most-stable (001) surfaces with considerable adsorption strength. Estimations of adsorption and desorption rates predict a capture of CO₂ at ambient temperature and even low partial pressures, suggesting TMCs as potential materials for CO₂ abatement.

Carbon dioxide (CO₂) is an active greenhouse gas with a rising atmospheric concentration currently approaching a value of 400 ppm. This raises concerns about serious consequences such as climate change and ocean acidification. As economies and energy demand are projected to grow in the future, a reduction of CO₂ emissions nowadays only seems possible with efficient technologies for carbon capture and storage (CCS). Employing absorbents or solvents, CO₂ re-use into a saleable product has been suggested to be an appealing route to contribute to climate change mitigation, though the field is still in its infancy and questions over overall system efficiency have been raised. Active catalytic systems for such processes are known, yet most of them are made of scarce and precious metals, normally supported on high surface area porous oxides or sulfides. The effective large-scale implementation on this basis then still remains a great challenge.

In the search for new active catalysts, numerous endeavours address the adsorption, activation, and subsequent catalytic conversion of CO₂ on different systems. Note that a relatively high adsorption energy is needed for CO₂ to stick on a catalyst surface, which is made difficult by the high stability of CO₂ molecules. Moreover, when possible, CO₂ adsorption normally goes without significant activation, a process that is known to require charge transfer from the substrate resulting in a concomitant bending. The possible capture/activation of CO₂ has been theoretically tackled via first principles calculations on metals, metal oxides, graphene-based materials, sulfides, zeolites, and metal–organic frameworks, to name a few.

While the surface chemistry of transition metal carbides (TMCs) was described as comparable to that of Pt-group metals by Levy and Boudart, these materials have received considerable attention in catalysis. TMC catalysts have been used in a wide range of reactions like methane dry reforming, conversion of methane to synthesis gas, desulfurization, the water-gas-shift reaction, and CO oxidation, demonstrating chemical robustness and oftentimes catalytic properties similar to precious Pt-group metals.

The interaction of CO₂ and TMC surfaces has been recently reviewed, although one must acknowledge that most of the studies so far focused on Mo₂C, MoC, WC, and TiC. From these studies it is now known that some surfaces of Mo₂C, δ-MoC, and TiC are highly active for CO₂ hydrogenation, producing methanol, methane, and carbon monoxide (CO) in different ratios. The catalytic activity of these TMCs has been found to be further enhanced by using them as supports for transition metal nanoclusters. Regarding CO₂ adsorption and activation, density functional theory (DFT) calculations have shown that upon interaction with these surfaces a surface-bound anionic CO₂ is formed. The key role of this activated intermediate in further reactions.

Apart from the above-mentioned extensively studied systems, information on the CO₂ surface chemistry involving other TMCs is rather limited. A recent study carried out by Porosoff and coworkers found CO₂ hydrogenation activities for TiC, ZrC, NbC, TaC, WC, and Mo₂C. These results largely motivated the present theoretical systematic investigation of CO₂ adsorption and activation on TMC surfaces. To facilitate a logical comparison of the results and to circumvent structural aspects and related effects, we restrained our study to TMCs with a 1 : 1...
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Table 1 Adsorption energies, \( E_{\text{ads}} \), and differences, \( \Delta E_{\text{ads}} \), of CO\(_2\) and TopC sites of different TMC(001) surfaces. All values are given in eV.

<table>
<thead>
<tr>
<th></th>
<th>TiC</th>
<th>ZrC</th>
<th>HfC</th>
<th>NbC</th>
<th>TaC</th>
<th>( \delta)-MoC</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMC</td>
<td>−0.81</td>
<td>−1.56</td>
<td>−1.62</td>
<td>−0.87</td>
<td>−1.21</td>
<td>−1.20</td>
</tr>
<tr>
<td>TopC</td>
<td>−0.83</td>
<td>−1.60</td>
<td>−1.65</td>
<td>−0.70</td>
<td>−0.94</td>
<td>−1.03</td>
</tr>
<tr>
<td>( \Delta E_{\text{ads}} )</td>
<td>0.02</td>
<td>0.04</td>
<td>0.03</td>
<td>−0.17</td>
<td>−0.27</td>
<td>−0.17</td>
</tr>
</tbody>
</table>

From the tabulated values, trends for adsorption energies can be withdrawn. For instance, on 3d, 4d, and 5d TMC(001) \( E_{\text{ads}} \) rise in magnitude when going down a group. Aside, the adsorption strength decreases when moving along a d series. Indeed, the almost vanishing interaction of CO\(_2\) on the VC(001) surface can be understood from the decrease of more than 0.7 eV when going ZrC \( \rightarrow \) NbC, applying the same trend in going from TiC to VC would result in an \( E_{\text{ads}} \) value of at most 0.1 eV at the PBE level, in full concordance with the obtained results. One must warn, however, that these trends may not hold for other stable surfaces, as the surface structure can strongly alter the adsorptive properties.\(^{15,17}\)

Last but not the least, it is mandatory to highlight that, compared to PBE \( E_{\text{ads}} \) values, PBE-D3 predicts more stable minima by 0.21–0.32 eV. Still, the main adsorption driving force comes from non-dispersive interactions between the surface and the adsorbate, as already found for other TMCs as well.\(^{10}\)

In the \( \delta\)-MoC(001) surface a fair comparison between D3 and D2 dispersion corrections reveals that, compared to PBE,\(^{33}\) the latter yields a slightly larger stabilization (0.4–0.6 eV).

Note that on both MMC and TopC sites, and regardless of the TMC, the adsorbed CO\(_2\) molecule exhibits a similar geometry, see examples of the adsorbate structures on the TiC(001) surface in Fig. 1. For each system, a complete list of structural descriptors, including bond lengths and molecular angles, is reported in Table S1 of the ESL.\(^\dagger\) Briefly, and in agreement with earlier results, C \( \leftrightarrow \) C and metal \( \leftrightarrow \) O interactions between CO\(_2\) and surface site atoms imply bond lengths on the order of 1.46–1.50 and 2.09–2.37 Å, respectively. Compared to MMC sites, the latter are circa 0.2 Å longer on TopC. The adsorbed CO\(_2\) species always bends with angles \( \gamma(\text{OCO}) \) between 120.1 and 128.8° which is a strong evidence of CO\(_2\) activation by charge transfer from the underlying TMC surface.\(^{10}\)

This is also consistent with C–O bond lengths between 1.29 and 1.32 Å, significantly longer than the 1.176 Å value of CO\(_2\) in a vacuum.

Given the relatively high adsorption energies, there is a clear interest in estimating the temperature fringe below which these
TMCs could capture and accumulate CO₂. This can be easily assessed from adsorption and desorption rates, \( r_{\text{ads}} \) and \( r_{\text{des}} \), respectively, which can be estimated in the framework of the transition state theory (TST). Hence, \( r_{\text{ads}} \) and \( r_{\text{des}} \) values have been calculated over a temperature range up to 1000 K with necessary quantities obtained from the above commented DFT calculations. The utilized equations and approaches are duly explained in the ESI.\(^{1} \) In a nutshell, the adsorption rate depends on the impingement of CO₂ to the surface and therefore, on the CO₂ partial pressure \( (p_{\text{CO₂}}) \). In this sense, and for each system, \( r_{\text{ads}} \) has been calculated for (i) the current atmospheric partial pressure of CO₂, \( p_{\text{CO₂}} = 40 \text{ Pa} \),\(^{44} \) (ii) for \( p_{\text{CO₂}} = 15,000 \text{ Pa} — 0.15 \text{ bar} — \), a benchmark value for postcombustion exhaust gases,\(^{5} \) and (iii) \( p_{\text{CO₂}} = 10^5 \text{ Pa} — 1 \text{ bar} — \), a partial pressure regime of interest for pure CO₂ stream generation from a CCS system.\(^{7} \)

The desorption rates largely depend on the adsorption strength, and so we obtained four curves for each TMC, two for MMC and two for TopC sites, each of them as evaluated using PBE or PBE-D3 calculations respectively. DFT calculations are widespread used for evaluating adsorption strengths on a plethora of substrates, whilst TST is frequently used for simulating desorption processes as in temperature programmed desorption; a recent study for CO adsorption/desorption on a similar system (TiC nanopowders) provides a paradigmatic example.\(^{45} \)

The estimated \( r_{\text{ads}} \) and \( r_{\text{des}} \) rates are graphically shown for the two fringe cases TiC and ZrC, see Fig. 2a. These are representative TMCs with low and high \( E_{\text{ads}} \) values. Note that intersection points of \( r_{\text{ads}} \) and \( r_{\text{des}} \), define temperatures below which adsorption prevails and, consequently, CO₂ accumulates on the TMC surface. Thus, for TiC(001) at 40 Pa CO₂ partial pressure, the lowest and highest intersection temperatures are 211 K for the MMC site as calculated using PBE and 329 K for the TopC site (PBE-D3), marked as \( T_1 \) and \( T_2 \), respectively. At a CO₂ partial pressure of 0.15 bar the intersection temperatures are shifted to higher values of 262 to 414 K (\( T_3 \) and \( T_4 \)), and so compare to the desorption temperature of 323 K reported for the zeolite 13X benchmark material.\(^{5} \) At a CO₂ partial pressure of 1 bar, the desorption process will likely occur in the 285–452 K range (\( T_5 \) and \( T_6 \)), temperatures above which a surface regenerate for further use becomes feasible.

Thus \( T_1 \rightarrow T_2, T_3 \rightarrow T_4, \) and \( T_5 \rightarrow T_6 \) can be interpreted as temperature ranges in which the TMC(001) surface looses its ability to initially capture and accumulate CO₂ when annealing. This seems reasonable as dispersive force contributions to the adsorption energies might be overestimated by PBE-D3 and we expect the real desorption rate to lie somewhere in between. The analysis for the rest of investigated TMCs is summarized in Fig. 2b allowing for an easy comparison. Thus, ZrC- and HfC(001) surfaces display very elevated temperature ranges, in accordance with their high adsorption energies. The temperature ranges for TiC, NbC, TaC, and δ-MoC are lower but still in the range or even well above room temperature. Even though no CCS experiments on TMCs are available, it has been experimentally proven, as mentioned above, that some TMCs catalyse the CO₂ hydrogenation to methanol at 500–600 K,\(^{33–35} \) implying that CO₂ must become an adsorbed surface activated moiety.

![Fig. 2](image_url) (a) Calculated rates for desorption and adsorption of CO₂ on TiC and ZrC(001) surfaces. On TiC, marked points with \( T_1–T_6 \) labels in (a) and (b) show how desorption temperature ranges in (b) have been obtained. Legend for (a): green, grey, and blue, adsorption rates on a single site per time unit for a CO₂ partial pressure of 40, 15 × 10³, and 10⁵ Pa, respectively. Black and red: desorption rates per site and time unit from MMC and TopC sites. PBE (solid) and PBE-D3 (dashed) values are provided. Legend for (b): green, grey, and blue bars belong to desorption temperature ranges for CO₂ partial pressures of 40, 15 × 10³, and 10⁵ Pa, respectively.

Note that, as stated above, in most of these cases, MMC sites show a stronger adsorption compared to TopC sites, with a concomitant widening of the calculated temperature range. One must keep in mind that upon annealing, these systems may well first partially lose their adsorption capability, i.e. those TopC sites would be firstly depopulated, yet here lateral interactions are being neglected, and so the obtained rates better correlate with the initial stages of CO₂ capture. Nevertheless, at low coverages on Ni surfaces,\(^{17} \) lateral interactions for adsorbed CO₂ have been found not to be repulsive and so initial desorption rates are likely not underestimated. All in all, most of the TMC(001) surfaces are envisaged to capture and accumulate CO₂ at ambient and even elevated temperatures.

In summary, periodic DFT calculations carried out at the PBE level and also including dispersion (PBE-D3) showed that CO₂ molecules adsorb and get activated on a range of stable (001) surfaces of transition metal carbides (TMC–TM = Ti, Zr, Hf, Nb, Ta, Mo). Two competitive adsorption sites are identified (MMC and TopC) where CO₂ chemisorbs in a bended (activated) geometry. Dispersion corrected adsorption energies are found to be favourable and lie in a range of −0.70 to −1.65 eV, depending...
on the TMC and surface site. These adsorption strengths are therefore considerably high, and despite dispersive forces are responsible of 0.21–0.32 eV, the major attachment force has a chemical nature. Adsorption and desorption rates, as predicted from DFT data, show that these materials theoretically can adsorb CO$_2$ up to elevated temperatures at even low partial pressures. Thus TMCs appear to be ideal materials for CO$_2$ capture and abatement, and actually given its activated adsorption appealing for using them as catalysts for CO$_2$ conversion as well.

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Notes and references