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Effect of Internal Donors on Raman and IR Spectroscopic Fingerprints of $\text{MgCl}_2/\text{TiCl}_4$ Nanoclusters Determined by Machine Learning and DFT

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Abstract: To go deep inside the origin of Ziegler-Natta catalysis we need to fully understand the structure and properties of $\text{MgCl}_2/\text{TiCl}_4/\text{ID}$ nanoclusters. In this work $\text{MgCl}_2/\text{TiCl}_4$ nanoplatelets derived by machine learning and DFT calculations have been used to model the interaction with ethyl-benzoate EB (as internal donor) with available exposed sites of binary $\text{Ti}_x\text{Cl}_y/\text{MgCl}_2$ systems. The influence of vicinal Ti_2Cl_8 and coadsorbed TiCl_4 on energetic, structural and spectroscopic behaviour of EB has been considered. The adsorption of homogeneous-like TiCl_4EB and $\text{TiCl}_4(\text{EB})_2$ at the various surface sites have been also simulated. Moving in the framework of Density Functional theory, we adopted B3LYP-D2 and M06 functionals. The adducts have been characterized by computing IR and Raman spectra that have been found to provide specific fingerprints useful to identify surface species; IR spectra have been successfully compared to available experimental data.

Keywords: polymerization catalysis, machine learning, nanoclusters, DFT, Raman spectrum, IR spectrum, Lewis bases

1. Introduction

In real heterogeneous catalysts the complexity of the system is crucial in determining activity, selectivity and life-time, that represent the main parameters defining the performance of the catalyst; hence together with active species, dopants, promoters or modifiers need to be carefully chosen and their role fully understood. With regard to MgCl_2 supported Ziegler Natta catalysts (ZNC) for olefin polymerization, the electron donors (Lewis base molecules) have been providing that supporting activity since 1960s with significant increase of both catalyst productivity and stereospecificity. [1–4]

This very diffuse practice has been continuously improved during last decades by industry mostly in a trial and error process.

The coadsorption of electron donors (generally bidentate organic molecules like diesters, diethers, and dialkoxysilanes) determine the local environment of the transition metal, behaving as ancillary ligand(s) in molecular catalysts able to enhance their regio- and enantioselectivity at the propene insertion in a defective context. They also determine the molecular weight distribution of the polymer, the distribution of active sites, the morphology of the catalyst [1,5–22] and the properties of the produced polymers. The industrial research in the last decade was devoted to the optimization of new couples internal donor/external donor (ID, ED) [21,22] because generally pairs of electron donors are used. [6,9,23,24]

Both experimental [25] and computational [26–32] research has been performed to get insights in the structure of ZNC at the atomic level, with main attention to the place and ways of binding of the ID, ED and their effects on the structure of the active sites.

Several contrasting hypotheses have been formulated about the role of the IDs. Some researchers claimed that the ID coordinates to a vacancy of the titanium centers, thus forming new active and stereo-specific sites;[33–36] it is also supported by the formation of homogeneous complexes between TiCl_4 and many Lewis base molecules. The other opposite hypothesis recall the historical one: the ID only interacts with the MgCl_2 support, indirectly determining the place and the distribution of the TiCl_4 moieties on the MgCl_2 surfaces and preventing the formation of aspecific sites.[33–40] Nowadays the most trusted and accepted hypothesis for ID, is the one known as the "coadsorption model": ID would coordinate to the MgCl_2 surfaces close to the titanium active sites, thus influencing their steric and electronic properties with no direct bonding.[22,27,28] We can assume a total consensus of scientific community on the nanostructured and disordered nature of the active δ form of MgCl_2 exposing surfaces and defective sites to adsorption of both TiCl_4 and internal donors to form Ziegler-Natta precatalyst. The hypothesis supported by experimental investigation, mostly by TEM microscopy, encountered a lot of limits in characterization due to these nanosize and disordered features, heterogeneity of sites, dilution of the active centres, air sensitivity. Theoretical and computational chemistry provided crucial insights in δ - MgCl_2 , made feasible the modelling of each component and the analysis of different catalytic sites.[41–43]

Molecular mechanics investigation identified the (110) and the (104) lateral surfaces as the surfaces for the adsorption of monomeric TiCl_4 and stereo-selective dimeric Ti_2Cl_8 species, respectively.[44] However, together with some experimental inconsistencies, DFT studies have definitely set aside those models, after puzzling DFT predictions of adsorption energies of Ti_xCl_y species on regular MgCl_2 surfaces and thanks to more recent reliable descriptions of the morphology of the δ - MgCl_2 particles. [45–53] Theoretical calculations on MgCl_2 bulk and surface structure exploiting periodic Density Functional Theory approaches including London interactions pointed out that MgCl_2 crystals in the absence of adsorbates mainly expose the (104) or other pentacoordinated surfaces, whereas the picture drastically changes in the presence of adsorbates (i.e. electron donors); in fact, regular MgCl_2 crystals expose the (110) surfaces at a relatively large extent when they grow in the presence of Lewis bases (small molecules such as methanol, ethanol, and dimethyl ether) or in presence of alkoxy silanes employed as external donors. [32,54] Thanks to Bravais' law it was possible to identify the most stable surfaces in terms of (surface) Gibbs Free Energy: (107), (012), (101), (015) and (110) ones. [55] The combination of a careful structural analysis of the supporting δ - MgCl_2 nanocrystals by synchrotron X-ray total scattering [42] with the definition of Wulff's polyhedron of MgCl_2 crystallites suggested that ball-milling of naked MgCl_2 produces a larger total surface area and modifies the relative extension of the surfaces, too. That would increase the stability of the lateral surfaces exposing strongly acidic tetracoordinated Mg^{2+} sites (i.e., the (110), (012) and (015) ones) at the expenses of the basal (001) one. The polar plot identified the presence of possible defective edges involving the (110) surface.[43] Inelastic Neutron Scattering (INS) spectroscopy,[56] able to sample all the first Brillouin zone (FBZ) and thus predicting the dynamical behaviour of materials, proved to be a powerful method to understand nano-sized systems and measure the degree of disorder of δ - MgCl_2 in terms of both translational and rotational contribution. A step forward to determine the morphology of platelets with dimensions comparable to those of primary particles, typically adopted in industrial catalysis (2-3 nm), have been made by us thanks to the adoption of a genetic algorithm; the calculations have been performed both in presence and in absence of TiCl_4 . [57] Differently than in the case of naked MgCl_2 where the evolutionary plots allowed us to identify structures where the majority of sites are pentacoordinated Mg^{2+} sites, in presence of TiCl_4 (in a ratio 50/3 $\text{MgCl}_2/\text{TiCl}_4$) couples of tetracoordinated sites are present after the reconstruction of nano-particles. That occurrence represents the topological requirement for the formation of octahedral Ti species that will start the catalytic process. That "motiv" confirms the presence of inter-surfaces edges as possible favorable place for stereo-selective active

sites as also reported in previous computational studies adopting traditional Density Functional methods. [31,43] FT-IR spectroscopy of probing carbon monoxide also was used to estimate the presence and the relative extension of MgCl_2 surfaces. [55,58] Very recently the vibrational Far-IR and Raman simulations have been carried out on large models of MgCl_2 nanoclusters identified through a non-empirical structure determination to discuss the effect of the binding of various precatalytic complexes Ti_xCl_y on IR and/or Raman features. IR features do not allow to clearly identify the presence of a certain adduct but in Raman a drastical change occurs after TiCl_4 adsorption. The spectroscopic features identified are specific of TiCl_4 adduct on 110-like tetracoordinated Mg^{2+} sites and different in shape and frequencies from those obtained in the case of the dimeric adduct on pentacoordinated sites. Hence Raman spectroscopy provides clear fingerprints to recognize each adduct on defective positions on nanoplatelet of Ziegler-Natta catalysts.

In this work, we make a step forward towards complexity and pass to ternary donor $\text{MgCl}_2/\text{TiCl}_4$ systems; we looked to different way of binding of ethyl benzoate (EB) in relation to available Mg^{2+} sites of attachment and respect to positions of Ti_xCl_y complexes, thus following the still open hypotheses on the mechanism that donors adopt for affecting polymerization catalysis. Differently than previous papers, mostly addressing perfect surfaces (or steps between them), we can here investigate ways of adsorption on defective positions on nanostructured support with sites formed after a reconstruction in presence of TiCl_4 .

Starting from model nanoplatelets (50MgCl_2 and $50\text{MgCl}_2/3\text{TiCl}_4$) identified by non-empirical structure prediction based on the genetic algorithm and DFT, as reported in recent papers of us [57,59,60] different way of binding were proposed and analysed, vibrational and Raman simulations were carried out to identify whether or not it is possible to identify clear fingerprints of many possible adducts donor/ TiCl_4 on nano-shaped MgCl_2 . The possible formation of homogeneous-like TiCl_4EB and TiCl_4EB_2 species was also considered. In the present paper, for simplicity, we address the location of ethyl benzoate (EB) on MgCl_2 and $\text{MgCl}_2/\text{TiCl}_4$ precatalyst, even if EB is not so largely adopted in industrial process. For sure, the number of experimental works pointed towards EB [61–65] by far overcomes studies on any other electron donor, make it easy to compare results with the literature reports at least in IR case. On the possibility to distinguish between different complexes, the Raman response for sure appears more informative but, at this very moment, experimental data are still not available on these systems even if groups are working on this topic and we foresee a future comparison between simulated and experimental data. Although the present findings are limited to the case of EB, the same approach can be easily applied to industrially more relevant electron donors and we are also immediately going to reiterate the analysis to ternary systems fully obtained by machine learning techniques that is nanoplatelet of 50MgCl_2 reconstructed in presence of both TiCl_4 and IDs.

2. Computational Models and DFT Calculations Details

As support for the adsorption of EB we adopted models of 50MgCl_2 and $50\text{MgCl}_2/3\text{TiCl}_4$ nanoplates, computed by employing non-empirical structure determination thanks to a software that combines global structure search, based on a genetic algorithm, and local geometry optimization by means of DFT[59,60]. The size of nanoclusters is about 3 nm in diameter, comparable with the experimental size of the catalyst primary particles (ca. 2.4-4.0 nm)[42]; at the meantime the ratio between TiCl_4 molecules and MgCl_2 units equal to 50/3 corresponds to 2.69 wt%, similar to Ti content in a typical Ziegler-Natta catalyst.[59,60,66,67] The DFT[68,69] calculations reported in this paper were mainly based on the B3LYP global hybrid functional,[70,71] as implemented in the CRYSTAL program.[72,73] London interactions strongly influence the description of these materials containing not only Ti_xCl_y species but also organic molecules; hence, accounting for long-range correlation, was mandatory. The semi-empirical DFT-D2 approach of Grimme

and co-workers [74,75] was adopted together with the hybrid B3LYP functional; it is well assessed that combined procedure may successfully describe surface interactions and adsorption process also including organic molecules.[76,77] Single point energy calculations were run with M06 functional [78] on B3LYPD-2 minima.

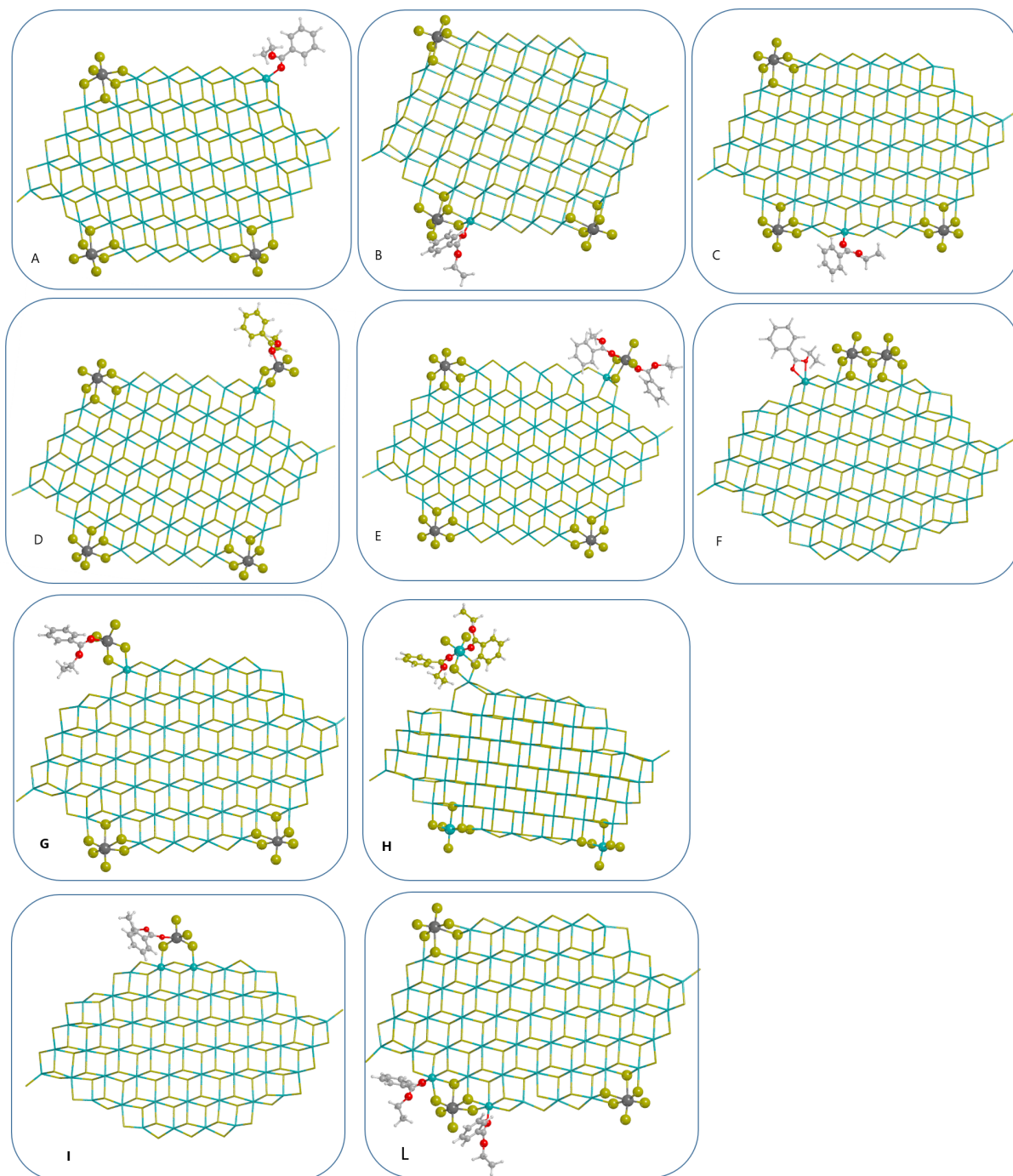


Figure 1. Coadsorption models of nanoplatelets ($50\text{MgCl}_2/\text{TiCl}_4$, see text for details) and EB with different way of binding (A-C); models of homogeneous-like $\text{TiCl}_4(\text{EB})_x$ ($x = 1, 2$) complexes on the same platelet (D, E, G, H) and models obtained after the coadsorption of EB and Ti_2Cl_8 on naked MgCl_2 nanoplatelet (F, I). Optimizations have been performed at B3LYP-D2/TZVP level. Chlorine, Magnesium and Titanium atoms are represented in green, yellow and dark grey, respectively. All generic atoms belonging to the nanoplatelet are represented as sticks, whereas atoms of TiCl_4 and EB molecules and Mg atoms involved in the adsorption process are reported as balls.

Split valence triple-zeta basis sets plus polarization (TZVP) of Gaussian type functions have been specifically customized and adopted for all the elements (Mg, Ti and Cl atoms).[51] Ahlrichs VTZ plus polarization quality [79] have been adopted for the adsorbed organic counterparts. The estimated energies have been corrected for Basis Set Superposition Error (BSSE). To inspect the molecular vibrational shifts of main modes involving CO group, O-C-O and TiCl_4 due to both the interaction with the support and the reciprocal interaction of organic and inorganic molecules, we run geometry optimizations followed by harmonic vibrational frequency calculations of the molecular adducts together with the corresponding IR and Raman intensities. Details about the computational set up and the calculations of the vibrational frequencies can be found in previous works by us [57,80,81].

3. Results and Discussion

3.1. Models of ternary systems $\text{MgCl}_2/\text{TiCl}_4/\text{EB}$

The models of coadsorption of EB and TiCl_4 on nanoplatelet of MgCl_2 may be collected in three groups.

We built a first group of models (A, B, C, L in Figure 1) where the EB is added in different positions with respect to TiCl_4 aiming to investigate the effect of reciprocal perturbation on the vibrational response of both internal donor and precursor of active species. Here, due to the heterogeneity of sites, the adsorption of both TiCl_4 moieties (and its dimeric form Ti_2Cl_8) and EB on defective sites has been possible with many combinations of reciprocal positions donor/precatalytic species: edge-like for one between them, EB next to Ti moieties, EB far enough from Ti species to remain more or less unperturbed.

A second group (models D, E, G, H) accounts for the possibility of adsorption of homogeneous-like complexes, already present in reaction medium, to nanoclusters with different Ti/EB ratio, i.e. TiCl_4EB and $\text{TiCl}_4(\text{EB})_2$ on various sites. Previous calculation on internal donor plus TiCl_4 adducts have been carried out by some of us on perfect surfaces at different degree of coverage, in that case we considered tetracoordinated Mg^{2+} ions of the type exposed on (110) surface [61] and (107) surfaces to adsorb in different way of binding EB on perfect surfaces.

A third group of models attains the $\text{Ti}_2\text{Cl}_8\text{EB}$ complexes (F, I) on pentacoordinated Mg^{2+} sites because historically these type of adducts emulate those formed on (104)-like surfaces claimed to be able to bind Ti_2Cl_8 , that was previously identified as the stereoselective active species for polypropylene.[38]. Although QM calculations proved that adsorption of Ti_2Cl_8 dimers on MgCl_2 surfaces is not feasible,[51] the presence of the donors might modify the energy of the adsorption process, giving a new chance for the adduct formation.

In the case of adducts $\text{EB}/\text{Ti}_2\text{Cl}_8$, after the adsorption on a penta-coordinated row of Mg^{2+} sites; the dimer breaks in an adsorbed TiCl_4EB complex and a TiCl_4 molecule physisorbed on the row, after the removal of physisorbed TiCl_4 molecule, the TiCl_4EB adsorbed on cluster has been investigated further as Model I. Model F presents a molecule of EB bound to four-coordinated Mg^{2+} and adjacent to Ti_2Cl_8 dimer.

For most reasonable models, the IR and Raman vibrational frequencies have been calculated and reported in Figure 2 whereas the energetics is reported in Table I (in terms of total electron energy) together with some reaction energies.

3.2. IR and Raman response of EB/TiCl_4 on nanoplatelet

3.2.1. IR simulations

We first recap main IR features of EB and $\text{MgCl}_2/\text{Ti}_x\text{Cl}_y$. The IR spectrum of EB is reported in panel a of Figure 1. The spectrum is overlooked by two intense peaks at 1766 and 1277 cm^{-1} ; the band at 1766 cm^{-1} is assigned to $\nu(\text{C}=\text{O})$ stretching mode, whereas that at 1277 cm^{-1} , coupled with the bands at 1130 and 1140 cm^{-1} , is due to a $\nu(\text{C}-\text{O})$ stretching mode in the (C-O-C). Most of the other weak bands in the spectrum are related to C-H stretching (about 3000 cm^{-1}) or deformation modes of the phenyl ring and of the

ethyl group, (e.g., the weak bands at 1200 and 1098 cm^{-1} is associated with -CH bending of the phenyl ring whereas 1334, 1349 cm^{-1} modes are due to ring deformation and the very weak bands at 1403 and 1432 cm^{-1} are associated to the wagging of CH_3). It is very widely reported that the positions of the absorption bands $\nu(\text{C-O})$ and $\nu(\text{C-O-C})$ change upon EB complexation, whereas the others are much less sensitive. Due to the fact that we investigate a coadsorption with Ti_xCl_y , also the region 400-650 cm^{-1} has been here reported for the first time since our recent investigations on binary systems $\text{TiCl}_4/\text{MgCl}_2$ revealed that the region cannot be neglected in IR spectra and even more in Raman analysis.

Concerning the IR response of $\text{MgCl}_2/\text{Ti}_x\text{Cl}_y$ nanoplatelets, it covers a region between 200 and 500 cm^{-1} [57]; although the 200-400 cm^{-1} region refers to bulk modes, strongly dependent on the particle shape and size, the 400-500 cm^{-1} region contains fingerprints that can be easily correlated to surface sites and adsorbed species: peaks at 429 and 445 cm^{-1} related to exposed tetracoordinated Mg^{2+} typical of 110 surfaces, a couple of bands at 465 and 485 cm^{-1} attributed to symmetric and antisymmetric stretching of Ti-Cl bonds in supported TiCl_4 and 458, 476 and 495 cm^{-1} bands for supported Ti_2Cl_8 . We considered now the adsorption of EB, according to the models sketched in Figure 1. IR spectra are reported in Figure 2, by considering separately the two vibrational regions 400-650 cm^{-1} and 1200-1750 cm^{-1} in order to better highlight the Ti_xCl_y and the EB response. Panels b, c and d of Figure 2 refer to model A-C that corresponds to EB coadsorption at almost isolated tetracoordinated and pentacoordinated Mg^{2+} (panels b and d) and EB coadsorbed at a pentacoordinated Mg^{2+} in proximity of a TiCl_4 surface specie of the precatalist (panel c). Panels e, and f of Figure 2 refer to models D-E of Figure 1. Panel g refers to model F. In Model D and F the TiCl_4/EB and $\text{TiCl}_4/(\text{EB})_2$ homogeneous-like complexes are loosely bound to the MgCl_2 surface through bridging chlorine atoms.

The main features of the spectra are:

- the $\nu(\text{C=O})$ vibration that dominates the IR spectrum and is located at 1683, 1697 and 1695 cm^{-1} for models A-C, downshifted by 83, 69 and 71 cm^{-1} in reference to gas phase, respectively. Two very weak satellite bands are also observed at 1638 and 1615 cm^{-1} that corresponds to C-C stretching modes of the phenyl ring. In the presence of a TiCl_4/EB adduct the C=O stretching mode undergoes a huge redshift to 1587 cm^{-1} (medium); it is further redshifted to 1556 cm^{-1} and coupled with a very weak band at 1585 cm^{-1} in the case of adduct E: they correspond to the antisymmetric and to the symmetric coupling of the two C=O stretchings;
- the $\nu(\text{C-O})$ signal drops its intensity with respect to free EB and is splitted into three components at 1308, 1323, 1336; 1312, 1332, 1348 and 1313, 1329, 1344 cm^{-1} for models A-C, respectively due to coupling with CH_2 twisting and phenyl H modes; however for model A the triplet of bands have comparable intensity, whereas for model B and C the band at higher wavenumber dominates in the triplet. A similar group of bands is observed for models D and E with the most intense peak located at 1362 cm^{-1} ;
- a new weak band appears at 1414, 1418, 1417 cm^{-1} for models A-C, and at 1427 cm^{-1} for models D-E, is associated to the wagging of $-\text{CH}_3$ and therefore slightly perturbed by different EB binding mode; the band is weak but it can be clearly identified in the IR spectra;

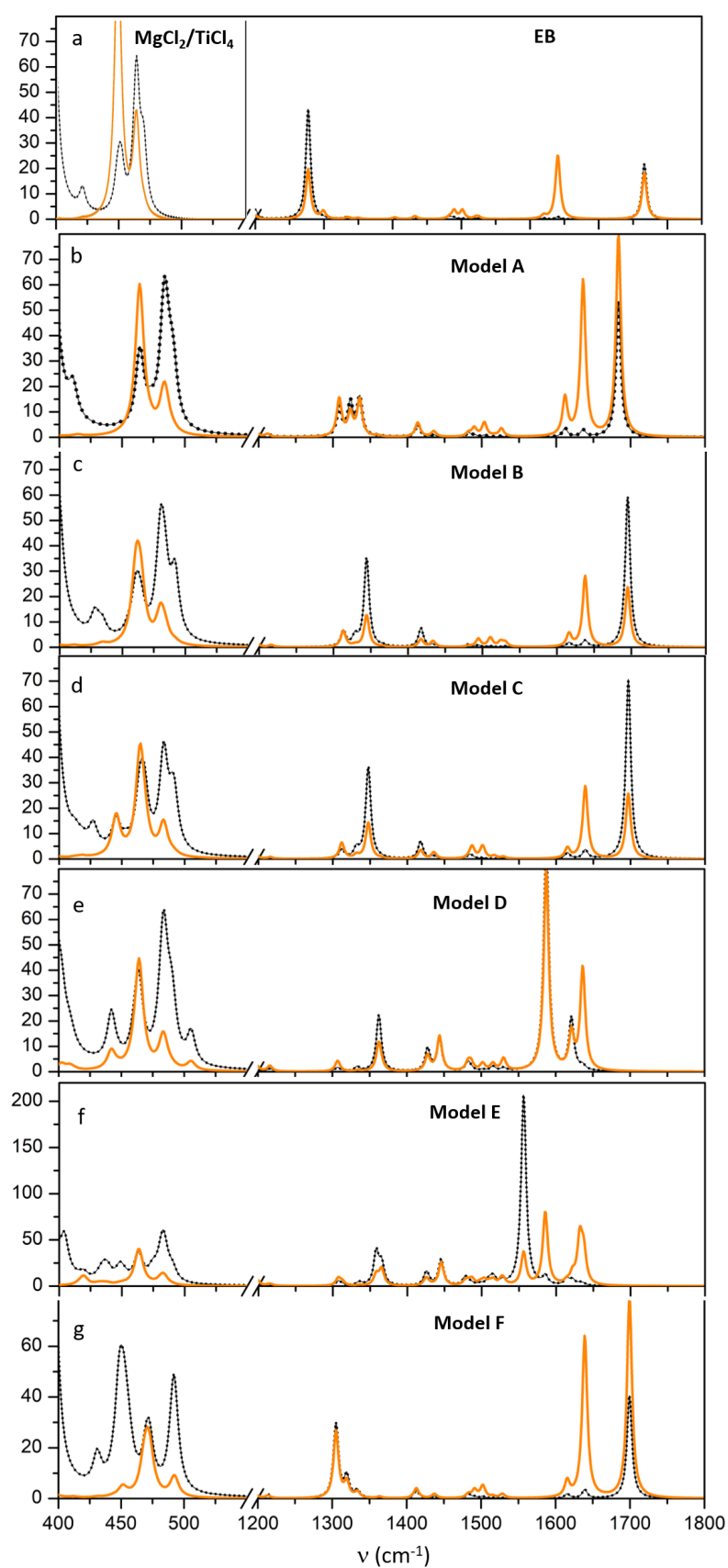


Figure 2. Simulated IR (black, dotted lines) and Raman spectra (orange, solid lines) at B3LYP-D2/TZVP for models proposed in Figure 1 in the spectral regions 400 - 550 cm^{-1} and 1200 - 1800 cm^{-1} . Intensity are in (Km/mol) and arbitrary units, respectively.

- the 400-500 cm^{-1} region. For systems A and B the Ti-Cl stretching mode at 465 and 485 cm^{-1} are only slightly perturbed by the presence of EB; for model C the coupling with EB mode causes the Ti-Cl symmetric stretching mode to split in components at 464, 465, 468 cm^{-1} and antisymmetric Ti-Cl stretching at 483, 484, also partially coupled with Mg-Cl modes of tetrahedral Mg^{2+} at 490 cm^{-1} . For models D-E two further bands, not visible in Raman, pop-up at 437 cm^{-1} and at 449 cm^{-1} due to bending modes of O-Ti-O and Ti-O-C, respectively. Weak bands at 588, 645 cm^{-1} assigned to Ti-O-C appear.

3.2.2. Raman characterization $\text{MgCl}_2/\text{TiCl}_4/\text{EB}$ adducts

Let us consider first the EB molecule by itself in gas phase, see Fig.2 panel a. The main differences with respect to the IR counterpart are: i) the much higher intensity of the 1640 and 1620 cm^{-1} bands, due to C-C stretching of the phenyl ring; ii) a band falling at 1019 cm^{-1} related to phenyl ring deformations; iii) a significant increase in intensity for the triplet 1483, 1489, 1501 cm^{-1} due to modes involving the O-C-O group. For EB supported adducts the main difference in the Raman spectra with respect to IR is the presence of the two intense bands at ~ 1636 and 1612 cm^{-1} related to the C-C stretching of the phenyl ring that appear almost unperturbed in all the models, and the antisymmetric and symmetric coupling of the two C=O stretchings of model E that, at variance of IR, give rise to peaks of comparable intensity at 1556 and 1585 cm^{-1} , thus unambiguously probing the presence of $\text{TiCl}_4(\text{EB})_2$. All the other peaks are common in the IR and Raman spectra even if with different relative intensity.

3.3. Adsorption properties of adducts on MgCl_2 nanoplatelets

Adsorption energies of selected models are collected in Table 1. Models A, B, C, L represents different coadsorption possibilities for TiCl_4 molecules and EB molecules. In the case A, one EB molecule is bound to isolated T site and close to a row of pentacoordinated Mg^{2+} sites hence reproducing a sort of edge condition. Models B and C similarly reproduces different EB on a P site: the first is perturbed by the presence of TiCl_4 whereas the latter is almost unaffected by TiCl_4 . In all three cases the adsorption energies are close each other (78-79 kJ/mol) with only a tiny larger energy in the adsorption to T site (A) but we can assume difference falls in the error bar. In model L the co-adsorption involve two EB molecules next to TiCl_4 , the energy involved in the adsorption of a second molecule on an adjacent site to TiCl_4 , as indicated in the reaction reported in Table I, is expected to be highly favored from an enthalpic point of view ($\Delta E = -121.9$ kJ/mol). In case F, the perturbation comes from the dimer Ti_2Cl_8 on sites of P type; the adsorption energy predicted for EB in presence of Ti species strengthens the binding of EB up to -99.3 KJ/mol, in fact in this case a chelation through O(R) occurs resembling a diester mode coordination. The other models D, G, E, H, I attain the adsorption of complexes TiCl_4EB_x ($x=1,2$), on defective T positions and only in the last case (I) on P position close to a couple of T Mg^{2+} sites. The mono-EB specie is weakly bound to T sites whereas the largest binding energies belong to the di-EB homogeneous-like adducts. A stronger interaction is predicted in the case G respect to D due to the presence of an adjacent most acidic T site. In the case G, the complex TiCl_4EB is bound to one between a couple of adjacent tetracoordinated Mg^{2+} sites in a 110-like manner; that differs from the outcome of previous periodic 2D calculations where the structure $\text{TiCl}_4\text{EB}/(110)$ evolves towards a new one with EB directly bound to support by carbonyl oxygen and at the meantime to a five coordinated TiCl_4 (Mg-O-Ti binding). This can be rationalized in terms of donor-donor interaction when periodic boundary conditions are assumed (even at low degree of coverage) and donor-adjacent layers of MgCl_2 . In the models presented here bulky donors have larger degrees of freedom to keep a Mg-Ti-O binding. TiCl_4 is partially detached from surface and that leaves coordination vacancies around titanium where a further EB molecule can easily insert to get a significant gain in energy because

Ti keeps a six-fold coordination due to two EB molecules bonded through carboxyl oxygen; that is the case of models E and H. In model I the adsorption energy of TiCl_4EB on P-type is close to case G, supporting the hypothesis that the presence of donor allows the binding of TiCl_4 on rows of 104-like Mg^{2+} sites contrary to an effective no binding of single TiCl_4 predicted in literature on 104 surfaces. For all the above-mentioned cases the adsorption energy has been computed by means of B3LYP-D2 method and in some selected cases, single points calculations have been carried out by employing the M06 functional as shown in Table 1.

In models close to truly nanosized structures, the clean break between the adsorption behaviour of T and P sites, emerging from a periodic approach, seems to become smoother; that can be due to the proximity of a few sites of the first type to the ones of the latter with consequent polarizing effect coming from the T sites on closer P ones. The adsorption energy reveals that the binding of EB is only tinely sensitive to the site of adsorption due to the frontier behaviour of sites. In particular, in the case of model B, where the proximity of TiCl_4 increases the polarizing ability of pentacoordinated Mg^{2+} (as we already found in probing cluster sites by CO [57]), the adsorption energy almost equal in model A and model B. In the case of model C the adsorption energy is only weakly affected due to the finite nature of support model and the fact that we are simply considering a monoester (for simplicity) with no strong ability to chelate the tetracoordinated sites present in type A models. In the case of homogeneous-like complexes, the complex with two EB molecules (model E) is by far more strongly bound to cluster, in fact in mono-EB case (models D, G), a large distortion occurs with respect to the geometry adopted from molecule of EB in the gas phase with a cost amounting to 34.5 kJ/mol. For model D, the significant difference with the results obtained for extended surfaces [61] is due to the fact that the minimum is unstable; it evolves towards a structure where EB is bound to an adjacent tetracoordinated site, in other words the TiCl_4EB is bonded to MgCl_2 through the oxygen of the ester group.

Table 1. Energetics of adsorption of EB, TiCl_4EB and TiCl_4EB_2 complexes for models reported in Fig.1 (energies in kJ/mol). All calculations were performed by adoption of B3LYP-D2 and M06 functionals and Ahlrichs VTZP basis set. In M06 case, single point energy calculations have been computed on geometries optimized at B3LYP-D2 level. All data are BSSE corrected.

Reaction	Model	$\Delta E(\text{B3LYP-D2})$	$\Delta E(\text{M06})$
$\text{EB}_{\text{gas}} + (\text{MgCl}_2\text{TiCl}_4)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\cdots\text{EB})_{\text{surf}}$	A	-79.3	-81.6
"	B	-78.9	-80.9
"	C	-77.5	-79.5
$\text{TiCl}_4\text{EB}_{\text{gas}} + (\text{MgCl}_2\text{TiCl}_4)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB})_{\text{surf}}$	D	-66.3	-75.4
$\text{TiCl}_4\text{EB}_2\text{gas} + (\text{MgCl}_2\text{TiCl}_4)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB}_2)_{\text{surf}}$	E	-181.2	-185.8
$\text{EB}_{\text{gas}} + (\text{MgCl}_2\text{Ti}_2\text{Cl}_8)_{\text{surf}} \rightarrow ((\text{MgCl}_2\text{Ti}_2\text{Cl}_8\cdots\text{EB})_{\text{surf}}$	F	-99.3	-96.0
$\text{TiCl}_4\text{EB}_{\text{gas}} + (\text{MgCl}_2\text{TiCl}_4)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB})_{\text{surf}}$	G	-79.3	-94.9
$\text{TiCl}_4\text{EB}_2\text{gas} + (\text{MgCl}_2\text{TiCl}_4)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB}_2)_{\text{surf}}$	H	-172.1	-191.6
$\text{TiCl}_4\text{EB}_{\text{gas}} + (\text{MgCl}_2\text{Ti}_2\text{Cl}_8)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB})_{\text{surf}}$	I	-79.5	-95.3
$\text{EB}_{\text{gas}} + (\text{MgCl}_2\text{TiCl}_4\cdots\text{EB})_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\cdots\text{EB}_2)_{\text{surf}}$	L	-121.9	-123.6
$(\text{MgCl}_2\text{TiCl}_4\cdots\text{EB})_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB})_{\text{surf}}$	G	35.5	60.9
$(\text{MgCl}_2\text{TiCl}_4\cdots\text{EB}_2)_{\text{surf}} \rightarrow (\text{MgCl}_2\text{TiCl}_4\text{EB}_2)_{\text{surf}}$	H	55.5	67.8

3.4. Discussion and Conclusions

EB is adsorbed at T and P isolated sites (models A and C) and at T and P sites adjacent to preadsorbed TiCl_4 and Ti_2Cl_8 (models B and F) giving rise to surface adducts of comparable stability (~ 80 kJ/mol, B3LYP-D2 and M06 results, Table 1). Physisorbed

EB on TiCl_4 and Ti_2Cl_8 cannot be identified since EB on Ti_2Cl_8 causes the fragmentation of the Ti cluster into TiCl_4/EB (model G) and physisorbed TiCl_4 . The formation of the MgCl_2/EB adducts is monitored by the IR spectra.

Line a of Fig. 3 reports a convolution of IR spectra of the most relevant MgCl_2/EB surface structures in presence of TiCl_4 (sum of simulated intensities for models A, B, C and F). Inspection of the spectrum shows that the two main features are the $\nu(\text{C}=\text{O})$ peak, sharp and narrow (its width is $\sim 12\text{ cm}^{-1}$), centered at 1678 cm^{-1} and the $\nu(\text{C}-\text{O}-\text{C})$ vibration that leads to a group of signals in the $1275\text{--}1350\text{ cm}^{-1}$ range; $\nu(\text{C}=\text{O})$ peak appears to be sensitive to the coordination number of Mg^{+2} , but scarcely affected by the chemical environment of the surface cations, and similarly the characteristic shape of the $\nu(\text{C}-\text{O})$ signal can unambiguously probe a simultaneous presence of tetra and pentacoordinated Mg^{+2} sites (compare panel d to panel f of Figure 2).

The homogeneous like TiCl_4EB and TiCl_4EB_2 complexes can easily bind at the the Mg^{+2} sites of MgCl_2 . However the only species present on P sites of the 104-like row is the TiCl_4EB (model G); on T sites (i.e. 110-like sites) both TiCl_4EB and TiCl_4EB_2 can be coordinated. The relative binding energies are quite similar for TiCl_4EB adducts, $\Delta E \sim 70\text{--}80\text{ kJ/mol}$ (B3LYP-D2), but much larger for TiCl_4EB_2 , $\Delta E \sim 170\text{--}180\text{ kJ/mol}$ (B3LYP-D2). Convoluted IR spectrum of adsorbed TiCl_4EB_x complexes (sum on spectra of models D and E) reported as line b of Figure 3, show that the TiCl_4EB complex is associated with two signals centered at about 1620 cm^{-1} due to ring deformations coupled to $\text{C}=\text{O}$ stretching whereas $\text{TiCl}_4/(\text{EB})_2$ is unambiguously characterized by a band at 1556 cm^{-1} , due to carbonyl couples, that dominates the IR spectrum.

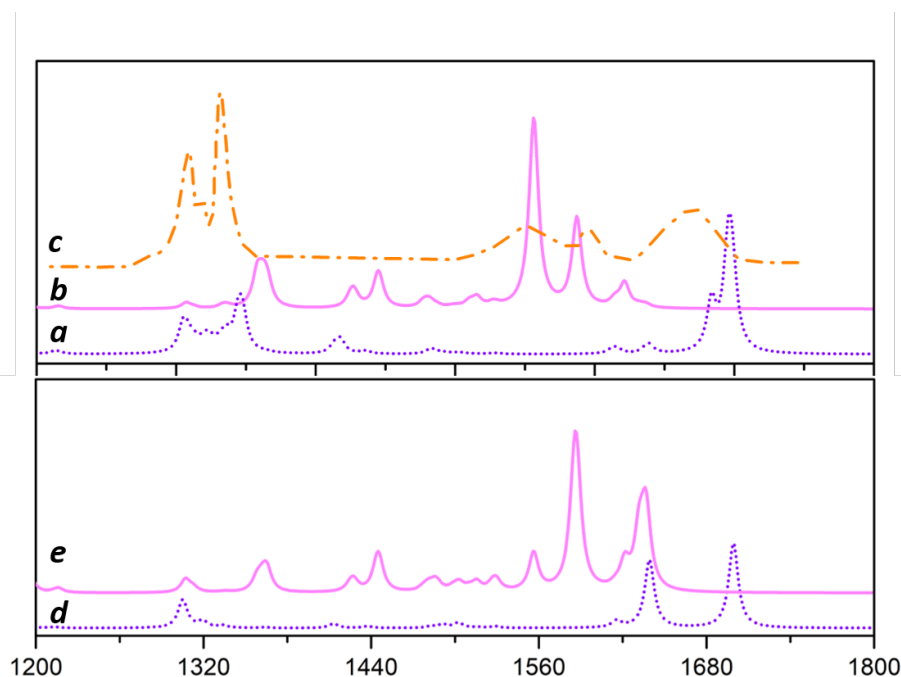


Figure 3. Sum of simulated IR intensity at B3LYP-D2/TZVP, for co-adsorption models A, B, C, F (line a) and for homogeneous-like adducts, models D, E, (line b). Experimental IR spectrum from Ref. [61] reported in the $1200\text{--}1350$ and $1550\text{--}1800\text{ cm}^{-1}$ spectral range (line c). Sum of simulated Raman intensity at B3LYP-D2/TZVP, for co-adsorption models A, B, C, F (line d) and for homogeneous-like adducts, models D, E, (line e). The spectral region considered is $1200\text{--}1800\text{ cm}^{-1}$, the intensities are in arbitrary units.

We can notice the excellent agreement between computed IR spectra of Figure 3a-b and the experimental one of $\text{MgCl}_2/\text{EB}/\text{TiCl}_4$ from Ref. [61] and reported in Figure 3b; the larger discrepancy can be found in the region of $\nu(\text{C}=\text{O})$ (CO stretching) with a shift

that can be easily attributed to anharmonicity of the mode, neglected in the harmonic approximation we adopted. Concerning the Raman response, convoluted spectra are reported in Figure 3d-e. The high frequency region is dominated by two intense peaks at 1720 cm^{-1} due to $\nu(\text{C}=\text{O})$ and at 1660 cm^{-1} associated to C-C stretching mode of the phenyl ring; the $1275\text{--}1350\text{ cm}^{-1}$ range shows a group of signals related to $\nu(\text{C}-\text{O})$. Adsorbed $\text{TiCl}_4\text{EB}_{1,2}$ provide clear fingerprints in the Raman spectra: the couple of bands at 1660 (with a shoulder at 1642 cm^{-1}) and at 1606 cm^{-1} identifies the mono-EB adduct, whereas the presence of an additional peak at 1576 cm^{-1} reveals the presence of the di-EB adduct.

The formation of homogeneous like $\text{TiCl}_4\text{EB}_{1,2}$ at the MgCl_2 nanoplatelets is thus unambiguously probed by IR and Raman spectra. However we can wonder about the role, if any, of the MgCl_2 substrate in promoting the formation of those surface complexes. Starting from MgCl_2/EB , TiCl_4 can be bound on a couple of adjacent T sites even in the presence of preadsorbed EB at one of the T centers ($\text{MgCl}_2/\text{TiCl}_4\cdots\text{EB}$, model B of Figure 1 and Table 1) with a binding energy of -91.6 kJ/mol , (the corresponding value on bare MgCl_2 is 69.0 kJ/mol). In the same way, starting from $\text{MgCl}_2/\text{TiCl}_4$, EB can absorb at the one of the T sites that binds TiCl_4 (again model B) with a coordination energy of -106.9 kJ/mol (M06). Despite sharing the same Mg^{+2} center, both TiCl_4 and EB are strongly bound to the substrate; thus a surface reaction starting from $\text{MgCl}_2/\text{TiCl}_4\cdots\text{EB}$ (Model B) and leading to $\text{MgCl}_2/\text{TiCl}_4(\text{EB})$ (model E) is energetically not favored by $\sim 61\text{ kJ/mol}$, Table 1. We consider now a couple of EBs adsorbed on adjacent T sites; coordination of TiCl_4 involving the same T sites is again energetically favored by -103 kJ/mol : hence, coadsorption, first with one and then with two EBs progressively improves TiCl_4 coordination to MgCl_2 . Similarly a second EB molecule successfully coadsorbs on $\text{MgCl}_2/\text{TiCl}_4\cdots\text{EB}$ leading to Model L with an associated energy variation $\Delta E = -123.6\text{ kJ/mol}$. Hence, EB and TiCl_4 act cooperatively and stabilize each other so that, the presence of a second coadsorbed EB further disfavors a spontaneous evolution to the homogeneous like $\text{TiCl}_4(\text{EB})_2$ surface adduct (Model H) and the corresponding reaction is disadvantaged by 68 kJ/mol , Table 1. Indeed, it is experimentally known that TiCl_4/EB complexes dissociatively adsorb on MgCl_2 . [82] The presence of donor allows the binding of TiCl_4 on rows of 104-like Mg^{2+} sites (model I) contrary to an effective no binding of single TiCl_4 predicted in literature on 104 surfaces.

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