Properties of \textit{Ir}_4 Clusters in the Gas Phase and on Oxide Substrates

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Résumé

Les résultats d’une étude théorique des propriétés de cluster d’Ir$_4$ dans la phase gazeuse et sur des surfaces d’oxyde sont présentés. Ce travail est basé sur la méthode de la théorie de la fonctionnelle de la densité (DFT) dans l’approximation du gradient généralisé (GGA) et des pseudo-potentiels ”ultrasoft”.

Les propriétés d’une petite particule telle que le cluster d’Ir$_4$ sont entièrement déterminés par sa géométrie. Le fait déjà connu que la forme la plus stable de l’Ir$_4$ dans la phase gazeuse est la structure carrée, qui est significativement plus stable que les structures en ”papillon” (butterfly) ou en tétraèdre, est confirmé. Ce résultat est en contradiction avec des expériences qui indiquent que l’Ir$_4$ déposé sur l’oxyde adopte une configuration en tétraèdre.

Il est montré, dans cette thèse, que l’environnement chimique exerce une forte influence sur la stabilité relative des clusters d’Ir$_4$.

Sur la surfaces MgO(110) l’isomère carré demeure la structure d’Ir$_4$ la plus stable, bien séparée en terme d’énergie des deux autres. De plus, le tétraèdre est fortement déformé de part l’interaction avec l’oxygène de la surface. La présence de défauts ponctuels (neutres et lacunes chargées O) affecte l’ordre énergétique, rendant tétraèdre et carré très proche en énergie, les déformations structurelles du tétraèdre étant même plus importantes, prédictions en désaccord avec les expériences.

Sur la surface TiO$_2$(110) les structures en tétraèdre et carré deviennent dégénérées et celle en papillon est l’isomère le moins stable. De plus, les déformations structurelles sont très petites, en bon accord avec les expériences. Il est montré que la surface de TiO$_2$ influence la stabilité relative des trois isomères par un champ électrostatique particulièrement fort.

Les interactions de l’Ir$_4$ avec les atomes H, C and O ainsi qu’avec des molécules de CO ont été étudiées. L’adsorption d’un unique atome C influence fortement la stabilité relative des trois isomères. Avec l’adsorption de C, la structure papillon devient le plus stable isomère de la phase gazeuse alors que sur les deux surfaces l’Ir$_4$ tétraèdre est la structure la plus probable. L’adsorption d’un seul atome C ou O ne produit pas le même effet.

L’interaction avec les molécules de CO est aussi important étant donné la procédure expérimentale pour produire les cluster d’Ir$_4$ déposés. Il est montré que sur la MgO(100) la dissociation du CO est autant probable que le processus concurrent de désorption du CO, justifiant la présence d’atomes de carbone sur les clusters d’Ir$_4$ et rapprochant ainsi les prédictions théoriques des données expérimentales.

Mots-clés: clusters, iridium, théorie de la fonctionnelle de la densité, surfaces, adsorption
Abstract

Results of a theoretical study on the properties of Ir\textsubscript{4} clusters in the gas–phase and on oxide surfaces are presented. The work is based on density functional theory (DFT) within the generalized gradient approximation (GGA) and ultrasoft pseudopotentials.

Properties of a small particle such as Ir\textsubscript{4} cluster are entirely determined by its geometry. The already known result that the most stable form of Ir\textsubscript{4} in the gas–phase is the square structure which is significantly more stable than the butterfly and tetrahedron is confirmed. This result is in contradiction with experiments which indicate that the oxide supported Ir\textsubscript{4} adopts a tetrahedral configuration. It is shown in this thesis that the chemical environment has a strong influence on the relative stability of Ir\textsubscript{4} clusters.

On MgO(100) surface, the square isomer remains the most stable Ir\textsubscript{4} structure, well separated in energy from the other two. Moreover, the tetrahedron is heavily distorted by the interaction with the surface oxygen. Presence of point defects (neutral and charged O vacancies) affects the energy ordering making tetrahedron and square very close in energy, but the structural distortion of the tetrahedron is even bigger and the predicted data do not correspond to experiments.

On TiO\textsubscript{2}(110) the tetrahedron and square structures become degenerate and the butterfly becomes the least stable isomer. Moreover, structural distortions are very small, in agreement with experimental data. It is shown that the TiO\textsubscript{2} surface influences the relative stability of the three isomers through a particularly strong electrostatic field.

Interactions of Ir\textsubscript{4} with H, C and O atoms as well as with CO molecules have been studied. Adsorption of a single C atom strongly influences the relative stability of the three isomers. Upon C adsorption, the butterfly becomes the most stable gas–phase isomer while on both surfaces the tetrahedron is the most probable structure. Adsorption of a single H or O atom does not produce the same effect.

The interaction with CO molecules is also important given the experimental procedure used for producing supported Ir\textsubscript{4} clusters. It is shown that on MgO(100), CO dissociation is as probable as the competing process CO desorption justifying the presence of carbon adatoms on Ir\textsubscript{4} clusters which brings theoretical predictions in better agreement with experimental data.

**Keywords:** clusters, iridium, density functional theory, surfaces, adsorption
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Chapter 1

Introduction

In atomic and molecular physics and in chemistry the term "cluster" refers to a group of bound atoms ranging from a few atoms up to several hundred or even thousand atoms. The term was introduced in early 1960s to describe compounds containing metal–metal bonds [1]. The definition has evolved and nowadays "cluster" refers to any group of atoms that contains element–element bonds.

Clusters can be both mononuclear or polynuclear. They are intermediate between atomic and bulk systems and as such, they are the true representatives of the "nanoworld", reduced both in size and dimensionality. Quantum confinement effects together with the lack of symmetry constraints make them extremely interesting since their properties may be completely different from the properties of their atomic and bulk counterparts (i.e. enhanced catalytic activity, magnetism in otherwise nonmagnetic systems,...). It has been found that the properties of clusters strongly depend on their size [2]. They may display abrupt changes with the cluster size, especially for small species (up to approximately several tens of atoms) while for bigger clusters the properties typically converge to those of the bulk.

These effects make clusters promising candidates for several technological applications such as catalysis, sensors, magnetic storage devices, solar cells, controlled drug delivery, etc. Potentially, clusters could also be used as the building blocks of composite materials whose properties can be tuned through selection of their size and composition [3].

As catalysts clusters are used both in solutions (homogeneous catalysis) or dispersed over an "inert" oxide support (for heterogeneous catalysis). The enhanced catalytic activity of clusters is usually associated with the large fraction of undercoordinated atoms due to their high surface to volume ratio [4]. For applications in catalysis an important family of elements used for cluster fabrication is comprised of the group 8–10 transition metals or the "platinum group" (Ru, Rh, Pd, Os, Ir and Pt).

Iridium is one of the "platinum group" transition metals with atomic number $Z = 77$, laying below rhodium and in–between osmium and platinum in the periodic table. It is very hard and brittle, and due to its very high melting temperature (2739K [5]) it is very hard to machine, form, or work. Iridium is the most corrosion resistant metal known, and was used in making the standard metre bar of Paris, which is a 90% platinum–10% iridium alloy\(^1\). However, its powder, if fine enough, is much more reactive and can even be flammable. Iridium is one of the least abundant elements in the Earth’s crust, but on the other hand, it is one of the cheepest precious metals, third after silver and palladium and three to four times cheaper than platinum.

\(^1\)This metre bar has since been replaced as a fundamental unit of length.
times less expensive than platinum and rhodium [6].

Iridium, as other group 8–10 transition metals, is also used as a catalyst. The surfaces of iridium have been shown to be a very effective catalysts for the C–H bond activation in alkanes and alkenes. Mechanisms of the dissociative chemisorption of both methane [7] and ethene [8] on iridium surfaces have been studied experimentally. Several reactions of industrial importance such as conversion of saturated hydrocarbons to aromatic compounds, steam reforming of methane and the conversion of methane to higher hydrocarbons or methanol involve the C–H bond activation. Furthermore, iridium complexes, mostly in the form of carbonyls, are also widely employed as catalysts in the chemical industry. The carbonyls are extensively used in homogeneous catalysis as promoters in the Cativa process for carbonylation of methanol to produce acetic acid [9]. Almost 60% of the production of acetic acid is done through the carbonylation of methanol and the Ir based catalysts offer significant improvements over conventional rhodium technology.

However, the need for more efficient heterogeneous catalysts, stimulated interest in studying small iridium clusters deposited on oxide surfaces. Several techniques for preparation of the supported species, such as size and energy–selected deposition of Ir\textsuperscript{m+n} (n = 1, 2, 5, 10, 15) clusters [10] or the vapor–deposition of Ir nanoparticles [11] have been used. Moreover, a method for producing nearly monodispersed Ir\textsubscript{4} particles over common oxide substrates, such as MgO, Al\textsubscript{2}O\textsubscript{3}, or TiO\textsubscript{2} was developed in the last 15 years [12]. The production procedure of supported Ir\textsubscript{4} clusters involves Ir\textsubscript{4}(CO)\textsubscript{12} carbonyls that are used as a source of iridium. They are deposited on an oxide surface, and decarbonylated in the flowing He at elevated temperatures. A combination of infrared (IR) and extended X-ray absorption fine structure (EXAFS) spectroscopies is then used for the characterization of the as–prepared species and for the study of their catalytic properties. A tenfold increase in the catalytic activity has been observed by replacing the MgO substrate with a \(\gamma\)–Al\textsubscript{2}O\textsubscript{3} revealing a strong influence of the substrate on the properties of the adsorbed species. The EXAFS results also showed that upon full decarbonylation the Ir–Ir coordination number remains three as in the precursor, suggesting that the carbonyl tetrahedral Ir\textsubscript{4} frame remains intact. This is an important result since the properties of such a small system like Ir\textsubscript{4} depend strongly also on the atomic configuration\textsuperscript{2}.

On the other hand, only a few theoretical studies on Ir clusters exist in the literature [64, 66, 71, 73] mostly treating isolated (gas-phase) clusters\textsuperscript{3}. In most of these studies [64, 71, 73] the results are obtained using state-of-the-art numerical implementations of density functional theory (DFT) which proved itself as a reliable theoretical tool for studying transition metal clusters both freestanding and on an oxide substrate [4, 13]. All authors agree that the most stable configuration of the gas–phase Ir\textsubscript{4} is the square (S) geometry. This configuration is significantly more stable than the tetrahedron (T) and the butterfly (B) structures. This result is in contradiction with the experimental findings for the geometry of supported Ir\textsubscript{4} clusters.

Results of an extensive theoretical study on the properties of different isomers of the gas–phase as well as oxide supported Ir\textsubscript{4} clusters are presented in this thesis. The influence of the chemical environment on the properties of the three most relevant Ir\textsubscript{4} isomers (T, B and S) is investigated. Special accent is put on the relative stability of the three isomers motivated by the already mentioned discrepancy between the theoretical predictions for the geometry\textsuperscript{2}

\textsuperscript{2}As it will be shown in this thesis.

\textsuperscript{3}Excluding the publications presenting the results obtained during this work [69, 70]
of the gas–phase Ir$_4$ and experimental findings for the supported species. As substrates the rocksalt MgO(100) and rutile TiO$_2$(110) oxide surfaces are chosen. Both of them are widely used in experiments and exhibit important differences in the structure and reactivity. Interaction of the three isomers, both free and supported, with H, C and O atoms and CO molecules is also studied. This choice of reactants is considered as the most relevant given the experimental procedure used to prepare Ir$_4$. Namely, hydrogen could come from the surface hydroxyl groups, while C and O atoms could be present as fragments left on the surface after decarbonylation of the Ir$_4$(CO)$_{12}$ precursor. In the case of partial decarbonylation also the CO molecules would be attached to the Ir$_4$ cluster.

In this work it is shown that chemical environment may cause drastic changes in both relative stability and reactivity of Ir$_4$ clusters [69, 70]. It is confirmed here that the S isomer is the most stable form of Ir$_4$ in the gas–phase. Moreover, deposition on MgO(100) surface does not alter the relative stability predicted for the gas-phase species. However, one adsorbed carbon atom is sufficient to do so in both environments [69]. While in the gas–phase the most stable structure upon C adsorption becomes B, on MgO(100) surface C adsorption changes the relative stability in favor of the T structure as observed in experiments [118, 119]. At variance to the adsorption of a single carbon atom, it is found that one H or O adatom does not modify the Ir$_4$ isomer stability observed for the pristine clusters [70]. Qualitatively, the same picture is accounted when H and O atoms are chemisorbed on Ir$_4$ clusters supported by MgO(100). The TiO$_2$ substrate also affects the relative stability of the three isomers. Upon adsorption of the pristine clusters T structure becomes almost degenerate with S, bringing theoretical predictions closer to the experimental results [12, 126]. Moreover, the reactivities of the clusters towards C adsorption as well as CO adsorption and dissociation are also strongly affected.

The thesis is organized as follows. In the first part the methods used in this study are briefly described. Foundations of density functional theory are presented and the main approximations are overviewed (Chap.2). Afterwards, properties of the gas–phase Ir$_n$ ($n = 1, \ldots, 8$) clusters are presented (Chap.3). Relative stability between different isomers given for a fixed cluster size is discussed as well as the differences in their properties. Special attention is on the Ir$_4$ isomers. Bonding in Ir$_4$ is analyzed together with the differences in the electronic structures of the three isomers that lead to a given energy ordering. The relative stability of the three Ir$_4$ isomers is than compared with other transition metal tetramers. The details of the interaction of the gas–phase Ir$_4$ clusters with H, C and O atoms as well as with a single CO molecule are presented in the following chapter (Chap.4). The pathways for the CO dissociation are also studied.

The thesis also covers properties of supported Ir$_4$ clusters (Chapter 5). Interaction mechanisms with two oxide surfaces MgO(100) and TiO$_2$(110) are discussed as well as their influence on the relative stability of Ir$_4$. Moreover, diffusion of Ir$_4$ on the MgO(100) surface is studied together with their interaction with neutral and charged oxygen vacancies. The influence of adsorbates on the relative stability of supported Ir$_4$ is the subject of the last chapter and it is followed by the conclusions and the list of references.
Chapter 2

Density Functional Theory and Methods

2.1 Density Functional Theory

Probably the most important and most used methods of all \textit{ab initio} electronic structure methods are based on Density Functional Theory (DFT). DFT is the theory of an interacting electron gas whose foundations lay in one theorem, the Hohenberg and Kohn (HK) theorem \cite{17}. This theorem states that "the ground state density of bound system of interacting electrons in some external potential determines this potential uniquely" \cite{18}, where uniquely means up to an additive constant.

The HK theorem has two important implications: (i) by determining the external potential ground state density determines the system completely together with all its properties; and (ii) there is a universal functional for the energy $E[n]$ in terms of the density $n(r)$ whose global minimum, for a given external potential, is the exact ground state density $n_0(r)$ \cite{19}. However, the theorem does not provide a way to construct the energy functional.

Kohn and Sham showed \cite{20} that it is possible to replace the problem of interacting many–electron system by an auxiliary system of non–interacting fermions. Therefore, minimization of the (unknown) energy functional with respect to the electron density can be recast into a set of self–consistent one–electron equations. The Kohn–Sham (KS) energy functional can be written as

$$F[n(r)] = T_s[n(r)] + \int V_{ext}(r)n(r)dr + E_H[n(r)] + E_{xc}[n(r)],$$

(2.1)

where $T_s[n]$ is the kinetic energy functional for the non–interacting many–electron system, $V_{ext}(r)$ is the external potential, $E_H[n]$ is the Hartree energy and $E_{xc}[n]$ is the exchange–corelation (XC) functional which is defined by Eq.2.1. The following set of one–particle, KS equations can then be derived by minimizing $F[n(r)]$ under the orthonormality condition for the wave functions

$$\left[-\frac{1}{2} \nabla^2 + V_{ext}(r) + \int \frac{n(r')}{|r - r'|}dr' + V_{xc}(n(r)) \right] \psi_i(r) = \epsilon_i^{KS} \psi_i(r).$$

(2.2)

The quantity in brackets is the Kohn–Sham Hamiltonian consisting of the kinetic energy operator, external potential followed by the Hartree and XC potentials. The XC potential is derived as the functional derivative $\delta E_{xc}/\delta n$ of the XC energy functional with respect to
the density. The eigenvalues $\epsilon_i^{KS}$ enter into the formalism as the Lagrange multipliers for the orthonormality condition.

The XC energy functional is not known and needs to be approximated. Once, an appropriate approximation for $E_{xc}$ is made, the KS equations are solved self-consistently starting from a certain $n(r)$ and constructing all terms in Eq.2.2. After the equations 2.2 are solved, the KS orbitals are filled according to the Pauli principle and the new density is constructed. For the set of doubly occupied KS orbitals $\{\psi_i\}$ (spin is neglected for simplicity) the total energy for the system of interacting electrons in the field of fixed nuclei can be calculated (in Hartree atomic units) as follows

$$E[\{\psi_i\}] = 2 \sum_i^{\text{occ}} \int \psi_i^*(-\frac{1}{2} \nabla^2)\psi_i d\mathbf{r} + \int V_{\text{ion}}(\mathbf{r})n(\mathbf{r})d\mathbf{r} + \frac{1}{2} \int \frac{n(\mathbf{r})n(\mathbf{r'})}{|\mathbf{r} - \mathbf{r'}|} d\mathbf{r}d\mathbf{r'} + E_{XC}[n(\mathbf{r})] + E_{\text{ion}}[\{R_I\}], \quad (2.3)$$

where all the terms are already known from Eq.2.1 except $E_{\text{ion}}[\{R_I\}]$ which represents the electrostatic interaction between nuclei (ions).

### 2.1.1 The LDA and GGA Approximations and Beyond

The simplest and the most used approximation to DFT for $E_{xc}[n(\mathbf{r})]$ is the Local Density Approximation (LDA). The XC energy functional is written within the LDA in the form

$$E_{LDA}^{xc} = \int \epsilon_{xc}^{\text{hom}}(n(\mathbf{r})) n(\mathbf{r}) d\mathbf{r} \Rightarrow V_{xc} = \frac{\delta E_{xc}}{\delta n} = \frac{d}{dn} \left[ n\epsilon_{xc}^{\text{hom}}(n) \right], \quad (2.4)$$

where $\epsilon_{xc}^{\text{hom}}(n(\mathbf{r}))$ is the exchange–correlation energy per particle of a uniform electron gas of density $n$. This approximation, expected to produce accurate results only for systems with slowly varying densities, works remarkably well also for inhomogeneous systems (i.e. isolated atoms). The usefulness of the LDA is not only in its accuracy, but also in its simplicity. Namely, the KS equations in the LDA approximation are only slightly more complicated than Hartree equations\(^1\) and are much more accurate. They are also much simpler than Hartree–Fock equations [67]. The success of LDA for the interacting electron systems (even inhomogeneous) can be, to a large extent, attributed to the fact that LDA satisfies the sum–rule for the exchange–correlation hole\(^2\) [18, 19].

Early attempts to improve LDA by including the terms dependent on the density gradients in Eq.2.4 have not shown improvements compared to simple LDA. One of the reasons for the failure of these "improvements" is the fact that simple gradient expansions do not obey the sum–rule for the exchange–correlation hole. Considerable efforts to construct the correct functionals have lead to Generalized Gradient Approximation (GGA). The GGA functionals can be written in the (general) form

$$E_{GGA}^{xc} = \int f(n(\mathbf{r}), |\nabla n(\mathbf{r})|) n(\mathbf{r}) d\mathbf{r}. \quad (2.5)$$

Many forms for $f(n, |\nabla n|)$ have been proposed up to now. Probably, the most used GGA functionals are those constructed by Perdew and Wang (PW91) [21], Perdew, Burke and

\(^1\)Which can be derived from the KS equations 2.2 by putting $V_{xc} = 0$.

\(^2\)Depletion of the two particle density $n(\mathbf{r}, \mathbf{r'})$ around $\mathbf{r}$, with respect to the average density $n(\mathbf{r'})$, which integrates to 1.
Enzerhof (PBE) [22], and Becke, Lee, Yang and Parr (BLYP) [23, 24]. Use of GGA instead of LDA has reduced errors of atomization energies of molecules, energy barriers and structural energy differences (for details check reference [22] and the references therein). Improved accuracy as well as the development of numerical algorithms which enabled investigations of the properties of relatively large systems have made GGA an important component of today’s theoretical solid–state physics and quantum chemistry.

However, both LDA and GGA functionals contain a fraction of the self interaction error that is comprised in the Hartree term and cancelled completely only by the “exact”, Fock exchange operator. Therefore, the so called hybrid XC functionals have been developed which include a portion of the Fock exchange added to the standard LDA/GGA functionals. They are typically constructed in the following way

$$E_{\text{hybrid}}^{\text{xc}} = E_{c}^{\text{GGA}} + (1 - \alpha)E_{x}^{\text{GGA}} + \alpha E_{x}^{\text{Fock}},$$

(2.6)

where parameter $\alpha$ determines what portion of the exchange will be ”exact”. If the hybridization is done with the PBE (GGA) functional than it has been justified that for molecular systems it is appropriate to put $\alpha = 0.25$ leading to the widely used PBE0 hybrid functional [25]. Although hybrid functionals do improve DFT results when applied to semiconductors/insulators or small molecules consisting of light atoms, their description of the electronic structure of metallic systems is rather poor (wrong bulk cohesive energies, gap opening, unphysical broadening of the transition metal $d$ bands, . . . ) [26]. This is not surprising since it is known that the self–interaction error is almost negligible in metals. Moreover, it is also shown that, when applied to transition metal dimers, hybrid functionals do not yield results in better agreement with experiments than those obtained with GGA methods [59, 60, 61]. Therefore, the most appropriate approximation to DFT for studying transition metal clusters and their interactions with surfaces and/or other molecules is still GGA which was used in the PW91 functional form throughout this work.

2.2 Methods

2.2.1 Pseudopotentials

Widely used approximation, where the Coulomb potential of nuclei is replaced by the pseudopotential, is based on the fact that the inner shell (core) electrons, in majority of cases, do not play a role in chemical bonding and are weakly affected by the environment surrounding the atom. It is a useful simplification to treat those electrons as ”frozen” and incorporate their contribution in the potential that acts on valence electrons. The gain when using this approximation is not only in reducing the number of electrons that are treated explicitly, but also in reducing the number of plane–waves that are needed for accurate representation of the valence wavefunctions. A huge number of plane waves is needed mainly due to the orthogonality constraints of the valence wave functions to the core orbitals. Price for using pseudopotential approximation is their nonlocality and the arbitrariness in dividing electrons into core and valence.

When constructing pseudopotentials one starts from the all–electron calculation for a given atom solving the KS equation (Eq.2.2). Approximation of spherical screening is usually

$^3$Dependence on the XC functionals used in simulations was verified by comparing GGA results with those obtained within LDA.
assumed leading to the radial equation

\[
\left[-\frac{1}{2}\frac{d^2}{dr^2} + \frac{l(l+1)}{2r^2} + V_{\text{eff}}[\rho; r]\right] r R_{nl}(r) = \varepsilon_{nl} r R_{nl}(r),
\]  

(2.7)

where \(V_{\text{eff}}[\rho; r]\) is the self-consistent one-electron potential (for example in the LDA approximation)

\[
V_{\text{eff}}[\rho; r] = -\frac{Z}{r} + V_{\text{H}}[\rho; r] + V_{\text{xc}}^{\text{LDA}}[\rho].
\]  

(2.8)

A following set of requirements is then imposed on the properties of pseudopotentials:

1. all–electron and pseudo–valence eigenvalues agree for chosen atomic configuration,
2. all–electron and pseudo–valence wavefunctions agree beyond a chosen core radius \(R_{cl}\),
3. the logarithmic derivatives of the all–electron and pseudo–wavefunctions agree at \(R_{cl}\),
4. the integrated charge inside \(R_{cl}\) for each wavefunction agrees (norm–conservation),
5. the first energy derivative of the logarithmic derivatives of the all–electron and pseudo–wavefunctions agrees at \(R_{cl}\) (transferability).

Pseudopotentials that fulfill these conditions are referred as norm–conserving. The above set of conditions does not determine them uniquely and in practice, firstly the pseudo–wavefunctions with the desired properties are constructed and then the screened pseudopotentials, containing the influence of other valence electrons, are obtained by inverting Eq.(2.7). Since the screening properties of valence electrons depend strongly on the environment it is necessary to “unscreen” the pseudopotentials to ensure their transferability. One way to do this is to divide the electronic density into contributions of core and valence electrons \(\rho = \rho_c + \rho_v\) and to subtract from the screened pseudopotential contribution to the electron–electron interaction coming from \(\rho_v\)

\[
V_{\text{ion},l}^{PP}(r) = V_{\text{scr},l}^{PP}(r) - V_{\text{H}}[\rho_c; r] - V_{\text{xc}}[\rho_v].
\]  

(2.9)

This is also an approximation since the \(V_{\text{xc}}[\rho]\) is not linear in \(\rho\). It is a fairly good approximation for systems whose \(\rho_c\) and \(\rho_v\) are spatially well separated, but for systems whose valence electrons penetrate considerably into the core region the nonlinearity of \(V_{\text{xc}}\) can produce serious errors. This problem is solved by modeling part of the core density that overlaps strongly with the valence and ”unscreening” the pseudopotential by subtracting \(V_{\text{H}}[\rho_v; r] + V_{\text{xc}}[\rho_c^{\text{mod}} + \rho_v]\) in Eq.(2.9). This, so called, non–linear core correction [42] improves significantly both accuracy and transferability of pseudopotentials.

In this way the norm–conserving pseudopotentials in the semi–local form (local in radial and non–local in angular variable) are obtained. This form is not suitable for practical applications and they are usually transformed into a fully non–local form using a procedure proposed by Kleinman and Bylander [43].

4This approximation is very useful, it simplifies considerably the equations and allows description of the electron dynamics in terms of hydrogen–like atomic orbitals since they can be labeled using the angular momentum quantum numbers. For open shell systems it relies on the assumption that the atomic core is spherical and that the effects of the reduced symmetry are significant only for the valence electrons.

5Formulation of these requirements from Reference [19] is adopted in this work.
On the other hand the norm-conservation condition imposes a lower limit to the number of basis functions because of the difficulty in representing the pseudo-wavefunctions in a plane-wave basis set for systems containing highly localized atomic orbitals (i.e. first row elements and transition metals). It has been shown that this condition can be omitted [44], recasting the original eigenvalue problem into the generalized one (involving the overlap operator). These pseudopotentials, often referred as the "ultrasoft", can also be written in a fully non-local form (with the help of the bracket notation)

\[ V = V_{\text{loc}} + \sum_{i,j} B_{ij} \langle \beta_i \rangle \langle \beta_j \rangle, \]  

(2.10)

where the \( \beta_i \) functions are angular momentum eigenfunctions in the angular variables, times a radial function which vanishes outside the core. Indices \( i \) and \( j \) go over total number of these functions. The \( \beta_i \) functions as well as \( B_{ij} \) coefficients characterize the pseudopotential and differ for different atomic species. As a consequence of abandoning the norm-conserving condition the pseudo-wavefunctions can be much smoother in the core region thus requiring much less plane waves to be accurately represented.

The pseudo-wavefunctions are no longer normalized and the electronic density is expressed in a more complex form

\[ n(r) = \sum_i \left| \phi_i(r) \right|^2 + \sum_{n,m} Q_{nm}(r) \langle \phi_i | \beta_n \rangle \langle \beta_m | \phi_i \rangle, \]  

(2.11)

where \( \phi_i \) are pseudo-wavefunctions and \( Q_{nm} \) are augmentation functions that are localized in the core region and are also provided by the pseudopotential. These augmentation functions serve to reproduce correctly the electronic density inside the core and can also vary rapidly in space. Therefore the problem of rapidly varying pseudo-wavefunctions is transferred to \( Q_{nm} \) functions also requiring large number of basis functions. Solution for this problem is to pseudize the "hardest" part of the augmentation functions. In practice it is done in such a way to preserve all of the charge moments inside the core [45]. This is one more advantage of the ultrasoft compared to the norm-conserving pseudopotentials that reproduce correctly only the total charge inside the core while the information about the core charge distribution is completely lost. This may lead to differences between those two types of pseudopotentials also outside the core\(^6\).

In this work the ultrasoft pseudopotentials are used. They are taken from the pseudopotential library of the DACAPO code [34]. These pseudopotentials are constructed using the code developed in the group of David Vanderbilt [48] and tested by a broader community of users. Of course, the tests have also been performed by the author of this thesis on a number of systems before any serious simulation required by the planning of this project took place. The test systems that were used are bulk fcc iridium and Ir(111) surface, bulk MgO and TiO\(_2\) as well as their surfaces (100) and (110), respectively. Molecular O\(_2\), CO and Ir\(_4\)(CO)\(_{12}\) have also been used for testing purposes. Good agreement with experimental results is achieved which confirmed applicability of these pseudopotentials.

\(^6\)Especially for open-shell atoms due to non-zero higher order charge moments.

### 2.2.2 The Supercell Approximation and Plane Wave Expansions

This thesis is a result of a theoretical study on the properties of the gas-phase Ir clusters as well as Ir clusters adsorbed on an oxide surface. The studied systems are not periodic in...
space. However, it is convenient to make the system artificially periodic by constructing an array of periodically repeated images (supercells) of the original system. This approximation allows one to use theoretical tools developed for the periodic systems such as the plane wave expansions. The plane wave basis set has the advantage of being easily controllable in the sense that the size of basis set can always be increased until the convergence of the desired physical property is reached. The second advantage is a relatively simple and computationally not too expensive algorithms for molecular dynamics (see Sec.2.2.4). Moreover, one could also take the advantage of the highly optimized and parallelized routines for the fast Fourier transformation. However, the disadvantage of the supercell approximation is undesired interaction between the periodic images. If the system is charge neutral and does not carry a dipole moment, these interactions can be suppressed by increasing the supercell size. However, charged or polarized systems suffer from the long range character of the electrostatic interactions regardless of the supercell size. This issue is further elaborated in Sec.2.2.3 and ways to overcome this problem especially for polarized systems are discussed.

The supercell approach is used throughout this work for simulations of small iridium clusters both isolated (in the gas–phase) and adsorbed on an oxide surface. Two existing computer codes DACAPO [34] and QUANTUM-ESPRESSO [35] with the state–of–the–art numerical implementation of the DFT using plane wave basis set are used for calculations.

When studying properties of crystalline (periodic) systems the problem of dealing with an infinite system with infinitely many electrons is simplified by using the Bloch theorem [27] which states that one–electron wave function for an electron in an infinite periodic potential must be of the form

\[ \psi_k(r) = Ce^{i\mathbf{k} \cdot \mathbf{r}} u_k(r), \]  

(2.12)

where \( C \) is the normalization constant and \( u_k(r) \) is a periodic function with the periodicity of the lattice. There are two main consequences of Bloch theorem which enable very efficient calculations of the properties of bulk systems. First, the periodic part of \( \psi(r) \) can be expanded into the plane–waves with the same periodicity

\[ u_k(r) = \sum_{\{G\}} c_{k+G} e^{i\mathbf{G} \cdot \mathbf{r}}; \quad c_{k+G} = \frac{1}{\Omega} \int_{\Omega} u_k(r)e^{-i\mathbf{G} \cdot \mathbf{r}} d\mathbf{r}, \]  

(2.13)

where \( \{G\} \) are the reciprocal lattice vectors and \( \Omega \) is the unit cell volume. Analysis show that the plane–waves with relatively small kinetic energies \( (\hbar^2/2m)|k+G|^2 \) typically contribute more to the wave functions than those with large kinetic energies (see reference [19], p. 93). Therefore, the plane wave basis set can be truncated to include only those plane waves with kinetic energies below a certain cutoff energy. The truncation of the basis set produces errors in the computed properties of the system. However, these errors can be controlled by increasing the cutoff energy until the convergence of the desired property is reached.

Second, and not less important consequence of the Bloch theorem is that solving of the one–electron Schrödinger equation can be simplified by reducing it only to the volume of one unit cell of the crystal. However, it is still necessary to diagonalize one–electron Hamiltonian at infinitely many \( k \) points in the first Brillouin zone of the reciprocal lattice. The fact that the wave functions do not differ much for the two \( k \) points which are sufficiently close, allows a discrete sampling of the Brillouin zone. Moreover, it is possible to define a single \( k \) point (Baldereschi point or the mean value point [28]) at which the value of any function of \( k \) can be used as a good approximation to the integral of the same function over the whole Brillouin zone. Generalization of this idea lead to the special sets or grids of \( k \) points in the
Brillouin zone, two of which are still in use in today’s computer codes (Chadi–Cohen [29] and Monkhorst–Pack [30]).

When a plane wave basis set is used equations 2.2 take very simple form [31]

\[
\sum_{\{G\}} \left[ \frac{1}{2} |k + G|^2 \delta_{GG'} + V_{ion}(G - G') + V_H(G - G') + V_{XC}(G - G') \right] c_{i,k+G'} = \epsilon_{KS}^{i,k+G}.
\]

When expanded into a plane wave basis set the kinetic energy is a diagonal matrix and all potentials are expressed in terms of their Fourier components. Equations 2.14 are then solved by diagonalizing Hamiltonian matrix \( H_{k+G,k+G'} \) given in the brackets above. Although the form of equations 2.14 is simple, matrix diagonalization is not an easy task. Especially for systems that contain localized electronic orbitals (\( i.e. \) core electrons of heavy atoms, valence \( d \) electrons) the number of plane waves needed to make the basis set sufficiently complete grows immensely and those calculations are not tractable even using today’s computers. This problem can be overcome by use of the pseudopotential approximation as discussed in Sec.2.2.1.

The last term \( E_{ion}(\{R_I\}) \) in Eq.2.3 is also not easy to compute. It is very difficult to deal with it by using direct real–space summation due to the long range nature of the Coulomb interaction. Transferring the problem into the reciprocal space also does not help since the Coulomb interaction is long ranged in the reciprocal space, too. The problem was solved by Ewald [32] who divided the Coulomb energy of a periodic array of point charges in two parts, one short and one long ranged. The first part converges fast in the real space while the second converges fast in the reciprocal space. Each of them is then computed in the corresponding fast–converging space as follows \(^8\)

\[
E_{ion} = \frac{1}{2} \sum_{I,J} Z_I Z_J \times \\
\left[ \sum_l \frac{\text{erfc}(\eta |R_I + l - R_J|)}{|R_I + l - R_J|} - \frac{2\eta}{\sqrt{\pi}} \delta_{JJ} + \frac{4\pi}{\Omega} \sum_{G \neq 0} \frac{1}{|G|^2} e^{-\frac{|G|^2}{4\eta^2}} \cos[(R_I - R_J) \cdot G] - \frac{\pi}{\eta^2 \Omega} \right].
\]

(2.15)

The term in brackets is the Coulomb energy due to the interaction between an ion positioned at \( R_J \) and an array of of ions positioned at \( R_I + l \) and is valid for any positive value of \( \eta \). The two summations in brackets become rapidly converging for an appropriate value of \( \eta \). \( Z_I \) and \( Z_J \) are the valences of ions \( I \) and \( J \), respectively. Since the ions do not interact with themselves the \( l = 0 \) term for \( I = J \) needs to be excluded from the summation.

### 2.2.3 Dipole Correction Method

The dipole correction method is used in periodic supercell calculations, when studying nonperiodic systems, to remove electrostatic dipole interactions between the periodic images. These

\(^7\)Divergent \( G = 0 \) term of both ionic and Hartree potentials is not included in Eq.2.14 since it cancells exactly with the same term of the ion–ion interaction when the total energies are computed.

\(^8\)The following formula, in which certain inaccuracies related to the \( G = 0 \) term are resolved, is taken from reference [33].
interactions appear when the system of interest has a nonzero dipole moment. An infinite periodic array of dipole moments leads to a macroscopic electrostatic field which influences the electrostatic potential of the system under consideration.

These problems typically appear when studying adsorption on surfaces. As a consequence of the charge transfer between the surface and the adsorbate the system develops a dipole moment in the direction perpendicular to the surface. One way of dealing with these kinds of errors is to duplicate the system in the unit cell in such a way that the dipole moments cancel. This is done in practice by using the symmetric slab with the adsorbate on both surfaces of the slab. However, the system size is doubled and the calculations are much more expensive which is not favorable for systems containing big number of atoms (of the order of 100 and more).

Second way of removing the dipole field in periodic supercell calculations is proposed by Neugebauer and Scheffler [36]. The idea is to add to the system an artificial planar dipole layer placed far away from the examined system inside the vacuum region, whose dipole strength is calculated selfconsistently in such a way that the adsorbate induced dipole field is compensated for (see Fig.2.1). In this way the asymmetric slabs with the adsorbates only on one side of the slab while the other side is kept fixed mimicking the bulk, can be used and the system size can be kept reasonably small. Moreover, the method can be generalized in order to treat the interaction of a true external electrostatic field with a surface or a molecule. Both possibilities are used in this work, the first to cancel adsorbate induced dipole fields and the second to study how the external field influences relative stability of Ir$_4$ clusters (check Sec.5.6).

### 2.2.4 Ionic Forces and Vibrational Analysis

The electronic structure calculations typically rely on a number of approximations. The first one presented in this thesis is the approximation for the exchange and correlation energy to density functional theory (LDA or GGA). However, the basic approximation used in the
majority of theories that deal with the electronic structure of matter, is the adiabatic or Born–Openheimer approximation [37]. This approximation allows to decouple the electronic degrees of freedom from the ionic ones because of the large difference in their masses. The ionic positions are then treated in the electronic structure calculations as parameters. The total energy of the system (Eq.2.3) can then be viewed as function of the ionic positions $E(\{R_I\})$.

Moreover, the adiabatic approximation also allows relatively simple treatment of the dynamics of the ionic subsystem. The ionic motion is typically described by the classical (Newton) equations of motion for which the forces that are determined by $E(\{R_I\})$ have quantum mechanical origin. The force acting on the ion positioned at $R_J$ is defined as

$$F_J = -\frac{\partial E(\{R_I\})}{\partial R_J} = -\frac{\partial}{\partial R_J} \langle \psi_0(\{R_I\}) | H(\{R_I\}) | \psi_0(\{R_I\}) \rangle - \frac{\partial E_{\text{ion}}(\{R_I\})}{\partial R_J},$$

(2.16)

where $\psi_0(\{R_I\})$ is the electronic ground state, $H(\{R_I\})$ is the Hamiltonian of the electronic subsystem and $E_{\text{ion}}(\{R_I\})$ is the electrostatic ion–ion interaction energy. All these terms depend explicitly on the ionic configuration. Computation of the derivative of the ground–state expectation value of the electronic Hamiltonian can be simplified by using the Hellmann–Feynman theorem that states

$$\frac{\partial}{\partial R_J} \langle \psi_0(\{R_I\}) | H(\{R_I\}) | \psi_0(\{R_I\}) \rangle = \langle \psi_0(\{R_I\}) | \frac{\partial H(\{R_I\})}{\partial R_J} | \psi_0(\{R_I\}) \rangle.$$

(2.17)

Within DFT the only terms in the total energy of the system (Eq.2.3) that depend explicitly on the ionic configuration are the terms that describe electron–ion and ion–ion interactions. Therefore, the expression 2.16 for the force acting on the ion positioned at $R_J$ after the implementation of the relation 2.17 becomes

$$F_J = -\int n_R(r) \frac{\partial V_R(r)}{\partial R_j} \, dr - \frac{\partial E_{\text{ion}}(\{R_I\})}{\partial R_J},$$

(2.18)

where $n_R(r)$ and $V_R(r)$ are the electronic ground state density and the electron–ion interaction potential, both corresponding to the nuclear configuration $R$. This expression allows very efficient simulations of the nuclear dynamics within DFT formalism. The equilibrium geometry of the system is given by the condition $F_J = 0$, that all forces acting on the nuclei vanish. It can be used for finding the equilibrium configurations of the system. Moreover, by introducing the concepts of the ionic temperature and the canonical ensemble or adding history dependant (penalty) potentials in order to forbid the system to adopt already "visited" configurations (scanning the potential energy landscape) various kinds of molecular dynamics can be studied.

The plane wave expansion (Sec.2.2.2) appears to be very useful in calculating ionic forces which can be computed analytically since the nuclear positions enter into the DFT formalism either in the ionic interaction energy (2.15) or as exponents in the structure factors that multiply Fourier components of the ionic potential. Moreover, the plane waves do not depend on the ionic positions inside the unit cell and the Hellmann–Feynman expression for the forces (Eq.2.18) is valid without any corrections (which is not true for the localized basis sets).

On the other hand, the second derivatives of the KS energy functional with respect to the ionic displacements, which are important for studying vibrational properties, cannot be computed analytically since they require calculations of the first derivatives of the ground state
The full vibrational spectrum of the periodic systems can be computed with much less effort using the density functional perturbation theory (DFPT) approach [38]. A linear set of equations similar to KS equations for the variations of the electronic wave functions due to the perturbation of the system (in this case nuclear displacements) needs to be solved in order to compute the (linear) response of the electronic density needed for calculations of the density derivatives with respect to the nuclear displacements. The advantage of the DFPT is that the responses to perturbations with different wavelengths can be decoupled. This allows computations of the phonon frequencies at arbitrary wave vector \( \mathbf{q} \) avoiding the use of the supercells.

Since this work is concentrated on the systems which are mostly nonperiodic (isolated clusters and clusters adsorbed on surfaces) calculations of the vibrational spectra only at the \( \Gamma \)-point are relevant. Therefore, the frozen phonon approach is used for all phonon calculations. At this point it is important to underline that the atomic displacements in these calculations need to be small enough to ensure the validity of the harmonic approximation. This implies that the equilibrium configuration needs to be very well converged. Therefore, in this work all configurations used for studying the vibrational properties are considered as equilibrium configurations when the sum of all the forces acting on ions does not exceed 0.01 eV/Å\(^9\). The atomic displacements used in calculations can then be of the order of 0.01 Å. Such a displacements produce the total energy differences between the ground state and the perturbed configurations of small Ir clusters of the order of 0.001 eV confirming the need for a very well converged structures.

### 2.2.5 Transition–state Theory and The Nudged Elastic Band (NEB) Method

Everything said up to this point is related mostly to the ground state properties of the system. DFT is a theory of the electronic ground state for a given atomic configurations which is not necessarily the ground state for the ionic subsystem. Therefore, the properties of the system that is far from equilibrium can also be studied under the constraint imposed by the adiabatic approximation\(^10\). This is typically done in the molecular dynamics simulations (see for example reference [31]). However, the activated processes in which the system undergoes a

\(^9\)This does not apply to the adsorption of Ir\(_4\) clusters at surfaces. The forces acting on the fixed ions from the surface bottom layer may be bigger and only the forces acting on nuclei which are allowed to relax are taken into account.

\(^10\)That the electrons remain at each moment at their ground state for any reasonable atomic configuration.
transition from one local minimum to another across an energy barrier are extremely difficult to study by the means of standard molecular dynamics. Processes with a relatively low activation energy of 0.5 eV would require years of computer time to simulate classical trajectory of the system in order for a single transition event to be expected to occur.

This problem can be overcome and accurate estimates of the transition rates can be obtained by using the transition state theory (TST) [39]. In addition to the adiabatic approximation TST relies on two assumptions: (i) the rates are slow enough and a Boltzmann distribution describes the reactant state; and (ii) the initial and final states can be separated in the configurational space by a D-1 dimensional surface (D is the number of degrees of freedom) in such a way that the trajectory going from one state to the other crosses this surface only once. Since the range temperatures of interest, when considering condensed matter systems, lays typically far below the melting temperature, the harmonic approximation to TST (hTST) can be used for studying diffusion or reactions at crystal surfaces. This simplifies the problem which becomes that of finding the saddle point (transition state) along the minimal energy path (MEP) connecting the two states (initial and final). The rate constant for the transition can be obtained from the energy and the frequency of the normal modes of the initial and transition state [41]

\[
k_{\text{hTST}} = \frac{\prod_{i=1}^{3N} \nu_i^{\text{init}}}{\prod_{i=1}^{3N-1} \nu_i^{*}} e^{-(E^{*} - E^{\text{init}})/k_B T},
\]

(2.19)

where \(E^{*}\) and \(E^{\text{init}}\) are the energies of the saddle point and initial state, respectively. The \(\nu_i\) appearing in the prefactor that multiplies the exponential term in Eq.2.19 are the corresponding normal mode frequencies. In the cases where transition occurs along one of the normal modes of the initial states the whole prefactor or the attempt frequency could be approximated with the frequency of that mode solely. Moreover, in the solid state systems the frequencies are of the order of \(10^{12} – 10^{13}\) Hz. These values are often used in the scientific community for rough estimations of the rate constants.

The MEP often has one or more minima in addition to those at the initial and final states which correspond to the (meta)stable intermediate configurations. Therefore, along the MEP there are also several saddle points, one between each pair of the neighboring minima. If a Boltzmann population is reached for the intermediate configurations the overall rate is then determined by the highest transition state.

Various methods are in use for calculating MEPs and finding the transition states. One of the most efficient is the nudged elastic band (NEB) method [40]. It has been widely used together with hTST. The idea is to discretize the initial path (guess) in the configurational space of the system between the two end points and to relax each of the resulting configurations only in the directions perpendicular to the path. This is done by putting all components of the forces that lay along the path to zero and performing standard minimization techniques. However, the definition of the tangent as well as the number of points used for discretization is crucial (see more about this in reference [40]). Since the relaxation of each point depends on the tangent and therefore on the configurations of other points, the relaxation towards the MEP is performed simultaneously for all points along the path. To prevent grouping of the images and other types of path instabilities the spring forces are added between neighboring configurations making an elastic band in the configurational space. The tangential components of the spring force are used to control the spacing between images (normal components are set equal to zero). Calculations of the energies and forces of different configurations are typically done using DFT methods.
The climbing image nudged elastic band method (CINEB) [41] can then be used to find the saddle points along the path. After the initial path has approached closer to the MEP the appropriately chosen configuration (or configurations) is detached from the springs and allowed to relax also in the tangential direction. This relaxation is done with inverted the tangential components of the forces. In this way the chosen configuration climbs to the point with the zero force, which is a minimum in all directions perpendicular to the path and a maximum in the tangential direction, the first order saddle point or the transition state.

In this work the CINEB method is mostly used to calculate barriers for structural transformations between different Ir$_4$ isomers. It is also used for calculations of the MEPs and finding the transition states for the CO dissociation on both gas-phase and MgO(100) supported Ir$_4$ clusters. Pathways between different local minima are discretized and minimal set of five intermediate configurations is used at the beginning of simulations. Saddle points are found, for a fixed number of images along the MEP, by relaxing appropriately chosen intermediate configuration (configurations) along the path in the direction opposite to the forces, until the total force in this configuration reached the threshold of 0.05 eV/Å. Afterwards, the number of images is increased and the convergence is achieved when energy of the saddle point (converged for a fixed number of images) changes less than 0.05 eV.
Chapter 3

Gas–phase Iridium Clusters

3.1 Iridium Atom

Iridium is a 5d transition metal with atomic number $Z = 77$ which, in addition to the electronic structure of Xe atom, has completely filled 4f and outer 6s subshell while the remaining seven electrons are accommodated in ten 5d orbitals ([Xe]4f$^{14}$5d$^7$6s$^2$) giving rise to magnetic moment of $3 \mu_B$. The 5d and 6s electrons are usually treated as valence when constructing iridium pseudopotential, as it is done in this work.

As already mentioned in Sec.2.2.1 the ultrasoft pseudopotentials (USPP) are used throughout this work. They are constructed within the GGA framework with exchange and correlation energies described by PW91 XC functional [21]. The non–linear core–corrections (Sec.2.2.1) are added to iridium PP in order to take into account the nonlinearity of the exchange interaction between the valence and the core electrons. Relativistic effects are treated at the scalar level [94] neglecting the spin–orbit interaction. Bulk fcc iridium is used as the test system. Sampling of the fcc iridium Brillouin zone is done with $10 \times 10 \times 10$ Monkhorst–Pack k–point grid [30]. The same plane wave and density cutoffs, which determine the size of the plane wave basis set, are used as for simulations of small Ir clusters (see below). Bulk parameters, the equilibrium lattice constant $a_0 = 3.85 \ \text{Å}$, bulk modulus $B_0 = 356 \ \text{GPa}$ and cohesive energy $E_c = 7.48 \ \text{eV}$, are obtained by fitting numerical results to the Murnaghan equation of state [49]. These results are in line with other calculations [50, 51]. When compared to experimental values ($a_0 = 3.84 \ \text{Å}, B_0 = 306 − 355 \ \text{GPa}$ and $E_c = 6.94 \ \text{eV}$ [52, 53]) it can be seen that the lattice constant is well reproduced (error $\sim 0.3 \%$) while $E_c$ is overestimated by about 8 % which are typical features of DFT methods.

In Fig.3.1 Kohn–Sham electronic spectrum of Ir atom is shown. Calculations are performed in a cubic unit cell with linear dimension $\sim 10.58 \ \text{Å}$ (20 Bohr). This cell size provides good convergence of the properties of small Ir clusters (cohesive energy, interatomic distances,...). Electronic wavefunctions are expanded in plane waves with the kinetic energy cutoff of $25 \times 13.606 \ \text{eV}$. Cutoff for expanding the electronic density is set to $140 \times 13.606 \ \text{eV}$ ensuring appropriate description of the pseudopotential augmentation functions (see Sec.2.2.1). These cutoffs provide sufficiently complete basis set for simulating small Ir clusters. Converged values of the physical quantities of interest, cohesive energies of small Ir clusters in particular, are obtained in this way.$^1$

$^1$Cohesive energy of iridium dimer changes by less than 0.02 eV when the cutoffs increase to $30 \times 13.606$ and $160 \times 13.606 \ \text{eV}$, respectively.
Calculated population of atomic orbitals of Ir atom \((5d^76s^2)\) compares well with experiment. However, this result is dependent on the super-cell symmetry. Due to the fact that \(d\)–shell is not completely filled there is a number of Slater determinants close in energy which are candidates for the atomic ground state. Different determinants are the ground state of Ir atom for different boundary conditions. This affects the atomic total energy, \(i.e.\) total energies of Ir atom calculated in hexagonal and triclinic super–cell differ by \(\sim 0.12\) eV for the same nearest neighbor distance. Energy of an atom is typically used as a reference for calculating cohesive energies of more complicated systems. Different reference atom is one of the reasons for differences in cohesive energies of transition metal clusters reported in literature.

Cubic super–cell is chosen in this work for simulating Ir atom since, due to symmetry reasons, it does not allow mixing of \(s\) and \(d\) angular momentum channels which is the closest possible to the situation of an isolated Ir atom. However, it leads to cubic splitting of \(5d\) orbitals into \(t_{2g}\) triplet and \(e_g\) doublet\(^2\). This splitting is also a consequence of the symmetry of \(d\) orbitals and the number of electrons filling them. In order to minimize Coulomb repulsion, two or three \(d\) electrons in the spin minority channel will always tend to occupy only one \(d\) orbital in \(xy\) plane \((d_{x^2-y^2} \text{ or } d_{xy})\) and one or two orbitals orthogonal to that \((d_{z^2} \text{ or } d_{xz} \text{ and } d_{yz})\) which will lead to electronic density with the octahedral or cubic symmetry and consequently to \(t_{2g}-e_g\) splitting of \(d\) orbitals.

3.2 Ir Dimer

Results for iridium dimer are presented separately from bigger \(\text{Ir}_n\) \((n = 3, \ldots, 8)\) clusters since the dimer presents a basis for studying \(\text{Ir}-\text{Ir}\) bonding mechanism. Two reviews on transition metal dimers appeared in literature in the last 25 years \([56, 57]\). Both authors find a lack of experimental data on iridium dimer. They report only empirical estimates of the dimer.

\(^2\)The term *cubic* is used in the sense that the splitting is produced by reducing the symmetry from rotational \(\text{SO}(3)\) to \(\text{O}_h\) group which is a symmetry group of both cube and octahedron. The \(t_{2g}\) and \(e_g\) denote respectively, triply and doubly degenerate irreducible representations of \(\text{O}_h\) group.
dissociation energy of 3.7 [56] and 3.5 eV [57] with the error bars as big as \( \sim 1 \) eV. On the other hand, several theoretical studies have been published recently treating the group VII transition metal dimers and comparing performances of different numerical approaches based on DFT [59, 60, 61]. It is shown that dissociation energies, magnetic moments as well as vibrational frequencies depend strongly on the XC functional used, while the dependence is less pronounced for the dimer bond distance. Moreover, in references [59, 61] the authors show that "pure" DFT methods (GGA) yield results in better agreement with experiments (in the cases where experimental results exist) than hybrid DFT schemes which are usually considered as more accurate. In addition to these, there is one GGA (PW91) study of the gas-phase iridium clusters consisting of 2–64 atoms [64] and one multiconfigurational self–consistent–field investigation on hydrogen activation by iridium dimers [62]. In both cases results for clean \( \text{Ir}_2 \) go in line with those obtained with corresponding method from the aforementioned references.

Properties of iridium dimer obtained in this work are presented in Tab.3.1 together with the bigger Ir clusters. Numerical setup is discussed in the previous section. Cohesive energy of 2.52 eV/atom, calculated as the difference \( E_c = -\left( E(\text{Ir}_2) - 2E(\text{Ir}) \right)/2 \), agrees well with the value of 2.53 eV reported in reference [64] obtained using similar approach (ultrasoft PP and PW91 XC functional). This value together with calculated frequency of the dimer stretching mode \( \hbar \omega_e = 33.6 \) meV lays within the range (\( E_c = 1.62–3.02 \) eV and \( \hbar \omega_e = 33.2–37.3 \) meV) of all–electron GGA calculations [60, 61]. Spread in the reported \( E_c \) values is typically quite big for transition metal clusters. There are two main reasons for this: (i) \( E_c \) is calculated using total energy of an atom as a reference and, as already mentioned in Sec.3.1, different approaches yield different atomic ground state; and (ii) since molecular orbitals in transition metal clusters are formed either exclusively of atomic \( d \) orbitals or of \( dsp \) hybrid orbitals (see later in the text and in Sec.3.4) differences also arise from different description of the level of \( dsp \) hybridization which is very sensitive to the details of numerical approach.

The dimer bond distance of 2.17 Å calculated in this work is shorter than those (2.24–2.29 Å) reported in references [60, 61] and considerably shorter than 2.43 Å from reference [62]. Since, the interatomic distances in small Ir clusters from this work agree well with results from other calculations using pseudopotentials (see for example reference [65]) it might be concluded that somewhat shorter distances result from the pseudopotential approximation. However, test calculations with the pseudopotential that includes semicore 5s and 5p atomic orbitals resulted in the dimer bond length of 2.19 Å, only slightly bigger. Therefore, the differences in the Ir–Ir distance between pseudopotential and all–electron calculations probably reflect differences in numerical approaches (i.e. local versus plane wave basis sets,...) and the sensitivity of the \( d–s \) hybridization level on the details of the numerical scheme3.

Bonding mechanism in \( \text{Ir}_2 \) is similar to other platinum group transition metal dimers with partially filled \( d \)–shell. As it is usually done when studying bonding in molecules, molecular orbitals (MO) formed upon interaction, can be classified (and labeled) according to irreducible representations of the corresponding symmetry group. If a spherical symmetry is assumed for Ir atom then, by forming the dimer, symmetry is lowered to \( D_{\infty h} \) axial point group containing rotational axis of the infinite order and two mirror planes, vertical \( \sigma_v \) and horizontal \( \sigma_h \). Since the vertical mirror plane associates atomic orbitals (AO) with opposite \( m_l \) values (angular

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3This can by supported by the fact that just by replacing the ultrasoft with norm–conserving pseudopotential results for the dimer change (\( d = 2.21 \) Å and \( E_c = 2.36 \) eV).

4Due to the infinite order of the principal rotational axis there are infinitely many equivalent vertical mirror planes, but only one horizontal.
Figure 3.2: Spin resolved DOS of iridium dimer (↑–spin maj. and ↓–spin min.). Gaussian broadening with the width of 0.07 eV is employed. The Fermi energy is positioned in the middle of the HOMO–LUMO gap. Projections onto orthogonalized atomic orbitals (see reference [55]) are also shown. Molecular orbitals, displayed in the left part of the figure, are labeled according to the symmetry properties with respect to $D_{4h}$ symmetry group (see text for details).

momentum along the principal axis) the 5$d$ multiplets on each atom will split to $m_l = 0$ singlets ($d_{z^2}$) and $|m_l| = 1, 2$ doublets ($\{d_{xz}, d_{yz}\}$ and $\{d_{x^2−y^2}, d_{xy}\}$). The consequence of $\sigma_h$ is mixing the orbitals from different Ir atoms. Therefore, AOs from both atoms with the same $|m_l|$ form a subspace that can be reduced, using the parity with respect spatial inversion (equivalently the parity with respect to $\sigma_h$ can be used), to one or two dimensional even (g) or odd (u) irreducible representations of $D_{\infty h}$ group.

Population of molecular orbitals (MO) reduces further the symmetry of the system. There is only one electron filling the antibonding spin minority $E_{2u}$ doublet of the $D_{\infty h}$ group. Given the shape of any of the two $E_{2u}$ orbitals (antibonding combination of either $d_{xy}$ or $d_{x^2−y^2}$ atomic orbitals from the two atoms) symmetry of the electron density is lowered to $D_{4h}$ group. This produces splitting of both $E_{2u}$ and $E_{2g}$ doublets to $B_{1u}$ and $B_{2u}$ or $B_{1g}$ and $B_{2g}$ singlets of $D_{4h}$ group, respectively. Therefore, the labeling of molecular orbitals is done according to the irreducible representations of $D_{4h}$ rather than $D_{\infty h}$ axial point group (see App.B). In Fig.3.2 electronic density of states (DOS) of Ir$_2$ is shown together with DOS projected onto orthogonalized atomic orbitals using the scheme proposed by Löwdin [55]. Reduction of $D_{\infty h}$ symmetry due to population of MOs can be seen.

Symmetry also provides an information about possible hybridizations. Only molecular orbitals belonging to the same irreducible representations are allowed to hybridize. In the case of Ir$_2$ this implies that hybridization is allowed between 5$d_{z^2}$, 6$s$ and 6$p_z$ as well as 5$d_{xz}$, 5$d_{yz}$, 6$p_x$ and 6$p_y$ AOs from the same atom. From Fig.3.2 it is clear that only the first type
of hybridization is realized. Moreover, contribution of $6p_z$ orbital in the electronic spectrum is appreciable only at higher energies while in the lower part mainly $6s$ and $5d_{z^2}$ orbitals are mixed. Two $\sigma$ bonds are formed, one strong (lowest $A_{1g}$ orbital) and the other weaker (higher $A_{1g}$). In addition to the $\sigma$ bonding there is also an interaction between $5d$ orbitals from the two atoms forming $\pi$ and $\delta$ bonds. This type of interaction is responsible for creating bonding $E_u$ ($\pi$ bond) doublet and two $B_{1g}$ and $B_{2g}$ singlets ($\delta$ bonds) as well as the corresponding antibonding $E_g$, $B_{2u}$ and $B_{1u}$ orbitals. If the simplest possible definition of the bond order is adopted:

$$B.O. = \frac{\text{number of bonding electrons} - \text{number of antibonding electrons}}{2},$$

then there is a triple bond connecting two Ir atoms.

### 3.3 Ir$_n$ ($n = 3, \ldots, 8$) Clusters

Free (gas–phase) iridium clusters were not extensively studied in the past, both experimentally and theoretically. While there are no experimental studies of the gas–phase as well as Ir clusters embedded in the rare gas matrices, several theoretical papers exist in literature. As already mentioned in the previous section only one systematic study on Ir$_n$ ($n = 2, \ldots, 64$) is published up to now [64]. The authors use method similar to that used in this work (PW91 and USPP). Two other more comprehensive studies of small Ir clusters also exist. They treat Ir$_n$ clusters with $n = 4, 8, 9, 12, 13$ [65] or $n = 4, 6, 8, 10$ [66]. While the former is a comparative study of the properties of Ru, Rh, Pd, Ir and Pt clusters performed by the same authors as of reference [64], results in the latter are obtained with the Hartree–Fock method [67].

Results for Ir$_n$ ($n = 2, \ldots, 8$) clusters, obtained in this work, are presented in Tab.3.1. Properties of the most stable isomers (with the lowest total energy) are presented together with the properties of their low–energy counterparts. For Ir$_4$ and Ir$_6$, except the data for the most stable structure and its closest isomer, the results for tetrahedral and octahedral structures, which are found in experiments treating supported Ir$_4$ and Ir$_6$ clusters [118, 119], are also given. Moreover, for Ir$_7$ three structures lay very close in energy and the properties of all three are presented.

Equilibrium atomic configurations are found by optimizing all degrees of freedom (including spin) of the certain number of starting configurations. For a small number of atoms there are several relevant starting configurations. However, as the number of atoms increases the number of possibilities grows immensely. Starting configurations are then chosen according to a certain group of rules (truncation from the bulk, symmetries different from the bulk, adding atoms to the stable configurations of smaller clusters,...). In the first stage, calculations are performed in the spin unrestricted fashion. Candidates for the local minima with their total energies and total $S_z$ ($z$–component of the spin) are obtained in this way. Afterwards, for each geometry the spin dependence of the total energy is investigated by performing geometry optimization with the fixed spin for several lower and higher values in order to refine the group of potential configurations. The vibrational analysis is then used for final verification of the stability of these configurations.

Configurations with the lowest total energy are, of course, the most probably realized in experiments and therefore are the most important to study. However, in the opinion
Table 3.1: The most stable Irₙ (n ≤ 8) structures are shown together with their isomers closest in energy. For Ir₄ and Ir₆ all structures between the most stable one and that found in experiments (tetrahedron and octahedron, respectively) are shown. Cohesive energies Eₖ, average nearest neighbor distances d₁ and total cluster magnetic moments per atom m are presented.

<table>
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<th>d₁ (Å)</th>
<th>m (µ₊/atom)</th>
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<td>D₂h</td>
<td>4.68</td>
<td>2.43</td>
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</table>

of the author of this thesis, importance of the configurations that are higher in energy is underestimated in the scientific community. Their properties are presented in the scientific publications mainly as a proof that all relevant configurations are taken into account and that the most stable one represents the true global minimum. Only in the cases where there is a discrepancy with experiments such as Ir₄ [69] or where several nearly degenerate configurations exist (like the situation with Pt₄ [77, 76]) properties of the low energy structures are discussed in more details. It is shown in this work (Chapters 4, 5 and 6) that the isomers which are as much as ∼1.5 eV higher in energy may become relevant due to the interaction with adsorbates and/or support.

For Ir₃ the linear configuration with the total magnetic moment of 1 µ₊ is found to be the most stable one. It is by 0.36 eV lower in energy than the isosceles triangle which carries a
magnetic moment of $3 \mu_B$. The equilateral triangle ($D_{3h}$ symmetry) is Jahn–Teller unstable due to a large number of electronic levels close to the Fermi energy. Differences in the Ir–Ir coordination of the two isomers are reflected in the different average interatomic distance which is by 0.19 Å shorter in the case of linear isomer. Results for the triangle agree very well with those reported in reference [64] ($E_c = 3.03$ eV and $m = 1.00 \mu_B$). However, the authors do not present at all, properties of the linear Ir$_3$. On the other hand, the accord is not that good with the results obtained with the relativistic configuration interaction (RCI) method [63]. The authors treated only triangular Ir$_3$ and found, in the absence of spin–orbit (SO) coupling, nearly equilateral triangle with $d_{sr} \approx 2.52$ Å as the ground state with the magnetic moment of $5 \mu_B$. The effect of SO coupling is mixing the $m = 5 \mu_B$ ground state with $m = 7 \mu_B$ first excited state which results in increasing $d_{sr}$ to $\sim 2.7$ Å. However, the average RCI Ir–Ir distance in triangular Ir$_3$ is close to the experimental values of the Ir–Ir nearest neighbor distance in bulk iridium and in Ir$_4$(CO)$_{12}$ molecule (2.72 and 2.69 Å, respectively) where iridium atoms are highly coordinated.

Results for Ir$_4$ are discussed in details in Sec.3.4. For Ir$_5$ two nearly degenerate isomers are found well separated in energy from all the others (about 0.9 eV). Both structures are formed by adding one Ir atom to the square configurations. The square pyramid carrying magnetic moment of $7 \mu_B$ is the lowest in energy, only by 0.05 eV more stable than planar configuration where the added Ir atom bridges one side of the square. The latter structure is also magnetic and carries the same magnetic moment. The average Ir–Ir first neighbor distance is longer in the case of pyramid by 0.09 Å. These results agree also very well with those from reference [64].

Total energy of the trigonal prism, the most stable Ir$_6$ isomer, is by 0.42 and 0.78 eV lower than that of planar and octahedral isomers, respectively. All three structures are magnetic with the total cluster magnetic moments of 8 (prism), 4 (planar) and 12 $\mu_B$ (octahedral). Both perfectly symmetric prism and octahedron are Jahn–Teller (JT) unstable and lower their symmetries to $D_{1h}$ and $D_{4h}$, respectively. Energies of JT deformations are about $\sim 0.1$ eV in both cases. Planar isomer has the lowest average Ir–Ir first shell coordination number and hence, the shortest average Ir–Ir first shell distance (2.31 Å). As the coordination number grows from 3 (prism) to 4 (octahedral structure) the Ir–Ir distance increases from 2.41 to 2.52 Å, respectively.

All theoretical studies treating the gas–phase Ir$_6$ [64, 66, 68] agree that the JT deformed trigonal prism is the most stable structure. However, details of the electronic ground state and the energy difference to the octahedral isomer are different. In reference [64] the prism has 8 unpaired electrons and is by 0.72 and 1.44 eV more stable than planar and octahedral structure each carrying magnetic moments of 8 and 6 $\mu_B$, respectively. On the other hand, scalar relativistic all–electron DFT calculations [68] yield the prism with the moment of 6 $\mu_B$ and $d_{sr} = 2.42$ Å which is by 1.06 eV more stable than octahedral isomer (trigonal antiprism). Hartree–Fock calculations [66] also yield trigonal prism as the most stable Ir$_6$ isomer, but the authors do not report values of total energies. Furthermore, calculated Ir–Ir distances are much bigger ($\sim 2.59$ Å for the prism) than those reported here. As it is already mentioned, complexity in transition metal clusters arises from the large number of low laying many–electron states and even small differences between computational approaches may give different ground states which consequently produce differences in isomer relative energies.

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5 They say for Ir$_4$ on the fourth page, sixth line "the calculated total energies are omitted here for simplicity".

6 Again, test calculations with norm conserving pseudopotential for iridium give difference between the
Three configurations very close in energy exist in the case of Ir$_7$. All three are obtained by adding one Ir atom to the Ir$_6$ trigonal prism. The most stable isomer is the one with C$_{2v}$ symmetry where the added Ir atom "sits" on one side of the prism while at the other two it is positioned either on top of the base (C$_{3v}$) or bridges one edge of the prism (C$_{2v}$). The lowest energy structure is only by 0.14 and 0.15 eV more stable than the other two (following the same order like in Tab.3.1). Each isomer carries a nonzero magnetic moment which decreases from 11 $\mu_B$ for the most stable structure to 9 and 7 $\mu_B$ for the less stable C$_{3v}$ and C$_{2v}$ configurations, respectively. The accord with the results from reference [64] is as good as for Ir$_6$. The most stable structure is the same, with the same magnetic moment, but the reported cohesive energy is by 0.05 eV higher. The second lowest structure is the C$_{2v}$ one (with the same magnetization), while the isomer with C$_{3v}$ symmetry is missing. Reported cohesive energy for that structure is only 0.01 eV higher than the value reported here which leads to considerably different relative energy of 0.42 eV.

Perfect cubic structure is by far the most stable Ir$_8$ configuration. The energy gain compared to eight isolated Ir atoms is 4.92 eV/atom, by 0.24 eV/atom higher that for the nearest (in energy) double–prism configuration. The difference between cohesive energies of the two isomers is similar to the Ir$_4$ case and bigger than for all the other clusters sizes. Hence, it implies that Ir prefers using the square geometry as the building block when forming small clusters. This is the point where all authors agree [64, 65, 68, 69]. The electronic ground state of the cube is nonmagnetic while the double–prism structure has eight unpaired electrons. Great stability of the cubic structure is also reflected by the drop of the average nearest neighbor Ir–Ir distance from 2.48 Å in Ir$_7$ to 2.34 Å. Results for Ir$_8$ agree well with those from references [64, 65].

In Fig.3.3 results from Tab.3.1 are represented graphically. Cohesive energy, average nearest neighbor Ir–Ir distance and magnetic moment per atom of Ir$_n$ (n = 2,...,8) are shown as a function of the cluster size. The dependence of the HOMO-LUMO gap is not presented here (at variance to other studies of the gas–phase clusters where it is usually done) since it is known that DFT methods underestimate this quantity (see Ref.[19]). It cannot be expected that the error is similar for all cluster sizes since the HOMO–LUMO gap depends strongly on the cluster geometry. Therefore, calculated trends in the HOMO-LUMO gap are not considered as reliable.

Cohesive energy grows monotonically with the cluster size. In reference [64] similar trend up to the eight atom cluster is reported. The authors also investigated bigger clusters and found much slower growth of $E_c$ when going from Ir$_8$ to Ir$_9$, from Ir$_{10}$ to Ir$_{11}$ and Ir$_{12}$ to Ir$_{13}$ which they attributed to the increased stability of the 8, 10 and 12 atom clusters having the simple cubic structure as the building block. They also found that up to 48 atoms iridium clusters prefer simple cubic instead of fcc arrangement which is much more than for other transition metals (only Ru shows similar behavior [82] while for Pt transition occurs at the size of 13 atoms [64]). In addition to the trend for the most stable isomers, it is reported here that the dependence of $E_c$ is similar also for structures nearest in energy. Moreover, as it is shown in Fig.3.3, low energy structures are quite close in $E_c$, except for Ir$_4$ and Ir$_8$ where strong preference towards square and cubic structures is found. In Sec.3.4 it is proved on Ir$_4$ clusters that the strong preference towards cubic–like structures is mainly due to very favorable d–s hybridization in iridium clusters.

However, results from EXAFS [133] measurements done on the MgO, TiO$_2$ and Al$_2$O$_3$ prism and octahedral structure of only 0.41 eV.


Figure 3.3: Properties of Irₙ (n = 2, ..., 8) clusters: cohesive energy $E_c$, average nearest neighbor distance $d_{sr}$ and magnetic moment per atom $m$, as a function of the cluster size. Data for the most stable isomers are presented as well as for those which are nearest in energy. For Ir₄ and Ir₆ properties of the tetrahedral and octahedral geometries, found in experiments for the supported clusters, together with experimental $d_{sr}$ are also given. Numbers in the middle graph denote average first shell Ir–Ir coordination numbers.

supported Ir₄ and Ir₆ clusters [118, 119, 126] do not correspond to the theoretically predicted stability. From the measured first shell Ir–Ir coordination number (CN) the authors concluded that, independently on the support, Ir₄ is tetrahedral while Ir₆ adopts octahedral shape. It also follows from the measured interatomic distances (check discussion below) that these discrepancies between theory and experiment may be attributed to interactions of the clusters with support and/or adsorbed atomic or molecular species. It is shown in this thesis that, indeed, the interaction with oxygen vacancies on MgO(100) surface (Sec.5.5) or clean TiO₂(110) (Sec.5.6) surface as well as with C adatom or CO molecules (Chapters 4 and 6) can
considerably affect relative energies between Ir$_4$ isomers making tetrahedral structure either degenerate or more stable than the square.

The average nearest neighbor Ir–Ir distance ($d_{sr}$) does not show monotonic behavior. The overall trend for the most stable isomers is growing with the bulk value as the upper limit, but in the cases of Ir$_3$, Ir$_6$ and Ir$_8$, $d_{sr}$ decreases. By analyzing correlations with the average coordination of Ir atoms in the clusters it can be concluded that $d_{sr}$ depends more on the average first shell Ir–Ir CN (shown in Fig. 3.3) than on the cluster size. For the most stable Ir$_6$ and Ir$_8$ structures CN decreases from the values it has for the corresponding smaller clusters which is followed by the drop in $d_{sr}$. Only in the case of Ir$_3$ the $d_{sr}$ shows opposite behavior. While first shell CN increases from 1 (dimer) to 1.3 (linear trimer) the average distance decreases from 2.17 to 2.15 Å. This is another manifestation of the strong $d$–$s$ hybridization which favors linear Ir$_3$ configuration more than the triangle. Particularly strong Ir–Ir bonds are formed in linear Ir$_3$ with better screening of the nuclei enabling shorter interatomic distances. However, the dependence of $d_{sr}$ on the first shell CN is supported by the fact that for a fixed number of atoms $d_{sr}$ is always the shortest for the isomer having smallest CN which is true also for Ir$_3$.

Experimental values of $d_{sr}$ for MgO, TiO$_2$ and Al$_2$O$_3$ supported Ir$_4$ and Ir$_6$ clusters [118, 119, 126] (Fig.3.3) lay in the range 2.62–2.73 and 2.61–2.62 Å for Ir$_4$ and Ir$_6$, respectively. They are noticeably longer than the calculated values of the corresponding gas–phase structures and it follows from the experimental results that do not depend appreciably on the type of the support. These values are close to the bulk nearest neighbor distance of 2.72 Å [52, 53] and $d_{sr} = 2.69$ Å in Ir$_4$(CO)$_{12}$ molecule [16]. In these systems the Ir atoms are highly coordinated suggesting that interaction of Ir$_4$ and Ir$_6$ clusters with their environment (support and/or ligands) is responsible for the differences between theoretically predicted isomer stability and experimental results.

Magnetization of small Ir clusters also does not show monotonic behavior. It is, even not possible to anticipate the decreasing trend towards nonmagnetic bulk fcc iridium from the values for Ir$_n$ clusters with $n \leq 8$. This is not surprising since the magnetic moment is highly sensitive to geometry. In reference [64] the authors show that starting from 9 atoms, magnetic moment is lower than 0.5 $\mu_B$/atom for all studied clusters. However, since iridium has odd atomic number the nonmagnetic solution is not possible for clusters with odd number of atoms and magnetic moment shows oscillatory behavior also for big $n$ values.

Bonding mechanism in Ir$_n$ ($n \leq 8$) is similar to that in Ir$_2$ that is explained in Sec.3.2. Molecular orbitals of the clusters are either of purely $d$ or mixed $dsp$ character. The level of $dsp$ hybridization strongly depends on the cluster geometry and is different for different isomers. This issue is elaborated in details on the example of Ir$_4$ clusters which are the subject of the following chapter.

### 3.4 The Case of Ir$_4$

Gas–phase Ir$_4$ clusters have been studied up to now only theoretically [64, 65, 69, 66]. Experimental studies treat only supported species [12, 118, 126]. Iridium tetrarners are prepared

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7Three $s$ orbitals with one electron per orbital (three H atoms) form linear trimer rather than triangular. This is due to the same number of bonding molecular orbitals that are formed in two structures (there is exactly one) and more destabilizing occupied antibonding orbital in the case of triangle. Look in Sec.3.4.1 for a more detailed discussion of this issue.
Figure 3.4: Three Ir$_4$ isomers (T, B and S) are shown together with the configurations of two transition states (TS1 and TS2) for structural transformation from T to B and B to S. Numbers beside each structure represent the Ir–Ir distances (in Å) or the butterfly bending angles. Relative energies (with respect to T) together with energy barriers are also displayed as well as vibrational spectra of the three stable structures. Length of the line representing certain vibrational frequency is proportional to the degeneracy of the corresponding mode. Vibrational modes are labeled according to their symmetry properties (see Tab.3.1 for the symmetries of Ir$_4$ clusters and App.B for labeling of their irreducible representations).

experimentally by decarbonylating Ir$_4$(CO)$_{12}$ complexes which are previously deposited over an oxide substrate. It follows from the EXAFS measurements (see reference [119] and the references therein) that, after the treatment, tetrahedral metal frame of the carbonyl molecules remains intact.

On the other hand, authors of theoretical studies on Ir$_4$ [64, 65, 69] agree that the most stable Ir$_4$ isomer is perfect square (S) followed by butterfly structure (rhombus which is bent around its shorter diagonal further denoted as B) and tetrahedron (T). All mentioned studies are done with DFT methods. Only one theoretical Hartree–Fock study of Ir$_4$ exists [66] whose authors also report that B isomer is more stable than T, but they do not present at all results for S structure.

Results for the gas–phase Ir$_4$ obtained in this work are presented in Tab.3.1 and Fig.3.4. Three relevant configurations (T, B and S) are found following the procedure for searching stable configurations explained in Sec.3.3. Namely, structures which are not stable with
respect to the magnetic moment variations (to the nearest possible values) are not considered as local minima.

It is confirmed here that S structure with eight unpaired electrons is the most stable form of the gas–phase Ir₄. It has cohesive energy 3.77 eV/atom and nearest neighbor distance 2.31 Å. The method that is used in this work for modeling iridium clusters (see Chapters 2 and 3) yields nondegenerate electronic ground state of S structure and hence, there is no Jahn–Teller distortion. Stability of this structure is also confirmed by the vibrational analysis (see Fig. 3.4). Six normal modes lay in the range 10.0–30.2 meV. Highest energy is the so called "breathing" mode (all interatomic distances increase) while the lowest in energy is antisymmetric out–of–plane mode which bends S structure towards B configuration.

The first less stable equilibrium configuration is B with the total cluster magnetic moment of 2 μₖ. It is by 0.93 eV less stable than S structure. Its average nearest neighbor Ir–Ir distance is 2.36 Å while the distance between two wing tip Ir atoms is 3.78 Å. The bending angle (angle between "wings" of the butterfly) is 138.8°. Range of frequencies of the vibrational modes is somewhat bigger compared to S isomer. The breathing mode (33.0 meV) is the highest in energy as well, but lays ∼3 meV higher, while the bending mode (7.5 meV) is also lowest in energy and lays ∼3 meV lower than the corresponding mode of S structure.

Authors of reference [64] report rhombus (butterfly with zero bending angle and D₄h symmetry) carrying magnetic moment of 6 μₖ as the only structure between T and S. It is found in this work that the rhombic configuration with 6 unpaired electrons is indeed one of the candidates for the local minima, but it is higher in energy than B isomer by 0.18 eV and unstable with respect to the change of magnetic moment. Increasing magnetic moment to 8 μₖ results in structural relaxations which transform rhombus into S configuration.

The least stable Ir₄ isomer whose properties are presented in this thesis is T structure, with the total energy by 0.58 and 1.51 eV higher than that of B and S, respectively. Its nondegenerate electronic ground state is also nonmagnetic. Of all three structures T isomer has the longest nearest neighbor Ir–Ir distance 2.44 Å. As discussed in Sec.3.3 this follows from the largest first shell Ir–Ir coordination number. Due to symmetry reasons (degeneracies) its low energy modes are shifted to higher frequencies compared to corresponding low energy sectors of the B and S vibrational spectra. The lowest frequency in the spectrum of T isomer is 17.3 meV and is triply degenerate. On the other hand frequency of the breathing mode, which is totally symmetric and therefore nondegenerate, remains close (by 0.7 meV higher) to frequency of the corresponding mode of B isomer.

Although, it is not possible to make direct comparisons between theoretical results for the gas–phase clusters and experimental ones for the supported species, the fact that T structure, that is by 1.51 eV less stable than S, is observed in experiments requires a closer inspection of the potential energy landscape around and between these configurations. It might occur that due to high energy barriers for structural transformations between these three isomers the S structure becomes kinetically inaccessible. Therefore, the pathways and energy barriers for structural transitions are investigated. Main tool in these kind of investigations is the nudged elastic band (NEB) method and its climbing image (CINEB) version. They are both described in Sec.2.2.5.

Results of the CINEB calculations are also presented in Fig. 3.4. Minimal energy paths (MEP) for the transformations are found and the saddle points corresponding to the transition states located. It turns out that the MEP for transformation between the T and S structures passes through the intermediate B configuration. Therefore, two transition states exist along this pathway, one for the transition from T to B (TS1) and the second one from B to S (TS2).
**Figure 3.5**: Total and projected DOS of the three Ir$_4$ isomers (T, B, and S) are shown together with low lying and molecular orbitals (MO) around the Fermi energy. Orbitals are labeled according to the symmetry properties with respect to the corresponding symmetry group.
Energies of these two transition states relative to the initial configuration of the corresponding path (energy barrier for the process) are 0.15 and 0.16 eV for TS1 and TS2, respectively. These barriers are very low and after applying Eq.2.19 at room temperature, even using the lowest frequency vibrational modes of the two initial structures as the prefactors (∼10^{12} Hz), the rate constants for these two transformations are of the order of 10^{10} Hz. Of course, if the system has enough energy to overcome the first it will have enough energy to pass the second transition state, too. The conclusion is that, at room temperature, gas-phase Ir₄ clusters quickly adopt S geometry.

Atomic structures of the two transition states are also shown in Fig.3.4. In both cases one Ir–Ir bond of the corresponding initial configuration is considerably elongated. By inspecting electronic densities of TS1 and TS2 it can be concluded (on the basis of Bader analysis [74] for example) that those Ir–Ir bonds are broken. Therefore, the two transition states are located in the points (in the configurational space) where one Ir–Ir bond of the corresponding initial configuration is broken enabling transition in the desired direction, while the other bonds are not strengthen enough yet. Pushing the system from both transition states in any of the two directions which lower its energy would result either in reestablishing the broken bond (going back to the initial state) or strengthening other bonds and stabilizing the system in the corresponding final configuration. Very small energy needed for breaking one bond in T structure is a consequence of the symmetry breaking which enables more efficient hybridization of atomic orbitals. Similar arguments can be used in explaining the transition from B to S.

In Fig.3.5 total and DOS projected onto orthogonalized ( Löwdin [55]) atomic orbitals are shown for the three Ir₄ isomers (T, B and S). Spectra are broaden using Gaussians with width of ∼0.07 eV (0.005 Ry). Low energy as well as molecular orbitals (MO) near the Fermi energy are also presented. In all three cases MOs are formed either purely of atomic 5d orbitals or by dsp hybridization. Calculated Löwdin charges (integrated projected DOS) on each atom are distributed like 5d^{8.03}6s^{0.59}6p^{0.35} for T structure, 5d^{7.82}6s^{0.75}6p^{0.40} for B^8 and 5d^{7}6s^{0.86}6p^{0.48} for S isomer. The contribution of 6s and 6p AOs grows from T to S indicating bigger hybridization of atomic orbitals. The projected DOS shows that in the low energy part of all three electronic spectra the d–s hybridization dominates, while the contribution of 6p orbitals becomes appreciable at higher energies. This is due to the big energy difference between 5d, 6s and 6p orbitals in Ir atom (see Sec.3.1). Therefore, the following discussion is restricted only to the d–s type of hybridization.

Different level of d–s hybridization in these three structures is a consequence of different cluster symmetries. Namely, being totally symmetric with respect to 3D rotations, s orbitals can form exactly one strongly bonding MO in any atomic configuration (this is a general conclusion and does not depend on the symmetry or number of atoms). All other linear combinations of s orbitals from different atoms are either nonbonding or antibonding. Good example is purely s–bonded H₄ molecule whose properties are presented in Sec.3.4.1 (the following one). As it is shown in Fig.3.8 in the case of tetrahedral H₄ all MOs higher in energy have antibonding character. On the other hand, in B and S configurations, strongly antibonding combinations are pushed to higher energies and remain unoccupied. Therefore, the binding in these two structures is considerably stronger. However, if the number of electrons is such that only bonding MO is filled (0.5 electrons/atom), relative energies change dramatically (H^4^+ in Tab.3.2). Therefore, H₄ in T geometry favors population of s orbitals

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8For B structure average population of two symmetry nonequivalent Ir atoms is presented.
of 0.5 electrons/atom, while for the other two configurations it is more favorable (relative to T) to have bigger population of s orbitals.

The same reasoning can also be applied to Ir₄. In T geometry the binding would be enhanced if the d–like MOs hybridize only with the strongly bonding s–like linear combination. Since the strongly bonding s–like orbital is totally symmetric, only MOs of the same symmetry can hybridize and the only possibility is the bonding combination of dₓ² AO (see App.B). In B and S configurations number of totally symmetric linear combinations of d AOs is bigger (three A₁ and two A₁g linear combinations in B and S structures, respectively). Moreover, participation of other s–like MOs (nonbonding or weakly antibonding) could also act stabilizing.

To extract the influence of d orbitals on the atomic structure of Ir₄ (and more general X₄ molecules) in similar way is more difficult. The system whose properties are governed only by d orbitals does not exist and the analysis based purely on the symmetry and population of d orbitals (without the real system as an example) would not lead to unambiguous conclusions. Therefore, another way of investigating the influence of d–s hybridization on atomic structure in Ir₄ clusters is followed here. The idea is to modify iridium pseudopotential in such a way to decrease the level of d–s hybridization. This can be done by making the d component of iridium pseudopotential more attractive which would shift the energy of d electrons deeper in the spectrum. The following transformation is applied

\[ V_{l=2}^{\text{loc.}} = V_{\text{loc.}} + V_{\text{nonloc.}}^{l=2}, \]

\[ V_{\text{nonloc.}}^{l=2} \rightarrow \lambda V_{\text{nonloc.}}^{l=2}, \]

in which the nonlocal d component \( V_{\text{nonloc.}}^{l=2} \) of the pseudopotential is multiplied by a constant factor \( \lambda \). In order to avoid complications due to the augmentation charges, the normconserving pseudopotential (NCP) in the Troulier–Martins form [46] is used for this purpose. It is
Figure 3.7: Total and projected spin majority DOS of T and S Ir₄ isomers calculated using modified norm-conserving PP (see Eq.3.3) are shown. Numbers denote values of modifying parameter \( \lambda \), energy difference \( \epsilon_d - \epsilon_s \) (in eV) between the KS eigenvalues of \( d \) and \( s \) electronic states in Ir atom calculated in the spherical screening approximation, total cluster magnetic moment \( m \), Löwdin charges \( \mu \) of each iridium atom and the total energy difference (in eV) of the two isomers.
constructed with the fhi98PP pseudopotential program [47]. Effects of the modification (Eq.3.3) on the Ir pseudopotential are shown in Fig.3.6.

Three NCPPs are constructed in this way ($\lambda = 1.00, 1.02, 1.03$). Each of them yields different relative $d$–$s$ energies in iridium atom and hence, different level of $d$–$s$ hybridization in Ir$_4$ clusters as shown in Fig.3.7. For the sake of simplicity results for only $T$ and $S$ isomers are presented. Notice that the true iridium NCPP ($\lambda = 1.00$) gives different relative energy of $T$ and $S$ structures$^9$. However, calculated population of different atomic orbitals is very close to that obtained by USPP. Results obtained using modified NCPPs indicate that in Ir$_4$ the level of $d$–$s$ hybridization is responsible for the energy ordering of the three isomers. As the hybridization becomes less favorable and the population of $s$ orbitals approaches 0.5 electrons/atom in both configurations, situation changes and the energy preference of 1.7 eV towards $S$ isomer decreases to very low or even negative values.

3.4.1 Stability of $X_4$ isomers ($X=H$, Au, Pt, Ir, Rh, Co, Ru)

Subject of this section is the influence of $d$–$s$ hybridization on the relative stability of the three $X_4$ isomers ($T$, $B$ and $S$), where $X$ is a transition metal. The most stable tetramer geometry for given transition metal is not necessarily among these three structures i.e. for Au$_4$ the most stable isomer according to reference [76] is a planar structure, triangle with the fourth atom bonded solely to one the other three. Reason for the restriction to only these three isomers is their relevance for Ir$_4$ which is subject of this thesis. Since bonding in late transition metal clusters is very complicated (they are in general multiparameter systems with a large number of parameters), simpler situation of purely $s$–bonded H$_4$ molecule and the relative stability of different H$_4$ isomers could give a better insight into the influence of $s$ atomic orbitals on bonding in transition metal tetramers.

Results for the three H$_4$ isomers ($T$, $B$ and $S$), both neutral and doubly ionized$^{10}$, are presented in Fig.3.8 and Tab.3.2. In the neutral case, the least stable H$_4$ isomer is $T$ structure that is by 2.72 eV higher in energy than $S$ and by 4.36 eV than flat $B$ configuration with $D_{2h}$ symmetry (rhombus). The energy ordering of the three isomers can be understood from the symmetry analysis. Namely, atomic $s$ orbitals are totally symmetric with respect to 3D rotations and as such they can form only one strongly bonding molecular orbital whatever is the symmetry and the number of atoms in a cluster$^{11}$. This MO is the lowest in energy and doubly occupied (see Fig.3.8) in all three cases. All other linear combinations of $s$ orbitals have either weak or strong antibonding character.

In $T$ configuration all Ir–Ir distances are equivalent and, except the totally symmetric MO, only strong antibonding combinations of $s$ orbitals can be formed. In $B$ and $S$ structures due to the existence of different Ir–Ir coordination shells, linear combinations of $s$ orbitals with less pronounced antibonding character can be constructed. After populating these orbitals,

9 This is due to differences between USPP and NCPP in the description of atomic core in the absence of spherical screening approximation which is used for construction of both pseudopotentials. Small differences will be present in their Hartree potentials also outside the core. The NCPP atomic core is slightly more repulsive leading to somewhat different cluster relative energies.

10 Charged systems are simulated using the Makov–Payne [75] corrections to remove spurious electrostatic interaction between supercells.

11 In fact, they can form more than one if there are nonequivalent (by symmetry) atoms in a cluster. However, those orbitals will all be totally symmetric with respect to the cluster’s symmetry group and consequently they will hybridize forming only one strongly bonding MO spreading over all atoms in a cluster. This situation occurs in $B$ configuration.
Figure 3.8: Occupied parts of the spectra of three $H_4$ isomers ($T$, $B$ and $S$) are presented together with the spin majority (minority) KS eigenvalues (in eV). Spectra are aligned on the same Fermi energy. Molecular orbitals, labelled according to their symmetry properties, are also shown.

$T$ configuration becomes much more destabilized than the other two. On the other hand, if two electrons are removed from the clusters ($H_{2+}^4$), the remaining two electrons occupy only bonding MO in a singlet spin state and the relative energies of the three isomers are completely different. The $B$ structure remains the most stable, but now only by 0.34 eV lower in energy than $T$, while $S$ becomes the least stable structure that is by 0.82 eV higher than $T$. However, as soon as the electrons begin to fill antibonding MOs energies change in favor of the two planar structures. When the number of electrons is $\sim$0.6/atom $T$ and $S$ become degenerate and about 1.4 eV less stable than $B$ configuration. This result is useful in understanding the influence that $s$ atomic orbitals have on relative energies of transition metal tetramers.

Beside iridium, four other platinum group transition metals (Pt, Pd, Rh and Ru) plus Co and Au are examined (see Fig.3.9). Their tetramers are simulated with the same theoretical approach used for Ir clusters (Chapter 3). The energy differences between the three cluster geometries ($T$, $B$ and $S$) as well as the population of their AOs are investigated. Results obtained in this work agree reasonably well with already published theoretical results on tetramers of Au [76], Pt [76, 77], Pd [78, 79], Rh [80, 81], Ru [82] and Co [84, 85]. Similarly to iridium, differences exist when relative energies of different isomers are compared.

As expected, energy ordering of the three $Au_4$ isomers is the same as in $H_4$. Symmetries of the three isomers are the same as in $H_4$ as well as their magnetic moments. Qualitative similarities to hydrogen come from the fact that Au atoms has completely filled 5$d$ shell and only one electron occupying the 6$s$ AO. However, the energy differences between isomers are much smaller and there is also some depopulation of both 5$d$ and 6$s$ and population of 6$p$ orbitals. Moving one place left in the periodic table to $Pt_4$, isomer relative energies become very small. The $d$ shell in Pt atom has one electron less than Au and influence of $d$ orbitals on binding is much bigger. Magnetic moments of the three structures are different (2, 4 and $6 \mu_B$ for $T$, $B$ and $S$, respectively) and the symmetry of $B$ isomer is changed. Flat rhombic structure bends (angle $158.9^\circ$) and reduces its symmetry to $C_{2v}$. The population of atomic
The three Rh and it seems that d orbitals towards T by 0.79 eV lower in energy than Pt. Fig.3.7 where, for binding was only due to 5s orbitals in Pt.

Table 3.2: Properties of the three X₄ isomers (T, B and S). The clusters’ energies relative to T configuration E - Eₜ (in eV) and Löwdin population LP of the valence atomic orbitals are presented. B configuration is not a stable structure of Pd₄ and therefore no results are presented.

<table>
<thead>
<tr>
<th>system</th>
<th>T</th>
<th>E - Eₜ</th>
<th>LP</th>
<th>B</th>
<th>E - Eₜ</th>
<th>LP</th>
<th>S</th>
<th>E - Eₜ</th>
<th>LP</th>
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<tr>
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<td>0.77</td>
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<td>0.74</td>
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<tr>
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<td>0.35</td>
<td>0.12</td>
<td>0.73</td>
<td>0.40</td>
<td></td>
</tr>
<tr>
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<td>-0.08</td>
<td>0.71</td>
<td>0.56</td>
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<td>0.72</td>
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<td></td>
</tr>
<tr>
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<td>0.20</td>
<td>0.79</td>
<td>0.31</td>
<td>0.16</td>
<td></td>
</tr>
<tr>
<td>Pd₄</td>
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<td>-0.72</td>
<td>0.80</td>
<td>0.29</td>
<td>0.04</td>
<td>0.79</td>
<td>0.26</td>
<td></td>
</tr>
<tr>
<td>Rh₄</td>
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<td>0.79</td>
<td>0.26</td>
<td>0.72</td>
<td>0.15</td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td>Ru₄</td>
<td>0.00</td>
<td>0.57</td>
<td>-0.52</td>
<td>0.76</td>
<td>0.54</td>
<td>-0.40</td>
<td>0.67</td>
<td>0.54</td>
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<tr>
<td>Co₄</td>
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<td>0.67</td>
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</tbody>
</table>

orbitals in Pt₄ is such that it is not clear what type of geometry would be favored if the binding was only due to 5s orbitals. On the other hand, there is a clear preference of iridium d orbitals towards S structure. It follows from similar population of s orbitals in Ir₄ (as in Pt₄) and a strong energy preference to S isomer. This is also supported by the results from Fig.3.7 where, for λ = 1.02 or 1.03, s population favors T structure, but the two isomer become almost degenerate.

In Pd₄ the B configuration is not stable structure and it relaxes to S isomer which is by 0.79 eV lower in energy than T. The population of 5s in Pd₄ prefers T configuration and it seems that d orbitals do not influence a lot the energy ordering of T and S isomers. The three Rh₄ structures are almost degenerate. Since population of atomic s orbitals in

Figure 3.9: Elements whose tetramers are used for the analysis. Their symbols, atomic numbers and calculated valence populations are shown.
**Figure 3.10**: Total and projected spin majority DOS of T and S isomers of Co₄, Rh₄ and Ir₄ (iridium column in the periodic table) are shown. Numbers denote values of the total cluster magnetic moment m (in µB). Similarities of Rh₄ spectra with those Ir₄ calculated using modified PP for λ = 1.03 (Fig.3.7) are obvious.
Rh\textsubscript{4} clusters should also have weak preference to T configuration it can be concluded that d–bonding favors (also weakly) S structure. In Ru\textsubscript{4}, on the other hand, interaction of d AOs clearly favors S geometry. Population of s is such that the two isomers should be close in energy, but S is by 0.72 more stable. Situation in Co\textsubscript{4} is different than in other X\textsubscript{4} clusters presented here. The B isomer is the most stable followed by S and T that are by 0.12 and 0.52 eV less stable, respectively. Calculated Löwdin charges of s atomic orbitals should produce stronger preference towards B and S configurations. However, all three structures have high magnetic moments (10 \(\mu_B\)) and the gain in exchange energy also influences relative energies of the three isomers.

Since the relative stability of different isomers in X\textsubscript{4} clusters depends strongly on the level d–s hybridization it also depends on the approximations used in DFT calculations, especially on XC functional. In the cases where relative energies are small (i.e. Pt and Rh) differences in the predictions of the most stable isomer can be expected. However, strong preference of Ir\textsubscript{4} clusters towards S structure guarantees that results obtained for Ir\textsubscript{4} should be robust enough that reliable conclusions can be drown. It is shown in Sec.3.4 that, in order to alter relative stability between T and S isomers, one needs to reduce the level of d–s hybridization. On the other hand, the method that yields that level of d–s hybridization would produce poor description of Ir atom (completely wrong population of atomic orbitals) and probably would fail in reproducing properties of the bulk iridium.
Chapter 4

Influence of Adsorbates on the Stability of Ir\textsubscript{4} clusters

Understanding the interactions of (potential) catalysts with atomic species is the first step in understanding their interactions with more complicated molecular systems. Since atomic H as well as O are always present in real situations and since carbon is a key ingredient of all organic molecules, interaction with these three atomic species is probably among the most important to study. Moreover, the experimental procedure used to produce supported Ir\textsubscript{4} clusters involves quite complicated decarbonylation of Ir\textsubscript{4}(CO)\textsubscript{12} molecules. Hence, the experimentally produced tetrairidium clusters could contain atomic or molecular impurities. To investigate the effect of adsorbates on structural properties of Ir\textsubscript{4}, atomic H, C, and O as well as CO molecules are added to the clusters. Results for adsorption of these adsorbates on the gas-phase clusters are presented in the following two sections.

4.1 Adsorption of H, C or O atoms

The most favorable geometries of Ir\textsubscript{4}X (X = H, C and O) complexes, their total energies and binding energies of the species adsorbed on the three gas-phase Ir\textsubscript{4} isomers are shown in Fig.4.1 and Tab.4.1, respectively. While interaction of atomic H and H\textsubscript{2} with Ir\textsubscript{4} has already been studied [71, 72], adsorption of atomic O and C on Ir\textsubscript{4} is a novel result. Computational method was tested against the already published data for the adsorption of atomic H, C and O over Ir(111) surface and the agreement with theoretical predictions as well as with available experimental results is confirmed (check reference [86] and the references therein).

It is found during this work (reported in reference [69]) that adsorption of one carbon atom in the most stable, bridging configuration, alters relative stability predicted for clean gas–phase Ir\textsubscript{4} clusters. At T and B isomers binding energies of the C adatom are 8.24 and 8.00 eV, respectively. These values, by 1.82 and 1.58 eV larger than that found at the S cluster, show that big differences in the C chemisorption lead to the change in the isomer stability in favor of B configuration. The energy preference of B compare to T isomer is 0.34 eV. On the other hand, differences in binding energies of both H and O adatoms are much smaller than the relative energies of the three isomers. Therefore, the energy ordering of the three structures remains the same as for the clean gas–phase Ir\textsubscript{4}. All three types of adsorbates bind stronger to T and B isomers than to clean defect–free Ir(111) surface. This is also true for the adsorption of atomic H and O on S structure. However, C adatom binds by 0.29 eV
Figure 4.1: The most favorable adsorption geometries of atomic H, C and O on the gas–phase Ir₄ clusters in T, B, and S configurations. Numbers denote interatomic distances (in Å).

Table 4.1: Properties of the three gas-phase Ir₄ isomers (T, B and S) interacting with atomic H, C and O. Total energies ∆E (eV) with respect to that of T configuration, binding energies \( E_b \) (eV) of the adsorbed species and total magnetic moments \( m (\mu_B) \) of the molecules are shown. Results for the adsorption on Ir(111) surface are also shown for comparison.

<table>
<thead>
<tr>
<th></th>
<th>T</th>
<th>B</th>
<th>S</th>
<th>Ir(111)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \Delta E )</td>
<td>( E_b )</td>
<td>m</td>
<td>( \Delta E )</td>
<td>( E_b )</td>
</tr>
<tr>
<td>H</td>
<td>0.00</td>
<td>3.12</td>
<td>1.00</td>
<td>-0.82</td>
</tr>
<tr>
<td>C</td>
<td>0.00</td>
<td>8.24</td>
<td>2.00</td>
<td>-0.34</td>
</tr>
<tr>
<td>O</td>
<td>0.00</td>
<td>6.47</td>
<td>2.00</td>
<td>-0.62</td>
</tr>
</tbody>
</table>

1 Values taken from reference [86]

weaker to S isomer than to Ir(111) surface. This indicates that it is not only the reduced coordination compared to the atoms at surfaces that influences stronger reactivity of clusters, but also geometry and type of adsorbates play an important role.

The carbon adatom in the bridging site induces much larger distortions when adsorbed on the T and B than on the S isomer, as shown in Fig.4.1. The Ir–Ir bond bridged by carbon,
in the T and B structures is elongated to the values of 2.96 and 3.19 Å, respectively. The corresponding distance of 2.54 Å calculated for the S isomer remains close to the value in the pristine cluster, indicating that the Ir–Ir bond is not broken. By looking at the electronic density of the three structures in the plane defined by the two Ir atoms and the bridging carbon (Fig. 4.2(b)), this residual Ir–Ir bonding in the S structure can clearly be seen. Even though in all three cases there is no critical point in the density along the line that connects two Ir atoms and, therefore no bonding line in the sense of Bader analysis [74], the gradient of the density in the direction perpendicular to that line is much larger for the T and B isomers than for the S one. Hence, change in the electron density between the two Ir atoms of S isomer is smaller, reflecting the residual interaction. This is further supported by the shape of the C–2p projected density of states shown in the Fig. 4.2(a). Larger part of the C–2p spectral weight is transferred to lower energies in the cases of T and B isomers than in the S one.

For all three isomers weak interaction of carbon 2s and stronger interaction of carbon 2p orbitals with the clusters’ MOs exist. The biggest contribution to the carbon binding energy comes from the interaction of its 2p_z orbitals (z-axis is as in Fig. 4.2(b)) with molecular orbitals (MOs) of the clusters. These MOs of the T and B structures show strong antibonding character. They mostly belong to the unoccupied parts of the two spectra and after the interaction with the carbon adatom they become occupied which facilitates breaking of the corresponding Ir–Ir bonds. This is easier to see for T isomer where the unoccupied MO of the cluster that interacts with C–2p_z orbital corresponds to one of the low energy (with respect to E_F) orbitals with mixed T_1 and T_2 character (shown in Fig. 3.5). The corresponding MO of S structure is a combination of half–filled B_{1g} (also shown in Fig. 3.5) with a number of occupied MOs. It has much less pronounced antibonding character.

Therefore, the strongly antibonding MOs of the T and B clusters are transformed by the interaction to strongly bonding and the energy gain is much bigger for those two isomers than for the S one. In addition, interaction of the C–2p_z atomic orbital with the filled MOs of T isomer is also significant (second peak in the projected DOS). The MO formed by the interaction of C 2p_y AO and MOs of S structure, represented in the DOS by the second smaller peak, clearly shows the residual Ir–Ir interaction. Shapes of MOs of intermediate B structure after the interaction ressemble features of the other two isomers. Strong interaction of the C–2p_z state with the antibonding MO of the free cluster gives the main contribution to the binding energy of 8.00 eV, while the interaction of 2p_x and 2p_y atomic orbitals is clearly a transition between T and S structures.

On the other hand, both H and O adatoms are not capable of altering energy ordering of the three Ir_4 isomers. As already reported [71] hydrogen binds to T structure in the bridging configuration as well, but due to the symmetry reasons it is not able to interact with the same strongly antibonding MOs of the cluster. In fact, H binds stronger to the single Ir atom, but this geometry is not stable and transforms to one of the butterfly Ir_4H configurations\(^1\). Oxygen adatom, due to differences in its electronic configuration compared to carbon (see Fig. 4.3), binds preferentially to a single Ir atom in all three cases.

\(^1\)In this case H adatom interacts with strongly antibonding MOs of the cluster. Resulting structure is not the most stable B configuration with the H adatom and therefore, is not presented here.
Figure 4.2: a) Spin majority C–2p Löwdin projected DOS of the three Ir₄C molecules. b) Contour plot of the electronic density of the three Clr₄ molecules in the plane determined by the two Ir atoms interacting with a bridging C adatom. c) Kohn-Sham molecular orbitals of the three molecules corresponding to the peaks in the C–2p projected DOSs.
stronger to geometry with the carbon atom in contact with the cluster (Fig. 4.4). The CO binds slightly than at Ir(111) terraces for all three cluster geometries (see Tab. 4.2). The binding energies are always calculated using the isolated CO molecule as the reference. Compared to the adsorption at $r_e = 11.17$ eV \cite{87}, the adsorption of a CO molecule on Ir(111) surface \cite{86} which is also confirmed in this work.

### 4.2 Adsorption of a single CO molecule

Many possible configurations of a CO molecule interacting with the three Ir$_4$ isomers (T, B and S) are carefully inspected and the properties of the most stable ones are presented in Tab. 4.2 and Fig. 4.4. Both molecular and dissociative adsorption are studied. The CO binding energies are always calculated using the isolated CO molecule as the reference.

Accuracy of the method is tested against available experimental data. Calculated CO atomization energy $E_a = 10.97$ eV, bond length $r_e = 1.155$ Å and the frequency of its stretching mode $\hbar \omega = 272$ meV (2198 cm$^{-1}$) agree reasonably well with the experimental values ($E_a = 11.17$ eV \cite{87}, $r_e = 1.128$ Å and $\omega = 2169.8$ cm$^{-1}$ \cite{88}). Moreover, DFT predicts correct adsorption site of a CO molecule on Ir(111) surface \cite{86} which is also confirmed in this work.

Molecular CO displays a range of bonding modes with transition metal clusters. The CO ligand is usually observed as terminal, edge-bridging or face-capping \cite{15}. When adsorbed to Ir$_4$ clusters a single CO adsorbed in the molecular form binds stronger in the terminal geometry with the carbon atom in contact with the cluster (Fig. 4.4). The CO binds slightly stronger to B than to T isomer, while the difference in binding energies is bigger $\sim 0.5$ eV compared to the adsorption at S structure. The binding to Ir$_4$ is stronger by 0.55–1.1 eV than at Ir(111) terraces for all three cluster geometries (see Tab. 4.2).

Interaction with the molecular CO does not induce significant deformations of the three Ir$_4$ structures. Certainly, the symmetries of the clusters are broken and the distortions exist,
Figure 4.4: The most favorable adsorption of a CO molecule on the gas-phase T, B, and S Ir$_4$ clusters. For all three structures adsorption of the CO both in molecular and dissociated form is presented. Numbers denote interatomic distances (in Å).

Table 4.2: Energetics of the three gas-phase Ir$_4$ isomers (T, B and S) interacting with a single CO molecule both in molecular form (full) and dissociated (diss.). Total energies $\Delta E$ (eV) with respect to that of Ir$_4$CO in T configuration with a CO molecule in the dissociated form and binding energies $E_b$ of the CO molecules are presented. Values in parentheses denote binding energies of O adatom to the Ir$_4$C molecule. All energies are in eV.

<table>
<thead>
<tr>
<th></th>
<th>$\Delta E$</th>
<th>$E_b$</th>
<th>$\Delta E$</th>
<th>$E_b$</th>
<th>$\Delta E$</th>
<th>$E_b$</th>
<th>$E_b^{1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO full</td>
<td>0.83</td>
<td>3.16</td>
<td>0.19</td>
<td>3.23</td>
<td>-0.20</td>
<td>2.68</td>
<td>2.13</td>
</tr>
<tr>
<td>CO diss.</td>
<td>0.00</td>
<td>3.99(6.70)</td>
<td>0.03</td>
<td>3.38(6.34)</td>
<td>0.75</td>
<td>1.72(6.25)</td>
<td>0.32</td>
</tr>
</tbody>
</table>

1 Values taken from reference [86]

but they are less than 0.1 Å on average for all three structures. Both B and S structures remain magnetic upon the interaction ($4 \mu_B$ in both cases), while the ground state of T isomer with one terminal CO ligand is, as for the free cluster, nonmagnetic.

Dissociated, a single CO binds considerably stronger to T (by 0.83 eV) and only slightly stronger to B isomer (by 0.16 eV). On the other hand, dissociative adsorption on S structure is by 0.95 eV less favorable (qualitatively similar to the adsorption at Ir(111) surface) than adsorption in the molecular form. By inspecting adsorption geometries (Fig.4.4) this huge difference in the energetics of the CO dissociation reaction on the three isomers can be associated to a particularly strong binding of carbon adatom in the bridging configuration to both T and B isomers. The coadsorbed oxygen binds almost as strong as without carbon (check Tables 4.1 and 4.2). Upon the interaction with a single CO molecule S isomer with a single
molecular CO is the most stable Ir$_4$ structure, but now only by 0.20 and 0.23 eV more stable than T and B structures with the dissociated CO, respectively.

Interaction of a single CO molecule, adsorbed in the molecular form, with the transition metal clusters is similar to the interaction with surfaces of transition metals. It can be understood in terms of the charge donation from the CO molecule to the metal and backdonation to the molecule [15, 89]. At surfaces, it is proved that highest occupied $\sigma$ molecular orbitals (HOMO) and lowest unoccupied $\pi^*$ molecular orbital (LUMO) of the CO (see Fig.4.5) are mainly responsible for the interaction [90, 89]. Charge is donated from the HOMO to the metal surface and backdonated to the LUMO. Backdonation to the antibonding $\pi^*$ orbitals increases the C–O distance and facilitates $\sigma$ donation which in turn increases backdonation and so on (cooperative $\sigma$–$\pi^*$ bonding).

Mechanism of binding to the clusters is similar. However, deeper CO orbitals also play a role in binding. In fact, due to reduced symmetry $\sigma$ and $\pi$ orbitals are able to mix. Only, the lowest $\sigma$ and the LUMO of the CO molecule do not hybridize much with the others because of the big energy difference (Fig.4.5). In Fig.4.6 (upper panel) the total and DOS projected (within the Löwdin scheme [55]) on the orbitals of CO molecule are shown. The lowest energy $\sigma$ orbital of the CO molecule (mostly of the O–2s character) at $\sim$ -23 eV below the $E_F$ is not shown. Each of the three spectra can be divided in two regions (shorter dashed line in

**Figure 4.5:** Schematic plot of the relative positions of the spin majority KS energy levels of a CO molecule and energy ranges of the occupied parts of electronic spectra of the three Ir$_4$ isomers (T, B and S). The same description as for Fig.4.3 applies also here. The corresponding CO molecular orbitals are also displayed.
Figure 4.6: Spin majority total and CO projected DOS (using the Löwdin scheme [55]) of the three Ir₄CO geometries (T, B and S). Densities of states are shown for both CO adsorption in the molecular form (upper panel–FULL) and dissociative CO adsorption (lower panel–DISS.). Dashed lines (shorter) in the upper panel delimitate σ–π and π* parts of the spectra.

Fig.4.6), low energy region that is of the CO mixed σ and π character, and high energy region which is exclusively of the π* character. Energies of the bonding π* combinations lay closer to the Fermi energy while parts of the spectra around and above 2.5 eV are composed of the antibonding π* combinations with the orbitals of the three Ir₄ clusters. By integrating π* projections up to the E_F it is possible to estimate the amount of the backdonation from the three Ir₄ isomers to the CO molecule.

The biggest backdonation occurs for the CO molecule adsorbed to B isomer. In total, 0.82 electrons are transferred from the CO molecule to the cluster², 0.44 and 0.38 electrons in the spin majority and minority channels, respectively. Somewhat smaller amount of charge 0.36 electrons in each spin channel is backdonated to the CO molecule from T structure. In case of the CO adsorption to S isomer, filled bonding π* combinations in the spin majority channel are shifted closer to the Fermi energy and due to the magnetization of the cluster (4 µ_B) these orbitals in the spin minority channel remain mostly empty. Therefore, the backdonation is the smallest for the adsorption to S structure and amounts to 0.34 and 0.1 electrons divided in two spin channels. Differences in the amount of the backdonation correspond well to the ordering of the CO binding energies to the three isomers proving that the proposed binding mechanism is correct.

²Since the total charge transfer is very close to zero for CO adsorption to all three Ir₄ isomers, this number also measures the charge donation.
In the lower panel of Fig.4.6 the total and projected DOS of the three Ir$_4$ isomers with a single CO molecule in the dissociated form are shown. The O–2s atomic orbital that does not interact with any of the clusters’ electronic states is not presented. It is obvious from the three spectra that coadsorbed C and O adatoms interact with different molecular orbitals of the three clusters. Only minor changes are produced in the binding energies of the two atoms compared to the binding energies of the single C or single O atoms. This can be proved by summing C and O binding energies from the Tab.4.1 and subtracting the energy of 10.97 eV needed for the CO atomization. Resulting energies are 3.74 eV (T), 3.54 eV (B) and 1.67 eV (S) proving that C and O coadsorption are to a good approximation two independent processes$^3$.

### 4.3 CO Dissociation on Ir$_4$

![Figure 4.7](image)

**Figure 4.7:** Initial, final and transition state (TS) configurations for the CO dissociation on the three Ir$_4$ isomers (T, B and S) are shown. Numbers denote energies (in eV) of the initial and final configurations relative to that of T final configurations (as in Tab.4.2). Energy of the transition state is determined by the three energy barriers. Notice that the transition state is the same for CO dissociation on any of the three Ir$_4$ isomers.

Dissociation of carbon monoxide is an important chemical reaction for industrial applications (see Chapter 1). It is also one of the important reactions in the transition metal carbonyl cluster chemistry. Moreover, stability of T isomer found in experiments on supported Ir$_4$ clusters can be explained by assuming the presence of adsorbates, C adatoms in

$^3$In the case of T calculated total binding of the two adatoms is by $\sim$0.2 eV stronger and in the case of B is $\sim$0.2 weaker than these numbers. This is only a small portion of the total binding energy, but it makes the two isomers with the dissociated CO molecule almost degenerate.
particular (see Chapter 6). One possible source of C adatoms are carbonyl ligands present during the preparation procedure.

The CO dissociation over the surfaces of transition metals has already been studied by the means of density functional theory. It is found that the reaction requires as much as 1.11 eV on Fe(100) surface [91], while the same reaction takes to occur as much as $\sim$3.20 eV on Rh(111) terraces, $\sim$2.0 eV on the stepped Rh(210) surface [92] and about 2.70 eV on Co(0001) surface [93].

On the three Ir$_4$ clusters the CO dissociation reaction is studied using the CINEB approach (see Sec.2.2.5) starting from the configurations with a CO molecule adsorbed in the molecular form (initial configurations) and taking the structures after the dissociative CO adsorption as the final configurations. Many different pathways have been inspected for the CO dissociation on all three Ir$_4$ structures. One dimensional energy diagram for the reaction is shown in Fig.4.7. The reaction is exothermic on both $T$ and $B$ isomers while it is necessary to invest at least 0.95 eV for the reaction to occur on $S$ structure.

It turns out that the minimal energy paths (MEP) for the reaction on all three isomers pass through the same transition state. It is located in the butterfly–like configuration with the carbon atom bridging the two "wing" Ir atoms, while the oxygen is also placed in bridging position, but between the C and the wing-tip Ir atom of the opposite wing. This configuration enables strong interaction between C–2$p$ orbitals, as already explained in Sec.4.1, which on the other hand, weakens the C–O bond through the backdonation mechanism. However, the situation is rather peculiar since it is not clear which route will be taken when going from the transition state in either direction.

Since the transition state is the same, energy barriers for the reaction on all three isomers are determined by the energies of the initial configurations. The biggest activation energy 3.70 eV is for the reaction on $S$ structure while the activation requires at least 3.35 and 2.67 eV for the CO dissociation on $B$ and $T$ isomers, respectively. These barriers are quite big and the CO dissociation requires, according to Eq.2.19 with the prefactor of $10^{13}$ Hz (the CO stretching frequency), temperatures of over 1000$^\circ$ C. However, the barrier on $T$ isomer is by 0.49 eV lower than the CO binding energy indicating that the CO dissociation is more favorable process (provided there is enough energy) on $T$ isomer than the CO desorption. Influence of possible promoters for this reaction, i.e. other CO molecules present on the cluster that would interact with the oxygen atom and form CO$_2$, have not been studied yet.
Chapter 5

Ir₄ Clusters Supported by MgO(100) and TiO₂(110) Substrates

Adsorption of the metal particles over an oxide support is an important step in preparation of the real catalysts. Understanding the interaction of the metal clusters with an oxide surface at the atomic scale enables better control of the process of preparation and better insight into the properties of the adsorbed species.

The adsorption of different Ir₄ isomers on two widely used (both experimentally and theoretically) oxide surfaces, rocksalt MgO(100) and rutile TiO₂(110) is studied in this chapter. The influence of the support on relative stability, atomic and electronic structure of the adsorbed species is analyzed and the obtained results are compared to the available experimental data. Diffusion mechanisms on defect–free MgO(100) surface are also studied as well as the influence of point defects (neutral and charged oxygen vacancies) present on MgO(100) on the properties of the adsorbed Ir₄ clusters.

At the beginning the properties of the clean surfaces are presented. The theoretical model used for predictions is introduced through the analysis of the clean surfaces as well as the adsorption of atomic iridium on defect–free MgO(100) surface.

5.1 Bulk Oxides and Their Surfaces

5.1.1 Bulk MgO and the MgO(100) Surface

Bulk MgO is a material extensively studied both experimentally [95, 96] and theoretically [98, 99, 100, 101] due to its importance in technological applications. In this work bulk MgO is modelled using similar numerical setup as that already used for gas–phase Ir clusters (Chapter 3).

The ultrasoft pseudopotentials are employed with the same cutoff energies (25 and 140 Ry for wavefunctions and augmented charges, respectively). Brillouin zone integration is performed over 6×6×6 Monkhorst–Pack mesh [30]. Converged structural parameters, lattice constant of the rocksalt MgO a₀ = 4.26 Å and bulk modulus B = 150 GPa lay within standard DFT error bars (1–2 % for lattice constants and ~10 % for bulk modulus) compared to the experimental a₀ = 4.21 Å and B = 155 GPa [95]. On the other hand band–gap of the
Figure 5.1: (a) Total DOS of MgO(100) surface, defect–free as well as with neutral (F) and charged oxygen vacancies (F$^+$). Insets show structure of the corresponding DOS around Fermi energy. The arrow indicates states in the bend–gap formed upon creating the vacancy. (b) From left to right: relaxations of surface atoms close to the vacancies with $(x,y,z)$ displacement vectors for both F (black) and F$^+$ (red) centers; electronic orbital of the neutral vacancy appearing in the surface band gap; and the difference of the electronic densities $\Delta\rho = \rho_F - \rho_{F^+}$, isosurfaces are at $\pm 0.08 \text{ e/Å}^3$ and negative value is represented by the blue isosurface.

system is severely underestimated 4.49 against experimental 7.8 eV [97]. This is a well known feature of DFT methods since the Kohn–Sham eigenvalues do not formally correspond to excitation energies. Different values obtained theoretically are reported in literature and the value calculated in this work agrees well with that from reference [99], obtained within similar theoretical scheme.

Clean, defect–free MgO(100) surface is modelled with a supercell periodic slab approach. Large $4\sqrt{2}/2 \times 4\sqrt{2}/2 \times a_0$ surface unit cell is used. Sampling of the surface Brillouin–zone is restricted to the $\Gamma$–point only. Population of the KS electronic orbitals is done according to the Fermi distribution function with $k_B T = 0.1$ eV. Periodically repeated slabs are separated by the vacuum region chosen in such a way that the perpendicular translational period equals to the thickness of ten MgO(100) layers (21.3 Å). Further increase of the vacuum did not result in any appreciable change of the physical quantities of interest, i.e. binding energies of adsorbates. It is shown in Sec.5.2 that only two layer slab with the bottom layer fixed in the calculated bulk configuration is sufficient to obtain converged binding energy of atomic iridium (check also reference [103]). The asymmetric two layer slab is also good enough to reproduce structural properties of MgO(100) surface. Almost zero relaxation of the top layer plane and $\sim 2\%$ positive rumpling$^1$ is in line with results reported in reference [100] obtained with the slab containing seven layers.

On the other hand, two layers of MgO(100) are not sufficient to calculate accurately the band gap of the surface. Moreover, the asymmetric configuration with a small number of layers is not suitable for calculating band gaps at all since the states localized at the top of the valence band typically belong to the unrelaxed bottom layer. Therefore, calculations

$^1$Rumpling is defined as the ratio $(z_O - z_{Mg})/d_{bulk}$, where $z_O$ and $z_{Mg}$ are $z$ coordinates of the top layer O and Mg atoms while $d_{bulk}$ represents the bulk interlayer distance.
using the symmetric slab are performed and converged surface band gap of 3.62 eV is obtained using five MgO(100) layers (equivalent to 3 layer asymmetric slab). The "unconverged" gap of our model does not influence binding energies of iridium clusters since they interact only with states from the surface valence band (see following sections).

The valence bands of MgO(100) surface can be associated to O 2s and O 2p orbitals (structures around -17 and -5 eV in Fig.5.1(a)). On the other hand states from the conduction band do not follow "simple" charge–transfer model of ionic materials. The nature of the states in the conduction band of bulk MgO as well as the reduction of the band gap upon forming MgO(100) surface is elaborated in details in reference [101] and will not be discussed further here.

Neutral oxygen vacancy (F center) is produced by removing one of the surface O atoms. Since the oxygen removal induces additional relaxations of both topmost and layers below, the three layer slab was used in calculations. Again, only the bottom layer is kept fixed and the other two are fully relaxed. As shown in Fig.5.1 removal of the surface O atom leaves two electrons localized both in energy (state marked by arrow appearing in the band gap) and in the space between neighboring Mg ions. The description of the neutral vacancy goes in line with the results reported in reference [102].

Charged O vacancy (F$^+$ center) is simulated, by replacing the substrate Mg cation, belonging to the bottom layer, with a Na cation which has one less valence electron. The charge that is concentrated in the vacancy is reduced, by this procedure, from two to one electrons (see Fig.5.1(b)). In this way the problem of electrostatic interactions between supercells is circumvented. Similar model was successfully employed by Molina et al. [103]. In order to avoid relaxations that could be induced by the presence of the Na cation calculations are performed with the four layer slab with positions of the two bottom layers fixed while positions of all the other atoms are fully relaxed.

Electronic state localized in the band gap is shifted, upon charging, closer to the valence band in agreement with results reported in reference [102]. It has been proved that the exact position of Na in the bottom layer does not influence energetics of Ir adsorbates. The DOS structure at the top of the valence band around -1 eV (Fig.5.1) is an artifact of the model and it is due to the presence of sodium. However, those states are localized in the bottom layer (O atoms around the Na cation) and do not influence interaction with adsorbates.

### 5.1.2 Bulk TiO$_2$ and the TiO$_2$(110) Surface

Bulk rutile TiO$_2$ is modelled with the ultrasoft pseudopotentials and $6 \times 6 \times 8$ k-point grid. Converged structural parameters ($a = 4.65$ Å, $c = 2.98$ Å and $B = 234$ GPa) are in a good accord with the experimental ones ($a = 4.587$ Å, $c = 2.954$ Å and $B = 211$ GPa [104, 105]). Calculated band–gap of the material $E_g = 1.92$ eV is underestimated compared to the experimental gap $E_g^{exp} = 3.0$ eV [106] and is in line with other theoretical predictions [107, 108].

Surface calculations are performed with a four layer asymmetric slab (Fig.5.2). Only the bottom layer of the slab is kept fixed in the theoretical bulk configuration, while all the others are fully relaxed. Monoclinic supercell is used with the in–plane dimensions $\sqrt{2}a^2 + 4c^2$ and $4c$ whose unit vectors form the angle $\alpha = 47.85^\circ$. Dimension of the supercell in the perpendicular direction equals to $5a\sqrt{2}$ (ten surface layers $\approx 32.88$ Å). The Brilloiuin zone is sampled with $2 \times 2 \times 1$ Mohknorst–Pack mesh [30]. This geometry ensures at least 6 Å distance between the periodic images of the adsorbed Ir$_4$ clusters. Further increase of the
Figure 5.2: The four layer asymmetric slab used to model stoichiometric TiO$_2$(110) surface (left). Arrows denote two types of the surface Ti (five and six fold coordinated) and O (bridge and basal) atoms nonequivalent by symmetry. The rectangle encloses the fixed fourth layer of the slab. The surface unit cell used in calculations (right up) and the total DOS of the TiO$_2$ surface (right) together with the projections on the atomic orbitals of the surface atoms.

Description of the atomic structure of stoichiometric TiO$_2$(110) surface obtained with the explained setup agrees well with other theoretical results available in the literature. Discrepancies of the surface relaxations compared to experiments are of the order of several $10^{-2}$ Å (reference [106] and the references therein). The asymmetric slab approach provides good convergence of the surface structural properties. In this way the even–odd oscillations of the surface relaxations with respect to the slab thickness are avoided (for details check reference [109]). On the other hand, as already discussed in previous section, the asymmetric slabs are not suitable for extracting unambiguously the electronic structure of the surface close to the valence band maximum as well as the conduction band minimum\(^2\). However, main features of the surface DOS are reproduced well. In Fig.5.2 the valence and part of the conduction band of the slab are shown. In agreement with the studies available in literature [110, 111] the surface valence band is mainly of the O 2$p$ character while the Ti 3$d$ character dominates the lower part of the conduction band. The $p$–$d$ mixing in both parts of the DOS is not negligible.

\(^2\)As it is also discussed in previous section this disadvantage of the asymmetric slab approach is not a problem for studying adsorption since adsorbates always interact with the states from the relaxed slab surface. It is just inconvenient for extracting the electronic properties of the surface that can be compared with experiments i.e. photoemission spectroscopy.
5.2 Adsorption of Atomic Iridium on MgO(100)

At variance to adsorption of Ir\(_4\) clusters on TiO\(_2\)(110), their adsorption on MgO(100) surface can be understood from the features of the adsorption of atomic iridium. Therefore properties and adsorption mechanism of the Ir atom on MgO(100) are presented before those of the tetramers.

Adsorption of platinum group transition metal atoms over the (100) surface of MgO has already been extensively studied with both all-electron and pseudopotential DFT methods. For atomic Fe [112], Ni [113, 116, 117], Pd [115, 116, 117], Pt [114, 115, 116, 117] and Ir [117] supported by MgO(100) a clear preference to the adsorption atop the surface oxygen is found. However, binding energies obtained by different methods may differ considerably\(^3\). These differences can be associated to different reference atom used to calculate binding energies.

It is confirmed in this work that Ir atom binds in the same way as the other transition metals. Converged values for the binding energy, Ir–O bond length and the induced dipole moment are achieved using the setup equivalent to that explained in Sec.5.1.1. As shown in Tab.5.1 the asymmetric two–layer slab, with the adsorbate only on one side, and the Γ–point BZ sampling with 4\(a\)\(_{\text{MgO}}\)×4\(a\)\(_{\text{MgO}}\) surface unit cell\(^4\) is sufficient for achieving the accuracy of 0.03 eV and 0.01 Å in binding energy and Ir–O distance, respectively. Similar numerical setup is used by other authors for the adsorption over MgO(100) surface (see for example reference [114]). A dipole correction method (Sec.2.2.3) is used to cancel the artificial electric field produced by the charge transfer and the asymmetry of the system. The surface dipole can be calculated as the difference

\[
P_{\text{Ir/MgO}(100)} - P_{\text{MgO}(100)}
\]  

between the total dipole moments of the slab with and without the adsorbate. Due to the geometry of the model the system can develop surface charges (on both surfaces of the slab) that may not be the same in the presence of adsorbates and hence does not cancel exactly when subtracting the dipole moments. To test validity of Eq.(5.1), calculations in the symmetric slab configuration with the adsorbates on both sides of the slab, which does not suffer from the abovementioned problems, are carried out. In this case surface dipole is proportional to the difference of the electrostatic potentials of the slab with and without adsorates (aligned at the middle layer) far away from the surface. The symmetric slab approach yields essentially the same results (Tab.5.1).

Optimized geometry of the Ir atom adsorbed atop the surface oxygen is shown in Fig.5.3. Calculated binding energy of 2.33 eV is by 1.66 eV higher than at the Mg site. The surface hollow site is a saddle point for diffusion between two O atoms and lays 0.74 eV above the atop oxygen adsorption energy. The Ir binding energy of 1.39 eV is reported in reference [117], calculated within the embedded cluster all–electron approach. As already said, this difference can be associated to the different reference Ir atom. On the other hand, at the most stable oxygen site, calculated Ir–O distance 1.91 Å is in reasonable accord with the value of 2.00 Å from the same reference.

\(^3\)For example binding energy of Pt atom 1.54 eV is calculated with the periodic supercell approach [114] while the value of 2.39 eV is obtained when the all-electron embedded cluster method is used [117].

\(^4\)Strictly speaking, the smaller cell would also be sufficient to simulate adsorption of Ir atom. Bigger one is used here only because of consistency with the simulations of Ir\(_4\) adsorption.
Table 5.1: Convergence tests of the binding energy $E_b$, Ir-O bond length and the $z$-component of the induced dipole moment $p_z$ of atomic Ir adsorbed on MgO(100) as a function of number of layers $n$ and a number of the special, Monkhorst–Pack (MP) $k$-points [30] used in simulations.

<table>
<thead>
<tr>
<th>$n$</th>
<th>MP grid</th>
<th>$E_b$ [eV]</th>
<th>Ir–O [Å]</th>
<th>$p_z$ [D]</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>$\Gamma$</td>
<td>2.33</td>
<td>1.906</td>
<td>-0.88</td>
</tr>
<tr>
<td>2 $\times$ 2</td>
<td>2.30</td>
<td>1.910</td>
<td>-0.92</td>
<td></td>
</tr>
<tr>
<td>3 $\times$ 3</td>
<td>2.30</td>
<td>1.910</td>
<td>-0.89</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>$\Gamma$</td>
<td>2.35</td>
<td>1.898</td>
<td>-0.87</td>
</tr>
<tr>
<td>3</td>
<td>$\Gamma$</td>
<td>2.32</td>
<td>1.909</td>
<td>-0.87</td>
</tr>
<tr>
<td>5</td>
<td>$\Gamma$</td>
<td>2.34</td>
<td>1.898</td>
<td>-0.90</td>
</tr>
</tbody>
</table>

The surface distorts upon adsorption in such a way that the Ir–bonded O atom is pushed down for 0.02 Å while the neighboring Mg atoms pop up from the surface for about 0.05 Å. These local distortions are not enough to change sign of the surface rumpling as in the case of Pt adsorption [114].

To rationalize binding of atomic Ir to MgO(100) surface analysis of the electronic structure is performed. Both charge transfer and projected density of states (PDOS) onto atomic orbitals are studied and the results are presented in Fig.5.3 and Fig.5.4. Charge transfer is estimated using the in-plane integrated electronic density difference along $z$ axis

$$\Delta \rho(z) = \int \int \Delta \rho(x,y,z) \, dx \, dy = \int \int \left[ \rho_{\text{Ir/MgO}} - \rho_{\text{Ir}} - \rho_{\text{MgO}} \right] \, dx \, dy,$$

(5.2)

where $\rho_{\text{Ir/MgO}}$, $\rho_{\text{Ir}}$ and $\rho_{\text{MgO}}$ are electron densities of the full system and the two non-interacting subsystems, Ir atom and MgO(100) surface (in their frozen interacting configurations), respectively. It is clear from Fig.5.3(b), in which only the region where $\Delta \rho$ is significantly different from zero is displayed, that the charge transfer occurs from the surface to the adsorbate. It can be estimated by integrating $\Delta \rho(z)$ up to the mid-plane between the two subsystems (red dashed line in Fig.5.3(b)) to 0.31 e resulting in the surface dipole of 0.88 D directed from the adsorbate to the surface. If a simple point charge model is constructed with the negative charge (-0.31 e) positioned in the center of $\Delta \rho(z)$ above the mid-plane and positive charge positioned in the center of $\Delta \rho(z)$ below, their interaction energy would be $\sim$1.3 eV (the distance between charges is about 1.05 Å). Therefore, more than half of the binding energy is due to electrostatics while the other part is a consequence of the energy gain due to the orbital overlap.

After the analysis of the shape of $\Delta \rho$ it is evident that density is depleted mostly from the oxygen out of plane electronic orbitals as well as the metal states pointing towards the oxygen whereas the Ir states parallel to the surface accumulate electrons. This is a consequence of the interaction of iridium atomic orbitals with the valence $p$–band of the surface. The $C_{4v}$ symmetry of the system (surface plus the adsorbate) allows the interaction only between states with the same symmetry properties. Thus only the iridium $s$, $d_{xz}$, $d_{yz}$ and $d_{yz}$ orbitals can interact with the filled $2p$ states of the Ir–bonded oxygen atom. In Fig.5.4 total and PDOS of the MgO surface prior and the full system after the interaction are presented,
Figure 5.3: (a) Top view of the Ir adsorption geometry over MgO(100) surface. Full 4a_{MgO}×4a_{MgO} simulation cell is shown. (b) Side view of the adsorbed Ir atom, Ir–bonded O and bottom layer Mg atoms as well as the electron density difference $\Delta \rho$ (Eq.5.2). Bright isosurface indicates a density gain while blue one indicates density depletion. The isosurfaces are at $\pm 0.035 \ e/\text{Å}^3$. The curve to the right is the in–plane integrated density difference $\Delta \rho_z$ whose integral below (above) the mid–plane, represented by the red dashed line, estimates total charge transfer $\Delta q$. Numbers in Å denote interatomic distances. Mirror plane $\sigma_v$ and the $C_4$ rotations belonging to the $C_{4v}$ symmetry group of the system are also presented.

respectively. Energy range which, for the oxide, is dominated by $2p$ and $2s$ states is displayed. MgO(100) surface without the adsorbate is “frozen” in the same atomic configuration as after the interaction. The PDOS clearly shows how the Ir $5d$ and $6s$ orbitals interact with the oxygen $2p$ ones. Deformation of the surface, that costs 0.33 eV, localizes $p$ states of the Ir–bonded oxygen energetically at the top of the valence band. Most of their spectral weight is pushed by the interaction to the bottom and smaller portion to the top of the $p$ band. The covalent type Ir–O bond is established. Bonding $\sigma$ orbital, formed by the interaction of Ir $5d_{z^2}$−$6s$ hybrids with O–$2p_z$ states, lays at the bottom of the new valence band. Next to it in the low energy part of the valence band the doubly degenerate $\pi$ bonding orbitals are positioned. They are formed by the interaction of iridium $5d_{xz}$ and $5d_{yz}$ orbitals with $2p_x$ and $2p_y$ from the interacting O atom. Corresponding antibonding orbitals are pushed to the top of the valence band or above the Fermi energy. The weakly interacting $5d_{x^2−y^2}$ (with neighboring O atoms) and non-interacting $5d_{xy}$ iridium orbitals are positioned in the vicinity of the Fermi energy, slightly below the antibonding $\sigma^*$ and $\pi^*$ combinations. Interaction is the strongest between oxygen $2p_z$ orbital and iridium $5d_{z^2}$−$6s$ hybrids. Thus, the charge transfer is mainly due to: (i) the antibonding $\sigma^*$ state in the spin majority channel, pushed above the Fermi energy causes the electron transfer to the spin minority Ir orbitals parallel to the surface; and (ii) the Ir $5d_{xz}$ and $5d_{yz}$ orbitals in the spin minority channel that were empty prior the interaction become partly occupied because of the interaction with oxygen $2p_x$ and $2p_y$. 
5.3 Adsorption of Ir$_4$ Clusters on MgO(100)

Interaction of the Ir$_4$ clusters with clean and defect–free MgO(100) surface is studied by placing the clusters, in their equilibrium gas–phase configurations, at the surface and performing the full geometry optimization. Many different adsorption geometries were carefully inspected for each of the three Ir$_4$ isomers, tetrahedral (T), butterfly (B) and square (S). Only results for the configurations with the lowest total energies are presented here. Details of the adsorption geometries as well as the relative energies of the three clusters are depicted in Fig. 5.5.

More quantitative description of the adsorption geometries and energetics is given in Tab. 5.2. Relative and binding energies, details of the electronic structure as well as the Ir coordination numbers and average distances for all iridium coordination shells up to 3.5 Å are given. As a coordination shell a group of distances to a specific type of neighbors spanned over the region not wider than 0.2 Å is considered$^5$. Available experimental data from EXAFS measurements (references [118, 119]) are also shown for comparison. Calculated distances are presented for only those shells with coordination numbers larger than 0.25 and the standard deviation (from the corresponding average distance) smaller than 0.1 Å. This is due to the fact that static disorder (measured by the standard deviation), together with the thermal disorder and inelastic effects, reduces the amplitude of the associated electron backscattering wave in EXAFS measurements (see Appendix A).

Similarly to the adsorption of Ir atom all three Ir$_4$ isomers bind to the surface oxygen. T isomer forms three contact points to three oxygen atoms with a total cluster binding energy of 3.83 eV. One side of the basal Ir triangle is considerably elongated (∼3.3 Å). The resulting configuration is similar to a butterfly geometry with two different Ir–Ir coordination shells.

\(^{5}\text{Value taken from the Ref. [133]}\)
The support relaxes as well increasing its total energy for 0.65 eV. Surface atoms neighboring to those which interact with the clusters move in–plane as well as out–of–plane for about several hundredths of an angstrom from their equilibrium positions at MgO(100). The interacting oxygen atoms and the Mg atom positioned below the cluster are displaced somewhat more, as shown in Fig.5.6.

Both B and S structures bind to defect–free MgO(100) surface with similar total binding energies of 3.35 and 3.33 eV, respectively. Hence, after the adsorption S isomer remains the most stable form of Ir₄. Both B and S interact with only two surface O atoms. Configurations where two isomers interact with more than two surface O atoms are not favorable due to the large mismatch between the surface O–O distance (3.01 Å) and the average bond lengths of the two isomers (2.35 and 2.31 Å for B and S, respectively). Interatomic distances in the two clusters are barely altered due to the interaction with the surface. Average Ir–Ir distances of S (B) isomer are 2.31 Å (2.38 Å) and 3.27 Å (3.68 Å) for the two Ir–Ir coordination shells.

Since T isomer distorts noticeably upon adsorption, the term tetrahedron (or the T symbol), when speaking about supported clusters, refers to the lowest total energy configuration formed after the relaxation of the gas–phase T on the MgO surface.
Table 5.2: Calculated properties of T, B and S Ir₄ isomers adsorbed on MgO(100) surface. Total energy $\Delta E$ relative to that of T configuration, cluster binding energy $E_b$, total magnetic moment $M$, charge transfer $\Delta q$ (in electrons), $z$-component of the induced dipole moment $p_z$, coordination number $N$, average distance $R$ of the corresponding coordination shell for all shells up to 3.5 Å and all types of Ir neighbors (see the text for details). Available experimental data are also presented.

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<tbody>
<tr>
<td>T</td>
<td>0.00</td>
<td>3.83</td>
<td>3.46</td>
<td>-0.50</td>
<td>-4.31</td>
<td>2.5</td>
<td>2.43</td>
<td>0.75</td>
<td>2.15</td>
<td>0.5</td>
<td>2.72</td>
<td>0.5</td>
<td>3.00</td>
<td>1.0</td>
<td>3.34</td>
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<td></td>
<td></td>
<td>0.5</td>
<td>3.29</td>
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<td></td>
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<tr>
<td>B</td>
<td>-0.14</td>
<td>3.35</td>
<td>2.04</td>
<td>-0.34</td>
<td>-3.60</td>
<td>2.5</td>
<td>2.38</td>
<td>0.5</td>
<td>2.06</td>
<td>1.0</td>
<td>2.88</td>
<td>0.5</td>
<td>3.00</td>
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<td></td>
<td></td>
<td>0.5</td>
<td>3.68</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>S</td>
<td>-1.06</td>
<td>3.33</td>
<td>2.31</td>
<td>-0.38</td>
<td>-3.88</td>
<td>2.0</td>
<td>2.31</td>
<td>0.5</td>
<td>2.07</td>
<td>1.0</td>
<td>2.88</td>
<td>1.0</td>
<td>3.27</td>
<td>1.0</td>
<td>3.19</td>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td>1.0</td>
<td>3.27</td>
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<tr>
<td>Exp. Ref.[118]</td>
<td>2.6–3.1</td>
<td>2.62–2.73</td>
<td>0.6–1.1</td>
<td>2.07–2.16</td>
<td>2.5–3.0</td>
<td>2.59–2.73</td>
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<tr>
<td>Exp. Ref.[119]</td>
<td>2.9</td>
<td>2.62</td>
<td>1.1</td>
<td>2.05</td>
<td>0.6</td>
<td>2.45</td>
<td>0.5</td>
<td>1.91</td>
<td></td>
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</table>
Table 5.3: Deformation energies $E_d = E_{deformed} - E_{ground \ state}$ (in eV) for the adsorption of Ir$_4$ clusters over MgO(100). Values are for the three Ir$_4$ isomers ($E_1^d$) as well as for MgO(100) support ($E_2^d$). $E_{int}$ is purely electronic interaction energy (in eV) that can be calculated as the total cluster binding energy (Tab.5.2) plus the total deformation energy $E_1^d + E_2^d$.

<table>
<thead>
<tr>
<th></th>
<th>$E_1^d$(Ir$_4$)</th>
<th>$E_2^d$(support)</th>
<th>$E_1^d + E_2^d$</th>
<th>$E_{int}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>-0.28</td>
<td>0.65</td>
<td>0.37</td>
<td>4.19</td>
</tr>
<tr>
<td>B</td>
<td>0.10</td>
<td>0.63</td>
<td>0.73</td>
<td>4.09</td>
</tr>
<tr>
<td>S</td>
<td>0.03</td>
<td>0.52</td>
<td>0.55</td>
<td>3.88</td>
</tr>
</tbody>
</table>

Figure 5.6: Deformations of MgO(100) surface after the adsorption of the three Ir$_4$ isomers (T, B and S). Arrows denote displacements that are shown only for the interacting O as well as the neighboring Mg atoms. Numbers in parentheses are the $(x, y, z)$ displacement vectors (in Å). They are given only for those atoms nonequivalent by symmetry.

respectively. Angle between the wings of B structure changes from the gas–phase 138.13° to 133.13°. This slight closing of B structure increases its energy by 0.1 eV. Distortions of the support, especially neighboring cations, are similar in both cases (Fig.5.6). Differences in deformation energies of the support come from different displacements of the interacting O atoms due to different symmetries of the two systems. Both clusters are magnetic upon adsorption, but while the magnetic moment of B is unchanged (2.04 $\mu_B$) with respect to its gas–phase value, moment of S structure is lowered to the value of 2.31 $\mu_B$.

As in the case of free clusters, CINEB calculations showed that S isomer, the most stable form of Ir$_4$/MgO(100), is not kinetically inaccessible starting from the other two. The numerical setup used for calculations is explained in Sec.2.2.5. The minimal energy path (MEP) for the transformation from T to S structure passes, similarly to the gas–phase MEP (Sec.3.4), through B configuration (see Fig.5.5). Activation energy for the transformation from T to B is calculated to be 0.16 eV while from B to S 0.15 eV is needed. However, main features of the transformations between T and B structures are different. One Ir–Ir bond of T structure between the two basal Ir atoms is already broken. Therefore, to transform into B configuration only one Ir–O bond is needed to brake. The activation of that Ir–O bond needs similar energy as one Ir–Ir bond in the gas–phase. The fact that very low energy is required for activation of the Ir–O bond can be attributed to the stronger bonding (binding energy per Ir–O bond is 1.28, 1.68 and 1.67 eV for T, B and S, respectively) of both B and S isomers to the surface. Activation energy of the Ir–O bond of T structure is to a good extent
compensated by the energy gain due to the strengthening of other two and the energy of the transition state is low. For further transformation along the MEP from B towards S isomer there is only one Ir–Ir bond that breaks, as in the gas–phase, which requires similar activation energy. Finally, the same conclusion can be drawn that at room temperature, both T and B configurations supported by defect–free MgO(100) remain metastable structures with a short lifetime.

Even though the gas–phase energy ordering of the three isomers remains the same upon adsorption, relative stability is considerably changed. As a consequence of different binding, relative energies of S and B isomers, calculated with respect to T one, are lower for 0.42 and 0.44 eV, respectively. If the interaction with the surface is divided in two contributions (i) deformations and (ii) purely electronic interaction; it can be shown that a good part of the reduction in relative energies comes from the different deformation energies of the three clusters. When differences between adsorption of T and B isomers are analyzed it can be seen that only a small portion (0.1 eV) of the change in their relative energy is due to purely electronic effects (Tab.5.3). This results from the unusual, negative deformation energy of T isomer.

From the analysis of the electronic structure of the three isomers (Fig.5.7) it follows that main features of the electronic interaction between the three isomers and MgO(100) surface are similar to those of Ir atom. Clusters interact mainly with the states from the surface valence band and the charge is transferred from the surface to the adsorbates. The charge transfer is estimated in the same way as explained in previous section. Due to the bigger number of interacting O atoms T isomer interacts with bigger number of O–2p orbitals (surface under the red curves on the left panel of Fig.5.7). This results in bigger charge transfer as well as larger surface dipole moment (Tab.5.2). However the electronic interaction is only by 0.1 and 0.27 eV stronger than that of B and S isomers, respectively. This can be attributed to
more favorable overlap of the molecular orbitals of \( B \) and \( S \) isomers with the states on the surface. Somewhat stronger interaction of \( B \) compared to \( S \) is mainly due to the different cluster symmetry and differences of the molecular orbitals near the Fermi energy.

From Tab.5.2 it is clear that geometry of none of the MgO(100) supported Ir\(_4\) isomers corresponds well to experiment. Intriguing is the fact that geometry of \( T \) configuration, predicted theoretically to be the least stable, is the closest to the experimental situation. It has the biggest Ir–Ir first shell coordination number and the longest Ir–Ir bond length. However, contributions coming from the second Ir–Ir shell are not measured and the calculated Ir–Ir distances are noticeably shorter than those reported from experiments.

When comparing theoretical results with those obtained by EXAFS it is necessary to take into account that the errors of EXAFS are not small. They are estimated to \( \sim 10\% \) for the coordination numbers and about 0.02 Å for the distances coming from the metal–metal contributions. For low–Z scatterers (carbon, oxygen,...) these errors can be even bigger [124]. Differences from the calculated results may also arise from temperature effects since the EXAFS spectra are recorded at room temperature. Finally, there are also differences between different measurements that are dependent on the preparation procedure of the substrate.

In Ref.[118] each coordination shell is characterized by a range of values depending on the calcination temperature of the substrate. No contribution from the Ir–Mg and Ir–C shells is reported. In a more recent publication [119] the same authors measure different longer Ir–O contribution as well as the contributions coming from the Ir–Mg and Ir–C shells. The latter could come either from the non–desorbed CO ligands (partial decarbonylation)\(^7\) or from the standalone C adatoms. As it is already shown for the gas–phase clusters (references [69, 70] and Chapter 4) and will be confirmed also for the MgO supported species (Sec.6.1 and Ref.[69, 70]) one C adatom is sufficient to stabilize \( T \) configuration.

### 5.4 Diffusion of Ir\(_4\) Clusters on MgO(100) Surface

Diffusion of supported atoms or small clusters is an important issue in understanding the processes related to their deposition. Low barriers for diffusion lead to aggregation of the supported species and island formation (sintering). In this case defects at surfaces act as the nucleation centers due to their ability to trap adsorbates. It has been found recently that small palladium clusters are highly mobile on MgO(100) surface [120, 121]. Moreover, by varying external parameters during the preparation procedure of MgO(100) supported Ir\(_4\) clusters signs of aggregation have been discovered when the supported clusters were treated in He gas at temperatures above 400\(^\circ\) C [123].

According to the results presented for clean Ir\(_4\) clusters adsorbed over defect–free MgO(100) surface (previous section) it makes sense to study diffusion only of \( S \) isomer. It is found in this work that any adsorption configuration of the other two isomers, different from those adsorption geometries shown in Fig.5.5 lays much higher in energy than the transition states for the structural transformations to \( S \) structure. Therefore, for \( T \) and \( B \) isomers it is energetically more favorable to transform to \( S \) configuration than anything else.

On the other hand, for \( S \) to diffuse over MgO(100) surface several possibilities exist: (i) to move, keeping the same plane normal to the surface, to the configuration with only one

\(^7\)The same Ir–C and long Ir–O coordination numbers go in line with this assumption, but in that case the CO bond would be considerably elongated (\( \sim 1.4 \) Å) which is not confirmed by our calculations and/or other measurements.
Ir in contact with the surface oxygen and then to continue "rolling" in the same direction to the next second O atom; (ii) to transform to geometry parallel to the surface and than to straighten itself up to the same configuration as the starting one, but now in contact with two different surface O atoms; (iii) to transform to the configuration with S structure also straighten up interacting with two surface oxygen atoms, but with one Mg atom below the cluster and then to continue to the configuration equivalent to the starting one with one of the interacting surface O atoms different from those at the beginning (cluster walk); and finally (iv) to transform from the same starting to the equivalent final configuration as under (iii) but not preserving the square shape and passing through B, T and then again B configuration. It turns out that the energies of the intermediate configurations along the pathways (i) and (ii) are by 1.42 and 2.39 eV higher than that of the most stable S adsorption geometry, respectively. On the other hand, the intermediate configuration for the transformation (iii) (S isomer positioned above one surface Mg atom interacting with two oxygen atoms) lays only 0.86 eV above the most stable S adsorption configuration. Calculated energy barrier for the structural transformation along the pathway (iii) is 1.32 eV. Transition state lays in the configuration with one Ir atom in contact with the surface and the other in–between the two potentially interacting O atoms. Calculated activation energy for the diffusion channel (iv) is
1.25 eV\(^8\) and is simply the energy needed to go backwards from \(S\) to \(B\) and \(T\) configurations\(^9\).

In Fig.5.8 energies of the stable configurations as well as of the transition states for the transformations \((iii)\) and \((iv)\) are shown together with the corresponding geometries. All energies are presented relative to the most stable \(S\) adsorption configuration (initial and final). According to the calculated energy barriers these two diffusion channels are relevant for the \(Ir_4\) clusters supported by MgO(100). The temperatures that could activate the diffusion can be estimated using Eq.2.19. The prefactor (attempt frequency) in Eq.2.19 is of the order of \(10^{12} - 10^{13}\) Hz since the highest vibrational frequencies of Ir clusters are \(\sim 30\) meV (\(\sim 10^{12}\) Hz) and it is known that vibrations of MgO(100) surface lay below 88 meV (\(\sim 10^{13}\) Hz) \([122]\). Both barriers of 1.25 and 1.36 eV are not small and for the temperature of 300\(^\circ\)C the rate constants for the two processes are in the range 1–10 Hz. With these rate constants the average half–life for diffusion of \(S\) configuration is \(\sim 0.07–0.7\) s. This is not much, since the probability of returning to the same starting configuration after two cycles is the same as displacing further, which reduces considerably the number of clusters that pass a certain distance. If the temperature increases to 400\(^\circ\)C rate constants become two orders of magnitude higher \(100-1000\) Hz which significantly changes the timescale for the diffusion (half–life is 0.0007–0.007 s). Increasing further the temperature from 400\(^\circ\)C the half–life decreases rapidly. This result is in a good agreement with experimental findings \([123]\) and confirms that \(Ir_4\) clusters prepared at and below 300\(^\circ\)C should be well separated, while the aggregation can be expected at higher temperatures.

5.5 Adsorption of \(Ir_4\) Clusters on MgO(100) Surface With a Neutral and Positively Charged Oxygen Vacancy

As in previous section, properties of only the most stable adsorption configurations for each of the three \(Ir_4\) isomers on MgO(100) surface with neutral (F) as well as with positively charged oxygen vacancy (\(F^+\) center) are presented here (Fig.5.9). Modelling of the defected surface is already explained in Sec.5.1.1.

The three isomers bind stronger to the neutral vacancy site than on defect–free MgO(100) surface. Stronger binding of \(d\)-metals to the O vacancy than at defect–free MgO(100) surface is a well known feature \([125]\) that is attributed to the charge transfer from the support to the adsorbate. Again \(T\), the least stable gas–phase structure, binds with the highest binding energy of 6.32 eV followed by \(B\) and \(S\) with binding energies of 5.21 and 5.06 eV, respectively (Tab.5.4). Increase in binding energies is the biggest for \(T\) isomer (2.49 eV), while this shift amounts to 1.86 and 1.73 eV for \(B\) and \(S\) structures, respectively. The Energy ordering is now different. The most stable isomer is still \(S\) with the total energy for only 0.25 eV lower than \(T\) while \(B\) isomer becomes the least stable structure with the energy for 0.53 eV higher than \(T\) configuration.

The adsorption geometries are similar to those at the defect-free surface and can be obtained by removing one of the interacting O atoms (with additional relaxations). At a variance to the adsorption of \(S\), where there are two equivalent (by symmetry) interacting O

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\(^8\)One could also imagine a two step process \(S\rightarrow B\rightarrow T\) which would require 1.10 eV. However, the probability of going from the intermediate \(B\) backward to \(S\) is much higher than going forward. Therefore, the rate constants are determined by the highest transition state which is 1.25 eV above the initial configuration.

\(^9\)Already calculated for the purpose of investigation of the relative stability of the three isomers and presented in Sec.5.3.
Figure 5.9: Top and side views of the most stable adsorption configurations of the three Ir$_4$ isomers T, B and S at MgO(100) surface with neutral oxygen vacancy. Only the topmost MgO layer is shown. Color scheme is the same as in Fig.5.3. Mirror planes, the only symmetry operations, are also displayed. Upon charging the vacancy configurations remain similar to the displayed ones.

atoms, in the cases of T and B two possibilities exist. The most stable adsorption configuration of T is obtained by removing the oxygen atom that lays at the $\sigma_v$ mirror plane. In case of B the vacancy lays next to the oxygen that interacts with one of the wing–tip iridium atoms (Fig.5.9). Structures of all three isomers relax from their configurations when adsorbed on the defect–free surface bringing the Ir atom, that was interacting with the missing oxygen, closer to the vacancy. These relaxations are very close to a rigid shift of atomic positions towards the vacancy for both B and S. Average Ir–Ir distances associated to the two coordination shells of both B and S are only slightly bigger (Tab.5.4) than at defect–free MgO(100) surface. The angle of B structure is also only slightly changed to the value of 131.9°. On the other hand T distorts considerably. Distances between the two basal Ir atoms which interact with the remaining oxygen atoms are somewhat smaller (3.20 Å) as well as distances between them and the two other Ir atoms (average value is 2.41 Å). Distortion comes from the motion of the Ir atom, that interacts with the vacancy, towards the vacancy elongating the distance to the topmost Ir atom to the value of 2.75 Å. Therefore the coordination number of the first Ir–Ir shell is lowered to 2 and another shell appears with coordination number 0.5 due to the distortion. Again, the deformation lowers the energy of T structure (Tab.5.5), but now for 0.18 eV, while the energy of B isomer is 0.1 eV higher and the energy of S does not change its gas–phase value.

The Ir–O distances in all cases remain unchanged (they are smaller only for 0.01 Å). This is mainly due to the relaxation of the support. While the magnitude and directions of these relaxations for T adsorption are almost the same as of the defect–free surface leading to the similar deformation energy, relaxations of the support for the adsorption of B and S isomers are bigger. Therefore, the relaxations of the defected MgO(100) surface in the two cases (adsorption of B and S) are more expensive as shown in Tab.5.5. Magnetic moments of the clusters are also affected by the vacancy and while moment of T isomer increases for 1.19 $\mu_B$ this change is +0.56 and +0.28 $\mu_B$ for B and S, respectively.

Binding of the Ir$_4$ clusters to the charged vacancy is weaker than to the neutral one, but while the binding energy of T structure decreases by only 0.16 eV, binding energies of B and S
Table 5.4: Calculated properties of $\text{T}$, $\text{B}$ and $\text{S}$ Ir$_4$ isomers adsorbed in the vicinity of the oxygen vacancy ($\text{F}$ and $\text{F}^+$ centers) at MgO(100) surface. Cluster binding energies $E_b$, total energy $\Delta E$ relative to that of $\text{T}$ configuration, total magnetic moment (M), $z$-component of the induced dipole moment $p_z$, coordination number (N), average distance (R) of the corresponding coordination shell for all shells up to the distance of 3.5 Å and all types of neighbors.

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<tr>
<th></th>
<th>$\Delta E$ [eV]</th>
<th>$E_b$ [eV]</th>
<th>M [$\mu_B$]</th>
<th>$p_z$ [D]</th>
<th>Ir–Ir</th>
<th>Ir–O</th>
<th>Ir–Mg</th>
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<tr>
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<td>6.16</td>
<td>2.69</td>
<td>-3.59</td>
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<td>2.38</td>
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<td>3.24</td>
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<tr>
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<td>2.5</td>
<td>2.42</td>
<td>0.25</td>
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<tr>
<td></td>
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<td>0.5</td>
<td>3.63</td>
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<tr>
<td>$\text{S}$</td>
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<td>2.59</td>
<td>-3.96</td>
<td>2</td>
<td>2.32</td>
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<td>3.28</td>
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Ir$_4$/F$^+$ isomers drop as much as 0.42 eV for both structures. The energy ordering of the three isomers changes even more. Two structures $\text{T}$ and $\text{S}$ become almost degenerate (they differ by 0.01 eV in favor of $\text{T}$) and the difference to the least stable $\text{B}$ isomer further increases to 0.79 eV. It can be seen from Tab.5.5 that $\text{T}$ isomer lowers its energy by relaxing for remarkable 0.40 eV (change of -0.22 eV compared to adsorption at neutral vacancy). The Ir–Ir distances of $\text{T}$ cluster are now shorter while the two basal Ir atoms that interact with two oxygen atoms become further apart (Tab.5.4). However, this gain is compensated by the stronger relaxation of the support and somewhat weaker (by 0.1 eV) electronic interaction leading to the small net decrease in the total cluster binding energy.

Although the $\sim$0.4 eV drop in binding of $\text{B}$ and $\text{S}$ isomers is the same it has different origin. While in case of $\text{B}$ it comes from bigger support relaxations, for $\text{S}$ isomer this difference comes from electronic effects (Tab.5.5). Both clusters do not relax almost at all, but at variance to the support deformations in case of $\text{S}$ which are more similar to those of the neutral vacancy, the support relaxes considerably in case of $\text{B}$ isomer. This is predominantly due to the relaxation of the remaining interacting O atom which is displaced in case of $\text{B}$ ($\text{S}$) for 0.35 (0.26)Å in $x$ direction (Fig.5.10) and 0.22 (0.18) Å out of plane. This increases the deformation energy of the support in case of $\text{B}$ for 0.5 eV which is very close to the difference of binding to the neutral vacancy.

The magnetic moments of the three isomers are also influenced by the charging. The moment of $\text{T}$ structure is lowered for almost 2 $\mu_B$ upon charging and moments of the $\text{B}$ and $\text{S}$
Table 5.5: Deformation energies \( E_d = E_{\text{deformed}} - E_{\text{ground state}} \) (in eV) for the adsorption of \( \text{Ir}_4 \) clusters over MgO(100) with the neutral (F) and charged (\( F^+ \)) O vacancy. Values are for the three \( \text{Ir}_4 \) isomers (\( E^1_d \)) as well as for MgO(100) support (\( E^2_d \)). \( E_{\text{int.}} \) is purely electronic interaction energy (in eV) that can be calculated as the total cluster binding energy (Tab.5.4) plus the total deformation energy \( E^1_d + E^2_d \).

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<th>( \text{Ir}_4 )</th>
<th>( E^1_d )</th>
<th>( E^2_d )</th>
<th>( E^1_d + E^2_d )</th>
<th>( E_{\text{int.}} )</th>
<th>( E^1_d )</th>
<th>( E^2_d )</th>
<th>( E^1_d + E^2_d )</th>
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<td>0.48</td>
<td>6.80</td>
<td>-0.40</td>
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<td>0.53</td>
<td>6.69</td>
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<td>B</td>
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<td>0.93</td>
<td>1.03</td>
<td>6.24</td>
<td>0.04</td>
<td>1.43</td>
<td>1.47</td>
<td>6.26</td>
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<tr>
<td>S</td>
<td>0.00</td>
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<td>0.83</td>
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<td>0.00</td>
<td>0.87</td>
<td>0.87</td>
<td>5.51</td>
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</table>

Figure 5.10: Deformations of the MgO surface with neutral and charged O vacancy caused by the adsorption of the three \( \text{Ir}_4 \) isomers (T, B and S). Arrows denote displacements that are shown only for the interacting O as well as the neighboring Mg atoms. Numbers in parentheses are the \((x, y, z)\) displacement vectors (in Å). They are given only for those atoms nonequivalent by symmetry. Black color is used for the displacements in the case of neutral while displacements in the case of charged vacancy are given in red.

are bigger by 0.26 and 0.85 \( \mu_B \), respectively. These changes can be attributed to the removal of an electron (charging) from different spin states of the three isomers and smaller portion to the electron transfer, upon charging, between the two spin channels.

The electronic effects of the neutral (as well as charged) O vacancy in the cases such as these, where vacancy introduces only relaxations of the same adsorption geometries as at the defect–free MgO surface, can be understood by analyzing changes in the electronic structure of the support and how it influences the interaction with adsorbates. Instead of having three oxygen 2p orbitals at the site from which the O atom is removed, there is only one surface state. This state is totally symmetric with respect to the site symmetry of the missing oxygen (in this case \( C_{4v} \) point group) as shown in Fig.5.1. It is shifted in energy compared to the analogous state (totally symmetric 2pz) of the missing oxygen, and its charge distribution is different as well. However, due to the relaxations, overlap of molecular orbitals of the \( \text{Ir}_4 \) isomers with the state in the vacancy remains similar, for all three isomers, to the that with the 2pz orbital of the missing oxygen. The fact that surface dipoles remain almost the same upon creating the vacancy goes in line with this assumption, since, as shown for the Ir adsorption (Sec.5.2), charge transfer occurs from the states perpendicular to the surface.
Creation of the vacancy affects, above all, the interaction of the cluster molecular orbitals with $2p_x$ and $2p_y$ states that are now missing. What really makes the difference between the electronic interaction of the three clusters (Tab.5.5) is the number of remaining surface states available for bonding. In case of $S$ isomer the number of O–$2p$ orbitals decreases from 9 to 7 (22 % change) while in the other two it goes from 6 to 4 (33 % ) upon creating the vacancy. In Fig.5.11 DOSs of the subsystems (clusters and the support) prior and of the full system after the interaction are shown. The vacancy state appears in the band-gap of the surface, as already explained in Sec.5.1.1. It interacts mainly with the molecular orbitals of the three isomers close to the Fermi energy. The difference in the number of O–$2p$ states that are interacting with the three isomers is clear. The 0.35 eV difference in electronic interaction between $B$ and $S$ can be again attributed to the lower symmetry of the $B$ isomer and the ability of its molecular orbitals to hybridize.

Upon charging, electronic interaction of both $T$ and $B$ isomers remains almost the same as when adsorbed to the neutral vacancy (Tab.5.5), while it further decreases in case of $S$. This can also be understood by analyzing electronic structure of the three systems. There is, due to similar geometry, one-to-one correspondence between electronic orbitals of the clusters adsorbed to the neutral and charged O vacancy. Then, charging of the vacancy can be viewed as removal of an electron from one of the states of the system when the cluster is adsorbed at the neutral vacancy site. On the far right in Fig.5.11 electronic orbitals, for each adsorbed isomer, from which the electron is removed are shown as well as their position in the
corresponding spectrum. Differences come from different character of those orbitals. In case of \( T \) the most favorable option is to remove an electron from a state, which almost does not interact with the surface. Induced relaxations of the cluster and the support cancel leading to only slightly changed electronic interaction energy (\( \sim 0.1 \text{ eV} \)). On the other hand in case of \( B \) the state is a weak bonding cluster–vacancy combination, while the corresponding state of \( S \) structure has the same, but much more pronounced bonding character. Small reduction in binding of \( B \) to the vacancy is compensated by strengthening other interaction channels due to bigger deformations of the surface resulting in the same electronic interaction energy. This is not the case for \( S \) isomer whose electronic interaction decreases for \( \sim 0.4 \text{ eV} \). In addition, charging is followed by a certain electron transfer between the spin channels for both \( T \) and \( B \) systems.

When comparing theoretical results for the three clusters adsorbed at the defected MgO(100) surface (Tab.5.4) with EXAFS measurements (Tab.5.2) it is evident that disagreement becomes even bigger. Calculated Ir–Ir distances are still too short and the Ir–O coordination numbers do not agree at all. Although, by introducing the vacancy relative stability of the three \( \text{Ir}_4 \) isomers is considerably affected, \( T \) structure becomes very close in energy to \( S \) at the neutral and degenerate at the charged vacancy, the distortions are so strong that it would not be possible to identify it in experiments as a tetrahedral–like any more.

### 5.6 Adsorption of \( \text{Ir}_4 \) on \( \text{TiO}_2(110) \)

Several experimental studies on adsorbed iridium nanoparticles and small Ir clusters on the surfaces of both anatase and rutile TiO\(_2\) have been performed recently [126, 10, 11]. Various techniques are used for preparation and characterization such as decarbonylation of \( \text{Ir}_4(\text{CO})_{12} \) carbonyl molecules dispersed on TiO\(_2\) powder (mixture of rutile and anatase phases) and EXAFS spectroscopy [126], size and energy–selected deposition of \( \text{Ir}_n^+ \) \((n = 1, 2, 5, 10, 15)\) clusters on rutile TiO\(_2\)(110) and X-ray photoelectron spectroscopy (XPS) [10] or the vapor–deposition of Ir nanoparticles also on rutile TiO\(_2\)(110) and scanning tunneling microscopy (STM) [11]. The authors did not observe any aggregation of the Ir clusters. Moreover, there is no oxidation/reduction reactions at the surface–cluster interface [10] similarly to the adsorption of other transition metals on TiO\(_2\)(110) surface (see reference [106] and the references therein). This is also supported by theory which does not predict any significant charge transfer between the stoichiometric surface and adsorbed late transition metal atoms [127, 128].

Similar kind of analysis as for MgO(100) (Sec.5.2) for the adsorption of atomic Ir on the TiO\(_2\) surface has been carried out. Convergence tests showed that the theoretical model presented in Sec.5.1.2 (unit cell size, slab and vacuum thickness, \( k \)-point sampling, . . . ) describes well the adsorption of atomic Ir. Equilibrium adsorption geometries of Ir\(_4\) clusters on TiO\(_2\)(110) surface are obtained in the same way as for adsorption on MgO(100) support (Sec.5.3). Details of the adsorption energetics, electronic structure and geometries of the three Ir\(_4\) isomers (\( T \), \( B \) and \( S \)) are presented in Fig.5.12 and Tab.5.6. Due to the large differences in the binding energies of the three isomers their relative stability is considerably altered. The \( S \) isomer is now only by 0.03 eV more stable, while \( B \) structure becomes 0.22 eV higher in energy than \( T \). The difference of 0.03 eV is very small and therefore \( T \) and \( S \) structures may be considered as two degenerate Ir\(_4\)/TiO\(_2\)(110) configurations. Magnetic moment of \( S \) isomer is reduced from the gas–phase 8 \( \mu_B \) to 0 \( \mu_B \), while \( T \) and \( B \) structures retain their gas–phase
magnetization upon adsorption of 0 and 2 $\mu_B$, respectively.

Geometries of $T$ and $S$ isomers are not altered much by the adsorption. Their average Ir–Ir distances are elongated only by 0.03 Å. Consequently their deformation energies are also small and amount to 0.11 and 0.17 eV, respectively (Tab.5.7). These deformations do not influence the gas–phase magnetization of the two isomers. On the other hand $B$ structure relaxes somewhat more. The average bonding Ir–Ir distance is longer by 0.08 Å and angle between the two wings is reduced to 127.68° (gas–phase value is 138.13°). This has almost no effect on the distance between the two wing–tip Ir atoms that are separated by 3.74 Å, very similar to the gas–phase 3.79 Å. The energy needed for the deformation of $B$ isomer is also bigger (0.29 eV) and its gas–phase magnetic moment increases to 4 $\mu_B$ due to the deformation. It decreases back to 2 $\mu_B$ upon interaction of the deformed cluster with the TiO$_2$ surface.

On the other hand, adsorption of the three Ir$_4$ isomers induces considerable relaxations of TiO$_2$(110) surface, much bigger than defect–free MgO(100) (Sec.5.3). The surface bridging O atoms that interact with the clusters (Fig.5.12) move parallel to the surface approaching the three adsorbates ($T$, $B$ and $S$) by, on average 0.18, 0.20 and 0.23 Å, respectively. At variance to $S$ which interacts only with bridging O atoms, $T$ and $B$ isomers interact also with one of the basal O atoms that is displaced by the interaction by 0.2 and 0.26 Å vertically out of the surface, respectively. In all three cases the nearby Ti(5c) atoms positioned below the three isomers move up from the surface. While for the adsorption of $T$ and $S$ isomers there are two cations which are displaced about 0.2 Å, in case of $B$ only one Ti atom relaxes considerably, by 0.3 Å in the same direction. The neighboring basal O atoms positioned below the adsorbates at the distance ≈2.9 Å (just one in cases of $T$ and $B$ and two in for $S$ adsorption) are pushed into the surface by ≈0.5 Å. The surface relaxations can be attributed to two main reasons: (i) chemical bonding between the three clusters and bridging O atoms and (ii) the electrostatic interaction between the Ti(5c) and basal O atoms that are in the high oxidation states and the charge from the clusters accumulated in the interfacial region.

**Figure 5.12:** Equilibrium atomic configurations of $T$ and $S$ Ir$_4$ isomers when adsorbed on stoichiometric TiO$_2$(110) surface. Numbers above each structure denote corresponding total energies calculated with respect to $T$ isomer. Energy barrier for the transformation between the two Ir$_4$ structures is also shown.
Table 5.6: Calculated properties of T, B and S Ir₄ isomers adsorbed on stoichiometric rutile TiO₂(110) surface. Cluster binding energies $E_b$, total energy $\Delta E$ relative to that of T configuration, total magnetic moment $M$, z-component of the induced dipole moment $p_z$ calculated using Eq.5.1, coordination number $N$, average distance $R$ of the corresponding coordination shell for all shells up to the distance of 3.5 Å and all types of neighbors. Available experimental data are also presented.

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</tr>
</tbody>
</table>

¹ Values taken from Ref.[126]

Table 5.7: Deformation energies $E_{1d}^i(Ir_4)$ and $E_{2d}(TiO_2)$ of the three Ir₄ isomers (T, B and S) and TiO₂(110) surface are shown together with the electronic interaction energies $E_{int}$. All values are in eV.

<table>
<thead>
<tr>
<th></th>
<th>$E_{1d}^i(Ir_4)$</th>
<th>$E_{2d}(TiO_2)$</th>
<th>$E_{1d}^i+E_{2d}^i$</th>
<th>$E_{int}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>0.11</td>
<td>3.49</td>
<td>3.60</td>
<td>9.18</td>
</tr>
<tr>
<td>B</td>
<td>0.29</td>
<td>3.88</td>
<td>4.17</td>
<td>8.95</td>
</tr>
<tr>
<td>S</td>
<td>0.17</td>
<td>3.76</td>
<td>3.93</td>
<td>8.03</td>
</tr>
</tbody>
</table>

Similarly to the analysis of the adsorption on defect–free MgO(100) surface, structural transformations between the three configurations are studied by performing the CINEB calculations (Sec.2.2.5). The minimal energy path between T and S structures is similar to that in the gas–phase¹⁰ and passes through B configuration. However, the energy barriers of 0.57 eV for the transformation from T to B and 0.91 eV from B to S isomer are much different from the corresponding barriers both in the gas–phase and on MgO(100) surface (Sections 3.4 and 5.3). At TiO₂(110) the differences can be attributed to the different number of bonds that need to be break simultaneously in order to perform the transformations. It is necessary, for both transformations, to break simultaneously one Ir–Ir bond (as in the gas–phase) and one bond formed upon adsorption between one of the Ir atoms and one basal O atom (check Fig.5.12). The transition states lay, in both cases, in the configurations where both Ir–Ir and Ir–O bonds are broken and the new ones with bridging O atoms are not fully established yet. This is the reason for a much higher activation energies for structural transformation between the three isomers.

¹⁰Of course, when considered only in the Ir₄ configuration space.
Even though, the barriers are much higher than in the gas-phase and on the MgO surface the rate constants are still large enough at the temperatures of several hundred degrees Celsius. Under the assumption that the prefactors in Eq.2.19 are $\sim 10^{12}$ the rate constants at 300° C (experimental conditions) lay in the interval from $10^2$ Hz for the backward $B \leftarrow S$ transition, to $\sim 10^8$ for the transformation $T \leftarrow B$. All half–lives are short enough compared to the time interval (3 h) of the treatment at 300° C that the equilibrium population of the three minima can be considered established. The equilibrium populations of the three local minima ($T$, $B$ and $S$) can be easily estimated, under the assumption that the populations are described well by Boltzmann distribution (which is the case in the transition state theory, Sec.2.2.5), to 34.7, 0.3 and 65.0 %, respectively. Therefore, when adsorbed on TiO$_2$(110), most of the Ir$_4$ still adopt $S$ geometry, but with the significant fraction of those in $T$ configuration. Of course, this distribution can be largely affected by the changes in relative energies of the order of 0.01 eV. Since the accuracy of DFT methods (or at least of the model used in this work) is also of this order it only possible to conclude that at TiO$_2$ a significant fraction of the adsorbed Ir$_4$ species will adopt $T$ geometry.

This result is in much better qualitative accord with experiment than for the adsorption on MgO(100) surface. The differences compared to the experimental results could also arise since Ir$_4$/TiO$_2$ clusters are produced using the Degussa P25 TiO$_2$ powder [126] which consists of 3/4 of anatase and 1/4 of rutile phases. Although rutile (110) surface that is used in calculations is similar to the most stable anatase (101) surface (they both have only Ti(5c) exposed to the vacuum) differences of the order $\sim 0.1$ eV in relative energies of different adsorption geometries are certainly possible.

Calculated Ir–Ir distances are also $\sim 0.2$ Å shorter than the measured ones and there are
some disagreements on the longer Ir–O coordinations. As already discussed in Sec.5.3 the discrepancies in the Ir–Ir distances may occur due to temperature effects while the longer Ir–O coordination could be different due to the fact that most of the adsorption occurs on different surface, anatase (101) instead of rutile (110).

As in previous sections analysis of adsorption of Ir4 isomers to TiO2(110) surface is done by dividing the process in two contributions, deformations of both adsorbates and the support and energy gain due to the purely electronic interactions. These contributions are presented in Tab.5.7. It is evident that the different binding of B isomer compared to T is mainly due to bigger deformations 0.57 eV, while a smaller portion 0.23 eV comes from the electronic effects. On the other hand, only 0.33 eV of the difference in binding energy of S isomer comes from the bigger surface relaxations and much bigger part 1.15 eV is due to electronic effects. The electronic interaction energies can be further reduced to energies needed to change the magnetic moments of the three isomers and the electronic interactions of the Ir4 clusters with fixed magnetic moments. The T isomer does not change its gas–phase magnetization (0 \( \mu_B \)) neither due to deformations nor adsorption. Reduction of the magnetic moment of S structure due to adsorption is 8 \( \mu_B \) and the energy needed for this change is large and amounts to 0.51 eV. Therefore, almost one half of the difference in \( E_{\text{int}} \) between S and T structures is due to the reduction of the S magnetization. On the other hand, the moment of B isomer changes upon the relaxation from the gas–phase 2 to 4 \( \mu_B \) for the deformed isomer and it changes back to 2 \( \mu_B \) upon adsorption. The energy needed for this \( \Delta m = 2 \mu_B \) change is only 0.05 eV.

In Fig.5.13 details of the electronic structure of the three Ir4 isomers and the support prior and after the adsorption are presented. The Fermi levels of the free clusters are positioned deep in the conduction band of the support. The three clusters with zero magnetization and the support are in their relaxed atomic configurations. Since the total energy of adsorbed B structure with zero magnetization lays only 0.05 eV above the 2 \( \mu_B \) ground state, the analysis is done on the non–magnetic system. Main features of the electronic interactions of the three isomers are similar. Spectral weight of the atomic orbitals located on the interacting O atoms is distributed mainly in the valence with very small, almost negligible, amount in the conduction band. As in the case of the adsorption on defect–free MgO(100) the deformations
localizes these states close to the top of the valence band and the interaction pushes them to the lower energies. This effect is less pronounced on TiO$_2$(110) surface due to the much bigger energy difference between the valence band maximum and the electronic states of the three isomers around the Fermi energy. In accord with experimental findings [10] there is no significant transfer of the spectral weight across the Fermi energy after the adsorption.

On the other hand, states from Ti(5c) atoms, localized by the deformations at the bottom of the conduction band, are shifted after the adsorption to higher energies. Again, there is no transfer of their spectral weight to lower energies in the occupied parts of the three spectra. The distances between interacting Ir and basal Ti atoms are 2.55–2.65 Å. These values are too big for any appreciable overlap of their d–orbitals. Therefore, the interaction of Ir$_4$ with surface Ti atoms is mostly electrostatic. Electrons from the adsorbates, attracted by the positively charged surface Ti ions, approach closer to the surface destabilizing 3d orbitals of near titanium atoms. These polarization effects can be observed from the charge difference $\Delta \rho(x,y,z)$ shown in Fig.5.14. Density is mostly depleted from the basal Ir atoms and accumulated between them and the interacting O atoms (formation of covalent bonds) as well as in the interfacial region between the two clusters and the surface Ti ions. Consequently, Ti cations, also attracted by the electronic density accumulation, relax out of the surface while the O anions laying below the clusters are pushed by the electrostatic interaction deep into the surface. Since there is no significant charge transfer between two subsystems (cluster and the support) the induced dipole moments are mainly due to the polarization effects. They are oriented from the surface to the adsorbates in all three cases. Differences in the values of dipole moments result mainly from a different charge displacement along the vertical axis. These strong electrostatic interactions on the TiO$_2$ surface are a consequence of the anomaly in the effective (Born) charges present in the rutile TiO$_2$. It is already known that the effective charges of the bulk rutile TiO$_2$, depending on the nuclear displacement direction, are 5.20–7.70 electrons for Ti cations (check reference [129] p.57 and also the references therein).

Polarization of the three isomers is not only responsible for the surface relaxations and induction of the dipole moments, but is crucial for altering relative stability between the three isomers. This can be proved by placing the free clusters with the same magnetization and the configurations as when adsorbed on TiO$_2$, in the external electrostatic field. In periodic–supercell calculations this can be done by adding a saw–like potential to the Kohn–Sham Hamiltonian, whose slope in the region of the cluster is determined by the value of the electric field (for the details check the manual of Quantum-ESPRESSO code [35]). Calculations
are performed for several electric fields and the dependence of the relative energies of B and S calculated with respect to T isomer is presented in Fig.5.15. The relative energies depend strongly on the applied field. They increase monotonically as magnitude of the field grows. For the value of the field of 0.015 a.u. (1 Rydberg a.u. = 51.44×10¹⁰ V/m) relative energy of B with respect to T configuration becomes -0.04 and that of S structure is -0.54 eV. When the support deformation energies are added to these values the relative energies of the three isomers are 0.23 and -0.15 eV for B and S, respectively. These values are very close to the relative energies from Tab.5.6. Further increase of the applied field leads to further increase of relative energies of both B and S structures. It is even possible, for a sufficiently high fields (above 0.03 a.u.), to reverse completely the gas–phase energy order of the three isomers.

Moreover, if the electrostatic potential of the support is averaged in the plane parallel to the surface, but only in the region of space occupied by the clusters¹¹ then the surface electric field in the normal direction can be estimated as the derivative of the averaged potential (see Fig.5.16). If an average slope of the potential is calculated from the z = 0 in Fig.5.16 to the point where the potential is 5 % of its V(z = 0) value (∼ 2.4 Å) the obtained electrostatic field is 0.013–0.015 a.u. (depends on the adsorbed isomer). Just for comparison, the calculated field in the vicinity of MgO(100) surface is 0.006 a.u. and the potential drops to 5 % of its lowest value very fast at 0.97 Å which is approximately around the positions of the basal Ir atoms.

If the gas–phase Ir₄ clusters are allowed to relax both their atomic coordinates and magnetic moments the situation would not be the same. They would orient differently with

¹¹This region is defined as the region of space whose points are closer to the Ir atoms of the three clusters then to any other atom in the system.
respect to the applied field and S isomer would remain the most stable structure. Therefore, the effect of the surface on the isomer stability is twofold. Chemical bonds ensure the appropriate geometry of the three clusters and then the surface field makes the change in their relative energies. It is not unknown that external electric field may influence structures of Si₇₆C₅₋₇ₘ clusters introducing additional relaxations [130] or assist in dissolution of metal clusters in metal island films [131]. However, this is the first report on the surface affecting relative energies of different configurations of adsorbed clusters through its electrostatic field.
Chapter 6

Influence of Adsorbates on the Relative Stability of Oxide Supported Ir$_4$ Clusters

Interactions of Ir$_4$ clusters with different types of atomic (H, C and O) and molecular species (CO) have already been studied in the gas–phase (Chap.4). Presence of the substrate may affect the electronic structure of the supported species and in that way influence their interactions with the environment. Equivalently, interactions with the environment may also induce changes in the electronic structure of the supported clusters and influence their interaction with the support.

Interactions of both MgO(100) and TiO$_2$(110) supported Ir$_4$ clusters with atomic (H, C and O) as well as molecular species (CO molecules) are the subject of this chapter. The result obtained for the gas–phase Ir$_4$ that one carbon adatom alters the relative stability of different isomers (Sec.4.1) motivated research in this direction. Moreover, the discrepancies with experimental results about the structure of the MgO supported Ir$_4$ (Sec.5.3) remain unexplained. On the other hand, for TiO$_2$(110) substrate it has been shown that strong electrostatic field present in the vicinity of the surface can, to a large extent, explain properties of the adsorbed pristine Ir$_4$ clusters (Sec.5.6). It is interesting to study how does this field, which influences considerably both relative stability and the electronic structure of Ir$_4$, affect their interactions with the environment.

6.1 Interaction of MgO(100) Supported Ir$_4$ Clusters With H, C or O atoms

Many different adsorption geometries of atomic H, C and O on the three Ir$_4$ isomers supported by MgO(100) have been inspected and the most stable configurations are displayed in Fig.6.1. Their properties are presented in Tab.6.1. Again, available experimental results [118, 119] for the MgO supported Ir$_4$ clusters are presented next to the calculated ones for comparison.

Effects of the interaction of the MgO supported Ir$_4$ with atomic H, C and O (Sec.4.1) are similar to those in the gas–phase. Upon interactions the relative stability of the three supported Ir$_4$X (X = H, C and O) structures remains close to that predicted for the gas–phase Ir$_4$X (Tab.4.1). Again, only the C adsorption affects considerably the relative stability, but
Figure 6.1: The most stable configurations of Ir₄/MgO(100) clusters when interacting with atomic H, C and O. The same color scheme is used as in Fig.4.1 and Fig.5.5 except for the interaction with oxygen (the lower panel) where the O adatom is colored in yellow.

now with one important difference. While in the gas–phase B isomer with a single C adatom becomes the most stable structure, on MgO(100) surface T structure, in accordance with experimental results [118, 119], is the configuration with the lowest total energy (reported in Ref.[69, 70]). Carbon binding energies remain very close to the gas–phase values pointing out that the surface has no effect on the interaction of Ir₄ with the carbon adatom. However, since the differences in carbon binding energies between the three isomers stay close to the gas–phase values, crucial effect in this change of stability in favor of T structure is the reduction in relative energies of the pristine clusters due to the adsorption on MgO(100) surface. Namely, upon adsorption energy of pristine B relative to T isomer is reduced from the gas–phase -0.58 to only -0.14 eV. The same happens with the relative energy of S structure which goes from -1.51 to -1.06 eV. With these reductions differences in the carbon binding energies are large
Table 6.1: Calculated properties of \( T, B \) and \( S \) \( \text{Ir}_4X \) (\( X = \text{H}, \text{C} \) and \( \text{O} \)) clusters when adsorbed on \( \text{MgO}(100) \). \( E_b \) refers to the binding energy of the adsorbates \( E_b = -E(\text{Ir}_4X/\text{MgO}) + E(\text{Ir}_4/\text{MgO}) + E(X) \) and the total \( \text{Ir}_4X \) cluster binding energy given in parentheses. All other quantities are the same as in Tab.5.2.

<table>
<thead>
<tr>
<th></th>
<th>( \Delta E ) [eV]</th>
<th>( E_b ) [eV]</th>
<th>( M ) [( \mu_B )]</th>
<th>( p_z ) [D]</th>
<th>( \text{Ir-Ir} ) N</th>
<th>( \text{R}[^{\text{Å}}] )</th>
<th>( \text{Ir-O} ) N</th>
<th>( \text{R}[^{\text{Å}}] )</th>
<th>( \text{Ir-Mg} ) N</th>
<th>( \text{R}[^{\text{Å}}] )</th>
<th>( \text{Ir-C} ) N</th>
<th>( \text{R}[^{\text{Å}}] )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Ir}_4\text{H} )</td>
<td>( T ) 0.00</td>
<td>3.21(3.94)</td>
<td>0.58</td>
<td>-5.62</td>
<td>2.5</td>
<td>2.42</td>
<td>0.75</td>
<td>2.14</td>
<td>0.75</td>
<td>2.74</td>
<td>0.5</td>
<td>3.46</td>
</tr>
<tr>
<td></td>
<td>( B ) -0.03</td>
<td>3.09(3.15)</td>
<td>1.06</td>
<td>-4.24</td>
<td>2.5</td>
<td>2.41</td>
<td>0.5</td>
<td>2.05</td>
<td>1.2</td>
<td>2.88</td>
<td>0.5</td>
<td>3.66</td>
</tr>
<tr>
<td></td>
<td>( S ) -0.90</td>
<td>3.06(3.29)</td>
<td>1.22</td>
<td>-4.05</td>
<td>2.0</td>
<td>2.32</td>
<td>0.5</td>
<td>2.06</td>
<td>1.0</td>
<td>2.88</td>
<td>1.0</td>
<td>3.28</td>
</tr>
<tr>
<td>( \text{Ir}_4\text{C} )</td>
<td>( T ) 0.00</td>
<td>8.16(3.57)</td>
<td>1.95</td>
<td>-3.86</td>
<td>2.5</td>
<td>2.52</td>
<td>0.75</td>
<td>2.21</td>
<td>0.5</td>
<td>2.59</td>
<td>0.5</td>
<td>1.79</td>
</tr>
<tr>
<td></td>
<td>( B ) 0.15</td>
<td>7.86(3.08)</td>
<td>0.00</td>
<td>-4.00</td>
<td>2.0</td>
<td>2.44</td>
<td>0.5</td>
<td>2.07</td>
<td>0.5</td>
<td>1.78</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( S ) 0.65</td>
<td>6.46(3.22)</td>
<td>2.54</td>
<td>-3.87</td>
<td>2.0</td>
<td>2.41</td>
<td>0.5</td>
<td>2.05</td>
<td>0.5</td>
<td>1.85</td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \text{Ir}_4\text{O} )</td>
<td>( T ) 0.00</td>
<td>6.00(3.36)</td>
<td>2.01</td>
<td>-7.54</td>
<td>2.5</td>
<td>2.44</td>
<td>0.75</td>
<td>2.14</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( B ) -0.48</td>
<td>6.33(3.22)</td>
<td>0.44</td>
<td>-5.14</td>
<td>2.5</td>
<td>2.41</td>
<td>0.5</td>
<td>2.06</td>
<td>1.5</td>
<td>2.93</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( S ) -0.97</td>
<td>5.91(3.08)</td>
<td>0.00</td>
<td>-6.29</td>
<td>2.0</td>
<td>2.32</td>
<td>0.5</td>
<td>2.09</td>
<td>1.0</td>
<td>2.90</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exp.</td>
<td>Ref.[118]</td>
<td>2.6-3.1</td>
<td>2.62-2.73</td>
<td>0.6-1.1</td>
<td>2.07-2.16</td>
<td>2.5-3.0</td>
<td>2.59-2.73</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ref.[119]</td>
<td>2.9</td>
<td>2.62</td>
<td>1.1</td>
<td>2.05</td>
<td>0.6</td>
<td>2.45</td>
<td>0.5</td>
<td>1.91</td>
<td>0.5</td>
<td>3.33</td>
<td></td>
</tr>
</tbody>
</table>
enough to make $T$ the most stable Ir$_4$C/MgO(100) structure and to push $S$ isomer further apart from $T$ than in the gas–phase.

The carbon adsorption geometries (Fig.6.1) are the same as those in the gas–phase (Fig.4.1). Carbon binds the most favorably to the bridge site between two Ir atoms and induces breaking of the corresponding Ir–Ir bond of both $T$ and $B$ isomers. Similarly to the gas–phase the interaction of the C adatom with $S$ is weaker and the Ir–Ir bond bridged by carbon is not completely broken. Of course, not all Ir–Ir bridging sites are equivalent on the supported clusters and it is energetically the most favorable for C adatom to bind in such a way to minimize the interference between the two interactions, cluster–support and cluster–carbon.

The explanation for the stronger carbon binding on $T$ and $B$ isomers than on $S$ remains the same as for the gas–phase species (Sec.4.1). In Fig.6.2 the projected DOS on 2$p$ atomic orbitals of the C adatom is shown for all three isomers. The DOSs are very similar to those in the gas–phase (Fig.4.2); of course, with the differences produced by the interaction with the surface (mostly broadening of the peaks). For both $T$ and $B$ isomers more of the carbon spectral weight is shifted to lower energies than in the case of $S$ structure. Electronic orbitals formed upon interaction with the carbon resemble those shown in Fig.4.2. The antibonding molecular orbital (MO) of both $T$ and $B$ isomers, responsible for the particularly strong interaction with C–2$p$ atomic orbitals (AO) in the gas–phase, remains above the Fermi energy for the pristine clusters on MgO(100) despite modifications in their electronic structure induced by the support. The interaction with the C–2$p$ orbitals pushes these electronic states to the bottom of the carbon spectra as for the gas–phase species. The other interaction channels for all three isomers also stay similar to those in the gas–phase.

The carbon adatom induces some modifications in the cluster–support interaction. The total cluster binding energies of $T$ and $B$ structures are by 0.26 and 0.27 eV lower than for the pristine clusters adsorbed on the MgO surface. Energy gain due to the adsorption of Ir$_4$C in $S$ configuration (3.22 eV) remains much closer to the value for the pristine cluster (3.33 eV). This effect of somewhat weaker binding to the support for $T$ and $B$ structures with and without carbon is accompanied by the change in the induced dipole moment. In the case of $T$, structural distortions induced by the C adsorption weaken the Ir–O interaction.
Interaction with the carbon adatom also induces modifications in the geometries of the three isomers. The first shell Ir–Ir coordination number remains the same upon carbon adsorption for both T and S structures while its value decreases to 2.0 for B due to the breaking of the Ir–Ir bond bridged by the carbon. The distance between these two Ir atoms increases to 3.16 Å, value very close to the gas-phase (3.19 Å), while the distance between the two wing-tip Ir atoms becomes 4.11 Å (not presented in Tab.6.1). Although the C adsorption also induces breaking of one Ir–Ir bond of T structure the coordination number remains the same since the two basal Ir atoms of the pristine cluster which are by 3.68 Å apart, approach closer to each other upon C adsorption (2.49 Å) reestablishing the broken Ir–Ir bond.

As for the MgO supported pristine Ir$_4$ the best agreement between calculated and measured structural data is for T structure which is now the most stable and hence, the most probable Ir$_4$C/MgO(100) configuration. The average first shell Ir–Ir distance of the supported T isomer with the C adatom is only by 0.1 Å (~4 %) lower than those reported from experiments (see Tab.6.1). Moreover, T structure is less distorted than without the carbon. Still, there are some differences in the first shell Ir–O coordination and distances, but the data for Ir–Mg as well as Ir–C shells agree reasonably well. The results for B and S structures do not correspond well to experiment.

Finally, the vibrational analysis is carried out for the T isomer with the carbon adatom both in the gas-phase and supported by MgO(100). Calculations are done in the frozen phonon approximation (see Sec.2.2.4) allowing only C, Ir and the surface atoms that are close to the cluster to move. Due to the large mass difference of the carbon and iridium atoms, purely carbon vibrational modes appear in both spectra$^1$. The two highest frequency modes,

$^1$Amplitudes of the C and Ir atoms along these modes differ by an order of magnitude.
appear to be of the carbon character both on MgO(100) and in the gas–phase. They are well separated from all the other modes in both spectra. For the MgO supported (gas–phase) T structure, the frequencies of these two carbon modes are 113.2 (103.5) and 87.6 (87.9) meV. Motion of the C adatom along these modes is the same for both environments. The highest frequency mode is longitudinal with respect to the line connecting the two bridged Ir atoms, while along the second mode the motion of C is transversal as depicted on Fig.6.3. All the other vibrations of the system involve large displacements of the Ir atoms and the frequencies approach the values for the pristine gas–phase cluster (10–40 meV). Since the highest frequency mode lays well above the vibrational frequencies of the clean MgO(100) surface (∼88 meV∼709 cm⁻¹ [122]) it could serve for the experimental verification of the carbon presence².

At variance to the C adsorption both H and O adatoms do not influence considerably the energy ordering of the three Ir₄ isomers. In both cases S isomer remains the most stable form also when supported by MgO(100) surface. While the energy of S relative to T structure remains close to the values for the supported pristine clusters the relative energy of B isomer approaches close to zero for the H adsorption and to -0.5 eV after the interaction with the O adatom. Binding configurations are the same as in the gas–phase and they also do not correspond well to the experimental results (in addition to the different relative stability of Ir₄ isomers). The binding energies of the H adatom and Ir₄H cluster lay very close to the corresponding values in the gas–phase and for the adsorption of pristine clusters on the MgO, respectively. On the other hand, the oxygen binding energies are lower than in the gas–phase and the adsorption is accompanied by the charge rearrangements (both different transfer from the surface and different distribution of that charge) that are followed by the considerable increase in the induced dipole moments. These charge redistributions influence also the total cluster binding energies, especially in the case of T structure.

### 6.2 Interaction of MgO(100) Supported Ir₄ Clusters With CO Molecules

The lowest total energy configurations for the adsorption of one and two CO molecules on the three MgO(100) supported Ir₄ isomers are presented in Fig.6.4. The structural data and energetics are presented in Tab.6.2 and Tab.6.3. For the supported T isomer the study of the interaction with CO molecules was continued to the adsorption of up to five CO molecules. The work is motivated by the experimental procedure for producing supported Ir₄ clusters which involves decarbonylation of the supported Ir₄(CO)₁₂ molecules (Ref.[119] and references therein).

Similarly to the gas–phase, interaction with one CO molecule alters considerably the relative stability of the three isomers, but on the MgO the most stable isomer becomes T. However, it is only by 0.01 eV lower in energy than S and given the accuracy of the theoretical model (check Tab.5.1) these two structures can be treated as degenerate. On the other hand, B isomer interacting with one CO molecule becomes the least stable structure whose energy lays by 0.34 eV higher than that of T.

The CO binding configurations are different than in the gas–phase. On T structure the dissociative CO adsorption is still more favorable than adsorption in molecular form (by 0.51 meV).

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²Everything said here is under the assumption that the adsorption of Ir₄ clusters whose vibrational frequencies are 10–30 meV, would not produce vibrations whose frequencies are higher than ∼88 meV.
Figure 6.4: Configurations with the lowest total energies for the three MgO(100) supported Ir$_4$ isomers (T, B and S) interacting with one (upper panel) and two CO molecules (lower panel). The same color scheme as in Fig.6.3 is used.

eV) while to both B and S one CO binds preferentially as a molecule, by 0.76 and 0.87 eV stronger than if dissociated, respectively. The carbon atom binds upon dissociative CO adsorption to the three Ir atoms of T structure while the oxygen is at bridging site between the fourth iridium and the closest surface Mg atom (Fig.6.4). Binding of the C adatom at the three–fold site is also very favorable C adsorption geometry (without the oxygen) both in the gas–phase and on the MgO. It is by 0.19 and 0.13 eV higher in energy than the adsorption at the bridging site, respectively. The explanation for the strong carbon bonding to T isomer at the three–fold site is similar to that for the bridging configuration$^3$. Adding one oxygen atom to the MgO supported cluster the configuration from Fig.6.4 becomes more stable, by 0.35 eV lower in energy than that with the carbon at the bridging site and oxygen interacting with one of the two other basal Ir atoms. Interaction of the oxygen atom with the surface cation is crucial for the increased stability of the geometry for dissociative CO adsorption on T structure from Fig.6.4.

Molecular CO binds to both B and S as a terminal with the carbon atom in contact with the cluster, similarly to the gas–phase. However, the sites and orientation of the CO ligand are different. It binds on B structure to a wing–tip Ir atom instead to a middle one, and on S isomer to one of the top Ir atoms, almost orthogonal to the plane defined by the cluster. This indicates that the binding of the three clusters to the substrate and interaction with the CO molecule are not two independent processes. The values of the CO binding energies confirm

$^3$Strong interaction with the empty antibonding MOs of the cluster which are transformed by the interaction with the C adatom into the strongly bonding orbitals.
Table 6.2: Calculated properties of Ir₄CO in T, B and S configurations when adsorbed on MgO(100) surface (Fig.6.4 upper panel). $E_b$ refers to the binding energy of the CO molecule $E_b = - E(\text{Ir}_4\text{CO}/\text{MgO}) + E(\text{Ir}_4/\text{MgO}) + E(\text{CO})$ and the total Ir₄CO cluster binding energy given in parentheses (calculated using the gas-phase cluster with the CO molecule in the same form). All other quantities are the same as in Tab.5.2.

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Note: All values in parentheses are calculated for the gas-phase cluster with the CO molecule in the same form.
Figure 6.5: Motion of the C and O atoms on the MgO supported T isomer along the highest frequency vibrational mode upon (a) the dissociative CO adsorption and (b) the molecular CO adsorption. Numbers denote frequencies (in meV) of the corresponding modes.

this fact. They are all lower than in the gas–phase. This is also true for the total cluster binding energies.

The biggest structural modifications as well as modifications of the binding energies are induced for the CO adsorption on T isomer. The Ir atom that interacts with the oxygen coming from the CO molecule, relaxes out of the surface to the distance of 2.67 Å from the surface oxygen (check Fig.6.5). This reduces both total cluster binding energy to the surface as well as the charge transfer. The adsorption of O adatom is not any more independent of the C adsorption (as in the gas phase) since all Ir atoms interact either with the carbon adatom or with the surface O atoms. At variance to T, the CO and the total cluster binding energies of B and S structures remain much closer to the values for the CO adsorption in the gas–phase and adsorption of the pristine clusters on the MgO. Therefore, the crucial effect that enables the CO molecule to alter the relative stability of the three isomers is again (as for the C adsorption) the reduction of the pristine cluster relative energies due to the interaction with the support.

Calculated structural data for Ir₄CO/MgO(100) in T configuration with the dissociated CO molecule (Tab.6.2) agree with the experimental data as well as for T structure with only the C adatom (Tab.6.1). Calculated Ir–Ir coordinations and distances are almost the same while there are some differences in the other shells. However, those differences are not sufficient to determine which structure suits more to the measured one.

Since the carbon binding geometry differs from that without the coadsorbed oxygen, the carbon vibrational modes could have different frequencies. Moreover, the modes of the purely O character could also be expected. Both effects are confirmed by the vibrational analysis. The highest frequency mode of 98.9 meV = 797 cm⁻¹ is now the oxygen mode, while the following three modes are purely of the carbon character with frequencies 88.4 meV (713 cm⁻¹), 86.2 meV (695 cm⁻¹) and 66.6 meV (537 cm⁻¹). Lower values reflect weaker binding at the three–fold than the bridging site. Calculated carbon frequencies lay close and below

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4This coordination shell is not presented in Tab.6.2 since its average coordination number is only 0.25.
the upper limit $88 \text{ meV} = 709 \text{ cm}^{-1}$ [122] of the MgO(100) vibrational spectrum and it is certainly more difficult to distinguish them experimentally. However, the oxygen mode should be possible to detect since it lays about $90 \text{ cm}^{-1}$ higher than those of the MgO surface. Motion of the C and O atoms along the two highest frequency modes is presented schematically in Fig.6.5.

Vibrations of the CO molecule when adsorbed on the MgO supported T cluster are also studied. The most favorable geometry for the molecular CO binding is shown in Fig.6.5 together with the motions of the C and O atoms along the two highest frequency modes. This configuration is by 0.51 eV higher in energy than that for the dissociative adsorption. The CO binding energy in this configuration is 2.98 eV. The CO stretching mode of 245.5 meV = 1980 cm$^{-1}$ is the highest in frequency. The frequency of the second highest mode, that involves motion of the C and O atoms, lays well below at 68.7 meV = 554 cm$^{-1}$. The C and O atoms move in–phase along this mode in the direction of desorbing from the cluster (see 6.5). The CO stretching mode is used in experiments for the verification of the presence of the carbonyl ligands during the preparation process [123]. Calculated CO stretching frequency agrees very well with the range of measured values $1884–2052 \text{ cm}^{-1}$ attributed to the same mode for the MgO supported Ir$_4$(CO)$_{12}$ molecules [123].

### 6.2.1 CO Dissociation

Until this point the issue of the CO dissociation, which would change the relative stability of the three Ir$_4$ isomers in favor of the T structure, has only been addressed in the gas–phase (Sec.4.3). It has been found that the barrier for the CO dissociation, although quite high, is the lowest on the T isomer (2.67 eV). This is due to the peculiar situation where the transition state for the CO dissociation is the same for all three isomers. The reaction requires the smallest amount of energy on the T isomer since the configuration with the molecular CO, is the highest in energy (closest to the transition state). However, main consequence is that on the T structure the CO dissociation requires less energy (2.67 eV) than the CO desorption (3.16 eV).

The CO dissociation on the MgO supported Ir$_4$ is studied only on the supported T structure since only in this case the reaction would be spontaneous (exothermic). Since the dissociative CO adsorption on the other two structures (B and S) is much less favorable, one cannot expect that the large fractions of CO molecules on these two isomers would be dissociated in real situation. Moreover, in experiments the decarbonylation is performed on the iridium carbonyl cluster which has a tetrahedral metal frame.

Predicted minimal energy path (MEP) for the CO dissociation occurring on the MgO supported T structure is presented schematically in Fig.6.6. Calculated energy barrier for the process is 2.97 eV which is only by 0.01 eV lower than the molecular CO binding energy (see previous section). Hence, on the MgO supported structure it is as favorable to dissociate one CO molecule as to desorb it$^5$. Still activation energy is quite high and would require (in the thermal equilibrium) very high temperatures to occur.

The crucial step for the dissociation is again, as in the gas–phase, the configuration in which the carbon atom resides at a bridging site. As shown in Fig.6.6 the geometry of the transition state is such that both C and O atoms are bridging the Ir–Ir bonds. As already

$^5$Calculations of the MEP for the CO desorption yield no barrier for the reaction. The energies of the intermediate configurations increase monotonically to the energy of the configuration with the CO molecule far away from the cluster.
explained for the gas-phase clusters this enables the 2p orbitals of the carbon atom to bind strongly to the cluster and weakens the CO bond. This effect is very strong since it lowers the energy needed for the CO dissociation from 10.97 eV for the free molecule (its atomization energy) to ”only” 2.97 eV. However, this does not mean that Ir\textsubscript{4} clusters are good catalysts for this reaction (as for example Ru, Co or Fe [93]).

A notion of the rate constants for the two reactions could be obtained by employing Eq.2.19. The vibrational frequencies of the molecular CO adsorbed on the MgO supported T structure (previous sections) could serve as the approximate prefactors. Since the CO stretching mode leads to the breaking of the CO molecule, the prefactor for the dissociation would then be of the order of $10^{14}$ Hz, while for the desorption (the other mode from Fig.6.5) it is $10^{13}$, by an order of magnitude lower. Similar energy barriers for both dissociation and desorption imply that the fraction of dissociated CO molecules would be an order of magnitude larger than the fraction of those which desorb. However, since the activation energies for the two processes are very high the timescale for the two reaction are very long compared to the experimental 3h, and by far the biggest fraction of the CO molecules, in the conditions of the thermal equilibrium with the atmosphere surrounding the system, would remain chemisorbed in the molecular form at typical experimental conditions (300° C). The issues of the influence of the environment on the number of chemisorbed CO molecules is discussed in the next section.

### 6.2.2 Multiple CO adsorption

In the case of adsorption of two CO molecules on Ir\textsubscript{4}/MgO(100) clusters, T isomer becomes the most stable structure (Tab.6.3). It is by 0.66 and 0.49 eV lower in energy than B and S, respectively. Moreover, the CO dissociation is no longer favorable on any of the three isomers (check the most stable adsorption configurations in Fig.6.4). The formation of CO\textsubscript{2} on T
structure is also not a favorable process. Total energy of the system with two adsorbed CO molecules is by $\sim 3.5$ eV lower than the configuration with the bridging C adatom and the CO$_2$ molecule in the vacuum.

Total cluster binding energies are not altered much by the interaction with the two CO molecules (less than 10% in all three cases). They are calculated using the gas–phase Ir$_4$ cluster with the same number of molecular CO attached to it, which is not necessarily the most favorable gas–phase Ir$_4$(CO)$_2$ geometry. This done in order to show that whatever is the number of CO molecules attached to the cluster (Tab.6.3) it is always energetically more favorable to desorb one CO molecule than the whole cluster (compare the values for the binding energy of nth CO molecule and the total cluster binding energies).

If the number of the adsorbed CO molecules increases the binding energy of the nth CO molecule, calculated as $E_b = -E(Ir_4(CO)_n) + E(Ir_4(CO)_{n-1}) + E(CO)$, decreases as it would normally be expected. However, the binding energy of the fifth CO molecule drops significantly compared to the value for the fourth ($\sim 1.0$ eV). This is accompanied by a similar drop of the total cluster binding energy. The decrease in the total cluster binding energy is a consequence of the modifications of the cluster geometry induced by the fifth CO molecule. Namely, the fifth CO binds as a terminal to one of the basal Ir atoms already having one CO molecule attached to it which produces a drop in the CO binding energy. Moreover, it is energetically more favorable for the cluster to interact with the CO than with surface and even repulsive interaction with the surface oxygen atom is produced that pushes the Ir atom away from the surface. The resulting Ir$_4$ configuration is very similar to B geometry with two contact points to the surface.

Structural data for the supported T isomer with up to five CO molecules agree relatively good with the measured ones. The average first–shell Ir–Ir distance increases even to 2.58 Å which is very close to the experimental values (2.62–2.73 Å). Of course, the Ir–C and longer Ir–O coordination numbers increase, but the distances remain relatively independent on the coordination. Only the Ir–Mg coordination shells are affected considerably. Since there are two Ir–Mg shells, for the pristine T isomer, whose average distances are very close ($\sim 0.3$ Å

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6This is true in the case of the gas–phase T isomer where it is still more favorable to have one CO dissociated and the second one in the molecular form.
Table 6.3: Calculated properties of Ir₄(CO)₂ in T, B and S configurations when adsorbed on MgO(100) surface. For T configuration the data for n = 3, 4, 5 are also presented. Eₘₚ refers to the binding energies of the nth CO molecule Eₘₚ = - E(Ir₄(CO)ₙ) + E(Ir₄(CO)ₙ₋₁) + E(CO) and total Ir₄(CO)ₙ cluster binding energies (in parentheses). All other quantities are the same as in Tab.5.2.

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difference, Tab.5.2) small cluster relaxations induced by the CO adsorption, for three and five CO molecules, join these two shells making one broad Ir–Mg shell which is not presented in tables. In the cases when this does not happen the relaxations increase shorter Ir–Mg distance which gets further away from the measured values. However, whatever is the number of the CO molecules adsorbed on each of the supported Ir₄ clusters, it would produce an infrared signal in the range of the CO stretching mode⁷ which contradicts the experiments where the full decarbonylation is performed [118, 119]. Therefore, it follows from both theory and experiments that the atomic species, very likely one C adatom or coadsorbed atomic C and O, change the relative stability of the three isomers in favor of T structure.

From the data presented in Tab.6.2 and Tab.6.3 it is possible to estimate, using simple thermodynamical considerations, what kind of external parameters are required for the decarbonylation process to occur. Of course, this kind of analysis is restricted to the case of the thermal equilibrium. One could imagine that the MgO supported clusters are surrounded by the CO atmosphere that is described by the temperature T and pressure p. By following the route of Reuter and Scheffler [132] one can define the adsorption free energy per cluster

\[ \gamma_n(T, p) = n \times (-E_b^{(n)} - \Delta \mu_{CO}(T, p)), \]  

where n is the number of adsorbed CO molecules, \( E_b^{(n)} \) is the average CO binding energy per molecule that depends on n (see Fig.6.8(a)) and \( \Delta \mu_{CO}(T, p) \) is the chemical potential of the CO molecules in the gas–phase at temperature T and pressure p. It is important to underline that all configurational and vibrational entropy effects for the adsorbed CO molecules have been neglected⁸. The dependence of \( \Delta \mu_{CO}(T, p) \) on p at different temperatures

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⁷The CO stretching frequency is very robust and although the its shift is used to identify whether the CO binds as a terminal or in a bridging configuration [15], the frequency remains close to 2000 cm⁻¹ and is easily detectable.

⁸The vibrational degrees of freedom could be neglected to a good approximation since the energies related to this type of motion (order of meV) are much smaller than the CO binding energies. On the other hand the CO configurational entropy is neglected due to practical reasons since it would require obtaining all possible CO
can be calculated using the chemical potential $\Delta \tilde{\mu}_{CO}(T, p^0)$ at the standard pressure ($p^0 = 1 \text{ atm}$)

$$
\Delta \mu_{CO}(T, p) = \Delta \mu_{CO}(T, p^0) + k_B T \ln \frac{p}{p^0},
$$

that on the other hand, can be calculated using the tabulated values for the CO formation enthalpy $H(T, p^0)$ at standard pressure and entropy $S(T)$ \[87\]

$$
\Delta \mu_{CO}(T, p^0) = H(T, p^0) - H(T = 0, p^0) - S(T) T.
$$

The value $\Delta \tilde{\mu}_{CO} = -1.17 \text{ eV}$ at $T = 600 \text{ K}$ is obtained. Temperature of 600 K is close to the experimental temperature of 300° C \[118, 119\].

Dependence of $\gamma_n(T, p)$ on pressure is plotted in Fig.6.8(b) for up to five CO molecules adsorbed on the MgO supported Ir$_4$ in $T$ geometry. The $\gamma_1(T, p)$ for both dissociated CO molecule and adsorbed in the molecular form are shown. The most favorable number of adsorbed CO molecules at a given pressure is determined by the lowest $\gamma_n$ value. It could have been anticipated that due to high CO binding energies it is not very favorable to desorb CO molecules. The data show that at pressures close to $p_0$ it is the most favorable to have five CO molecules adsorbed on the clusters\(^9\). The desorption starts at very low pressures about exp(-10)$p_0$ and the full decarbonylation requires pressures which are practically zero. This indicates that in experiments the crucial are the nonequilibrium conditions under which the decarbonylation of the deposited Ir$_4$ is performed. Namely, it is done \[118, 119\] in the atmosphere of the flowing He gas which removes all the desorbed CO molecules. In these conditions at the point where there is only one CO molecule per cluster it is certainly possible also to dissociate the CO ligands.

### 6.3 Interaction of TiO$_2$(100) Supported Ir$_4$ Clusters With a C Atom or a Single CO molecule

Interaction of TiO$_2$(110) supported Ir$_4$ clusters only with a C adatom and a single CO molecule has been studied. The adsorption geometries with the lowest total energies are displayed in Fig.6.9 and the corresponding calculated data presented in Tab.6.4.

Upon adsorption of a single C adatom, the relative stability of the three isomers changes in favor of $T$ structure. It becomes by 0.66 and 0.43 eV more stable than $B$ and $S$, respectively. However, the binding energies of the C adsorbate and binding energies of the total Ir$_4$C cluster become considerably different than those for the adsorption of the C adatom to the gas–phase clusters as well as those for the adsorption of the pristine clusters to the surface. Binding of the C adatom on the TiO$_2$(110) supported $T$ and $B$ isomers is as much as 0.81 and 1.00 eV weaker than in the gas–phase, respectively. Moreover, the total cluster binding energies of both structures become by 1.00 ($T$) and 1.22 ($B$) lower as well. On the other hand, binding of the C adatom to $S$ structure is by 0.55 eV stronger than in the gas–phase and the binding energy of the total cluster also increases by 0.36 eV.

It has been shown in Sec.5.6 that the pristine Ir$_4$ clusters to a good extent keep their gas–phase geometries. Therefore, the origin of the changes in the interactions with adsorbates

---

\(^9\)Or even six! The adsorption of more than five CO molecules has not been studied yet.
Figure 6.9: Adsorption configurations with the lowest total energies of a single C adatom and CO molecule on the three TiO$_2$(110) supported Ir$_4$ isomers.

lays in the differences in their electronic structures induced by the support. It is also shown in the same section that in the vicinity of TiO(110) there is a strong electrostatic field which polarizes the adsorbed clusters. The polarization affects the gas-phase relative stability of the three isomers and certainly affects their electronic structures. In fact, symmetry breaking accompanied by the surface field leads to mixing of the molecular orbitals of the three clusters which on the other hand influences their interaction with the carbon adatom.

This effect is particularly strong for T and B isomers. Carbon adatom binds in the same bridging configuration to all three structures which indicates that the bonding mechanism itself remains similar to that in the gas-phase (and on the MgO surface). However, the strongly interacting antibonding orbital of both T and B structures (see Sec.4.1) is now involved in linear combinations with other MOs of the two clusters and becomes partially filled. Moreover, the linear combinations in which this orbital participates located above the Fermi energy are shifted to higher energies. Therefore, the energy gain due to the interaction of the C–2p orbitals with this MO of the two isomers (which is now broadened over the electronic spectrum) is lower than in the gas-phase. In the case of B structure this interaction channel is even not energetically favorable. This is the reason why the Ir–Ir bond of B isomer bridged by the carbon is not broken and the C binding energy approaches that of S structure. The Ir–Ir bond of B isomer bridged by the carbon atom is elongated to the distance of 2.65 Å.
Table 6.4: Calculated properties of Ir$_4$C and Ir$_4$CO molecules in T, B and S configurations adsorbed on stoichiometric rutile TiO$_2$(110) surface. $E_b$ refers to the binding energy of either C adatom or the CO molecule and the total cluster binding energy (in parentheses). All other quantities are the same as in Tab.5.6.

<table>
<thead>
<tr>
<th></th>
<th>$\Delta E$ [eV]</th>
<th>$E_b$ [eV]</th>
<th>$M$ [$\mu B$]</th>
<th>$p_z$ [D]</th>
<th>Ir–Ir</th>
<th>Ir–O</th>
<th>Ir–Ti</th>
<th>Ir–C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ir$_4$C</td>
<td>T</td>
<td>0.00</td>
<td>7.43 (4.58)</td>
<td>0.00</td>
<td>2.21</td>
<td>2.5</td>
<td>2.51</td>
<td>0.75</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5</td>
<td>3.10</td>
<td>1.25</td>
<td>2.93</td>
<td>1.0</td>
<td>3.27</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td>0.66</td>
<td>7.00 (3.56)</td>
<td>0.00</td>
<td>0.84</td>
<td>2.5</td>
<td>2.53</td>
<td>0.75</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5</td>
<td>3.70</td>
<td>0.75</td>
<td>2.47</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>0.43</td>
<td>6.97 (4.46)</td>
<td>0.00</td>
<td>0.47</td>
<td>2.0</td>
<td>2.36</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.0</td>
<td>3.34</td>
<td>2.0</td>
<td>2.96</td>
<td>1.0</td>
<td>3.35</td>
<td></td>
</tr>
<tr>
<td>Ir$_4$CO</td>
<td>T</td>
<td>0.00</td>
<td>2.36 (4.77)</td>
<td>0.00</td>
<td>1.19</td>
<td>3.0</td>
<td>2.53</td>
<td>0.75</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.0</td>
<td>2.95</td>
<td>0.5</td>
<td>2.93</td>
<td>0.5</td>
<td>3.38</td>
<td></td>
</tr>
<tr>
<td></td>
<td>B</td>
<td>0.11</td>
<td>2.49 (3.70)</td>
<td>0.00</td>
<td></td>
<td>2.5</td>
<td>2.50</td>
<td>1.25</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5</td>
<td>3.69</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>-0.28</td>
<td>2.60 (4.01)</td>
<td>0.00</td>
<td>-0.22</td>
<td>2.0</td>
<td>2.42</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.0</td>
<td>3.42</td>
<td>2.5</td>
<td>2.89</td>
<td>1.0</td>
<td>3.30</td>
<td>0.5</td>
</tr>
<tr>
<td>Exp.</td>
<td>Ref.[126]</td>
<td></td>
<td></td>
<td>3.3</td>
<td>2.68</td>
<td>0.9</td>
<td>2.05</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.2</td>
<td>3.21</td>
<td>0.9</td>
<td>2.05</td>
<td>0.7</td>
</tr>
</tbody>
</table>
which is much less than for the C adsorption in the gas–phase (3.19 Å). The C adsorption on T structure does induce breaking of the bridged Ir–Ir bond (3.10 Å), but the energy gain due to the interaction is considerably lower than in the gas–phase.

Binding energy of the C adatom to S increases compared to the gas–phase value. The reason for this lays mainly in the energy cost for hybridization of the MOs of the cluster. Namely, it has been discussed in Sec.4.1 that the MOs of S structure, that interact strongly with the C–2p states, are formed by the hybridization of the several both empty and filled MOs of the pristine cluster. At variance to the gas–phase, on the TiO₂ surface the modifications of the cluster’s electronic structure induced by the support in fact, facilitate the C binding since the necessary hybridization requires less energy (it is already there).

The total cluster binding energies are also affected by the adsorption of carbon. The decrease in their values for T and B isomers is mainly due to the much weaker interaction between the bridged Ir atoms and the surface oxygen. In the case of T structure the interaction is with one of the bridging O atoms, while for B isomer it is with one basal surface O atom¹⁰ (see Fig.5.12). Somewhat bigger total cluster binding energy of S isomer with the C adatom than for the pristine mostly come from different surface relaxations. Chemical bonding of the cluster with the bridging O atoms is not affected since the orbitals involved in the C adsorption are oriented orthogonal to the Ir–O(b) bonds. Different surface relaxations are due to different "polarizability" of the Ir₄C cluster¹¹. Less charge is accumulated in the interfacial region which induces smaller displacements of both Ti(5c) and O(ba) atoms located below the cluster. The differences in the response to the surface field of B and S isomers upon C adsorption are reflected in the smaller values of the induced dipole moment than for the pristine clusters.

When compared with experimental results (Tab.5.6) the calculated structural data for the three isomers with the C adatom do not agree as well as the structural parameters of the pristine clusters. Although the Ir–Ir distances of the most stable T geometry increase upon the C adsorption the average first shell Ir–Ir coordination number decreases from 3.0 for the pristine cluster (Tab.5.6) to 2.5. As already discussed for the pristine clusters (Sec.5.6) the experiments are performed using the TiO₂ powders with the bigger fraction of the anatase phase and therefore, direct comparisons of the other coordination shells is not appropriate thing to do.

The most stable adsorption geometries of a single CO molecule on the three TiO₂ supported Ir₄ isomers are also displayed in Fig.6.9 and their properties presented in Tab.6.4. On the TiO₂ surface none of the three isomers favors the CO dissociation. The molecular CO adsorption is by 1.11, 1.17 and 1.16 eV more favorable than the dissociative adsorption on T, B and S structure, respectively. Moreover, the most stable Ir₄CO structure is S that is by 0.28 and 0.39 eV lower in energy than the other two (T and B, respectively). This is due to considerably lower CO binding energies on both T and B isomers than in the gas–phase. These changes as well as the changes in the CO adsorption configurations are due to changes in the electronic structures of the three isomers that are already discussed. Namely, to all three structures the CO molecule binds in a bridging not terminal configurations. However, the adsorption of a single CO molecule does not favor the CO dissociation as found for T

¹⁰Notice the lower first shell Ir–O coordination for B isomer (Tab.6.4) than for the adsorbed pristine cluster (Tab.5.6)
¹¹Strictly speaking the polarizability is defined as the linear response to the applied field. Here, I use this term to describe the response (not necessarily liner) of the three clusters to the surface electrostatic field which is too strong for any linear response treatment of the properties of the three isomers.
structure on MgO(100) and the C adatoms are not likely to be present on any of the three structures.

6.4 Influence of the Environment on the Properties of Ir$_4$ Clusters

After properties of Ir$_4$ clusters are being examined in three different environments, the gas–phase and supported by two different substrates MgO(100) and TiO$_2$(110), it is possible to conclude that the influence of the support is two–fold. Firstly, if the properties of the certain cluster geometry are followed, depending on the surface the electronic structure of the supported cluster can be more or less affected due to the electronic interactions with the surface atoms. These modifications of the electronic structure of the adsorbates are reflected in the change of their interactions with atomic or molecular species. This can be seen in Fig.6.10. Binding energies of the C adatom as well as the preference towards the CO dissociative adsorption depend strongly on the environment. At variance to the TiO$_2$ surface the influence of the clean MgO substrate on the properties of the supported Ir$_4$ clusters is

![Figure 6.10: Dependence of the properties of the three Ir$_4$ isomers (T, B and S) on the environment (GP refers to the gas–phase). Their energies relative to T configuration $E_{\text{rel.}}$, the binding energies of the C adatom $E_b(C)$ and the difference of the CO binding energies for the dissociative and molecular CO adsorption $\Delta E_b$(CO) (negative values mean that the dissociative adsorption is more favorable).](image-url)
not that strong. The binding energies of C adatoms as well as the adsorbed CO molecules on the TiO$_2$ supported Ir$_4$ clusters are considerably different than in the gas–phase and on the MgO as well. Moreover, the preference for the CO dissociation reaction accounted in the gas–phase is different for the supported Ir$_4$.

The second and, what has been proved in this thesis, equally important type of influence of the support on the adsorbed species is the ability to change the relative energies of the different isomers and to influence the geometry of the clusters not only through the relaxation effects. As shown in this thesis for the defected MgO(100) surface as well as for the defect–free TiO$_2$ surface the relative stability of different isomers can be considerably altered upon adsorption. In the gas–phase T isomer is as much as 1.5 eV less stable than S structure and normally it would be considered as irrelevant in any further investigation. However, it might become relevant depending on the chemical environment. Since the electronic structure of small clusters strongly depend on their atomic configuration this is another way how the substrate may influence electronic structure of the supported species.

Similarly to the effects of the support, the interaction with atomic or molecular species could also affect the relative energies of different isomers. It is shown in this thesis that a single C adatom (neither H nor C) changes the energy ordering of the three supported Ir$_4$ isomers in favor of T isomer bringing theoretical results in a better accordance with the experimental ones for the MgO(100) supported Ir$_4$. The origin of the carbon lays in the CO ligands which are present on the clusters during the preparation procedure. The CO dissociation is as favorable as the competing process CO desorption on the MgO supported T isomer.

Therefore, this thesis can be concluded with the statement that the properties of small transition metal clusters, Ir$_4$ in particular, greatly depend on the surrounding conditions. Understanding the relations between atomic and electronic structure of different isomers for a given cluster size, and their chemical environment, enables the control of their properties which is the ultimate goal for the applications.
Chapter 7

Conclusions

A theoretical study of the properties of Ir$_4$ clusters both in the gas–phase and when adsorbed on two commonly used oxide substrates MgO(100) and TiO$_2$(110), has been carried out. Theoretical tools based on density functional theory (DFT) within the generalized gradient approximation (GGA) and ultrasoft pseudopotentials have been used. Theoretical models are extensively tested and their applicability to studied problems verified.

Special emphasis is put on the relative stability of different Ir$_4$ isomers. The importance of this issue is, according to the opinion of the author of these lines, underestimated. The lowest total energy configurations are of course, the most relevant since they are most probably realized in experiments. However, which geometry will be the most stable one in a given situation depends both on the gas–phase relative stability and experimental conditions.

Properties of the gas–phase structures laying higher in energy are presented in the scientific publications mainly as a proof that the search for a global minimum is conducted carefully. Only in the cases of discrepancy with experiments (as for Ir$_4$) or where several nearly degenerate configurations are predicted, properties of the low energy structures are discussed in more details. It is shown in this thesis that the configurations which lay very high in energy\(^1\) and which would normally be treated as completely irrelevant may become relevant due to the interactions with the environment.

At the beginning, gas–phase Ir$_n$ ($n = 1, \ldots, 8$) clusters have been studied. Resulting energy orderings of different isomers for a fixed $n$ are in a relatively good accordance with other theoretical results treating gas–phase Ir clusters. However, variations in calculated relative energies between different isomers and, in some cases different ordering of the low energy structures can be found in the literature (see discussion in Chap.3). Agreement between different studies is much better when the geometries of the most stable configurations are compared.

In this thesis it is shown that the discrepancies in stability of different isomers are mainly due to different level of hybridization of atomic orbitals yielded by different theoretical models (Chap.3). The Ir$_4$ clusters are used for studying bonding mechanism in iridium clusters and it has been found that the relative stability of different isomers depends strongly on the level of 5$d$–6$s$ hybridization. This is also confirmed for other transition metal tetramers. The differences in published results are due to the fact that the level of hybridization is very sensitive on the approximations used in computations\(^2\). However, predicted energy ordering

\(^1\)Compared to the thermal energies at room temperature (~25 meV).

\(^2\)More precisely mostly on the exchange and correlation functional used in computations.
for the three relevant Ir$_4$ isomers is proved to be robust.

The gas–phase Ir$_4$ isomer with the lowest total energy is the square (S) structure that is as much as 0.93 and 1.51 eV more stable than the butterfly (B) and tetrahedron (T), respectively. This result is in contradiction with experimental findings which indicate that the oxide supported Ir$_4$ clusters adopt tetrahedral configuration.

Upon adsorption on the MgO(100) surface the S isomer remains the most stable Ir$_4$ structure, well separated in energy from the other two (Chap.5). Moreover, the tetrahedron is heavily distorted by the interaction with the surface oxygen. Presence of point defects (neutral and charged O vacancies) affects the energy ordering making tetrahedron and square very close in energy, but structural distortion of the tetrahedron are even bigger and the predicted data do not correspond to experiments. Diffusion of the S cluster on MgO(100) surface has also been studied. Two diffusion mechanisms are found the cluster "walk" and diffusion through structural transformations to B and T structures. It is estimated, based on the magnitudes of the calculated energy barriers and typical vibrational frequencies of the S structure, that temperatures of about 400° C could trigger the diffusion. This result agrees well with experimental findings.

On TiO$_2$(110) the T and S structures become degenerate and B becomes the least stable isomer (Chap.5). The structural distortions of the three isomers are small which is in accord with the experimental data. It is shown that the TiO$_2$ surface influences relative stability of the three isomers through a particularly strong electrostatic field acting on the adsorbates. This field is probably caused by a large anomaly in effective charges which is known to exist for bulk rutile TiO$_2$.

In addition to the oxide substrates interactions with H, C and O atoms as well as with CO molecules have been studied (Chap.4 and Chap.6). In all environments adsorption of a single C atom strongly influences the stability of the three isomers. Upon C adsorption, B becomes the most stable gas–phase isomer while on both MgO(100) and TiO$_2$(110) surfaces T is the most probable structure. Due to the big mass difference between carbon and iridium atoms, purely carbon vibrational modes appear in the spectra. Calculated carbon frequencies on T structure in the gas–phase (the MgO supported) are 103.5 and 87.6 meV (113.2 and 87.6 meV). These characteristic carbon frequencies offer a possibility for the experimental verification of the carbon presence. Adsorption of a single H or O atom does not produce the same effect as the C adsorption regardless of the environment.

Interaction with CO molecules is also important, given the experimental procedure for producing supported Ir$_4$ clusters out of Ir$_4$(CO)$_{12}$ carbonyl molecules. The dissociative CO adsorption is more favorable than the molecular adsorption on the gas–phase T and B isomers. On MgO(100) the dissociative CO adsorption is energetically more favorable only on T structure while on TiO$_2$(110) molecular CO binds stronger to all three isomers. Barrier for CO dissociation on T cluster supported by MgO(100), although quite high (2.97 eV), indicates that this reaction is as probable as the competing process the CO desorption (requiring 2.98 eV). Since the decarbonylation of Ir$_4$(CO)$_{12}$ is conducted in the nonequilibrium conditions (flowing He gas) the CO ligands could dissociate leaving carbon adatoms on Ir$_4$ clusters.

All these results indicate that chemical environment influences significantly both atomic and electronic structure of the supported Ir$_4$ clusters. Therefore, properties of the clusters

---

3In the bulk rutile TiO$_2$ the effective charges on Ti and O ions are considerably different from their formal +4 and -2 values. They lay, depending on the direction, in the range 5.20–7.70 and -1.35– -4.82 electrons, respectively [129].
cannot be treated independently on their surrounding. Knowledge of the relations between clusters and their environment is the key factor in engineering better catalysts based on oxide–supported small metallic particles.

There are also several issues that remain open. The first is related to the theoretical modeling of transition metal clusters. Although DFT is the method of choice it remains open which approximation to DFT (or which XC functional) describes the best properties of transition metals clusters. In these kind of investigations one usually faces a lack of experimental data for well characterized clusters in the gas–phase or deposited on well characterized substrates. In these cases, the results obtained with more accurate CI calculations are typically used as the benchmarks for DFT models. However, due to very high demands for computational power these results are available only for a number of systems typically very small in size (usually only for dimers).

Second, it is not yet clear what are the catalytically active, Ir$_4$ based species found in experiments. Again, well characterized samples are needed. This is necessary in order to be able to reach better understanding of the role of Ir$_4$ in chemical reactions. Without unified interpretation of the results coming both from experiments performed in controlled conditions and the theory, any theoretical modeling of processes such as chemical reactions on supported transition metal clusters remains mostly in the realm of the imagination of a theoretician.

Furthermore, none of the results, neither experimental nor theoretical, treating Ir$_6$ clusters have been presented here. In experiments [119] deposited Ir$_6$ clusters are prepared also using the Ir$_4$(CO)$_{12}$ as a precursor. Additional chemical treatments are then used, and from the EXAFS data it may concluded that the resulting supported species are octahedral Ir$_6$. Theory predicts trigonal prism as the most stable Ir$_6$ structure (Chap.4). The same result is obtained for the MgO supported species. Moreover, none of the studied adsorbates affects the stability of Ir$_6$ isomers in the same way like for Ir$_4$.

Finally, the issue of the electrostatic field of the TiO$_2$(110) surface needs to be further investigated as well as its influence on adsorbates. While it can be assumed that this field originates from the large anomaly in effective charges of bulk rutile TiO$_2$ there are no indications in the literature what are the effective charges of atoms at the surface.
Appendix A

Extended X-Ray Absorption Fine Structure (EXAFS) Spectroscopy

Extended X-Ray absorption fine structure (EXAFS) refers to the oscillatory behavior of the X-ray absorption as a function of photon energy beyond an absorption edge. This definition of EXAFS, taken from reference [133] describes the best what is EXAFS. The fine structure in the absorption coefficient $\mu$ is shown in Fig.A.1. It typically displays oscillatory behavior as a function of the incident photon energy. This effect is absent for an isolated atom and depends exclusively on the chemical environment, in particular on the number and type of neighbors as well as interatomic distances. This effect is widely used as an experimental tool for studying atomic structure of various systems (solids, molecules, liquids,...). The EXAFS spectroscopy is a very sophisticated tool and its development is greatly due to the availability of the synchrotron radiation which improved substantially the quality of EXAFS data. It is nowadays established as a powerful technique for structural studies. EXAFS is very useful for systems where single crystal X-ray diffraction is not applicable.

![Figure A.1](image)

**Figure A.1:** (a) Schematic figure of a typical X-ray absorption spectrum. (b) EXAFS above the Cu K-edge. Both absorption coefficient $\mu$ and and normalized oscillatory part $\chi$ (Eq.A.2) are shown. The values of $\chi$ are multiplied by 3. Both figures (a) and (b) are taken from Ref.[134].

The fine structure in the X-ray absorption spectra may extend up to 1000 eV above the edge and may have an amplitude in range of 1–20 % of the edge jump [133]. The explanation for this effect lays in the interference of the outgoing photoelectron wave with...
the wave backscattered from the neighbouring atoms. In fact, if the absorbing atom has a neighbouring atom the outgoing photoelectron wave will be backscattered by the neighbouring atom. The final state is then the sum of the outgoing and all the backscattered waves from each neighbouring atom.

For an isolated atom a photoelectron emitted due to the X-ray absorption will travel as a spherical wave with the wave vector

$$k = \sqrt{\frac{2m}{\hbar^2}}(E - E_0),$$

where $E$ is the photon energy and $E_0$ is the absorption edge energy. The EXAFS is usually described in terms of oscillatory part $\chi$ of the absorption coefficient $\mu$ normalized to the averaged absorption $\mu_0$

$$\chi(E) = \frac{\mu(E) - \mu_0(E)}{\mu_0(E)}.$$ (A.2)

Standard way is to express $\chi$ as a function of the wave vector using Eq.A.1 and to write it as a sum

$$\chi = \sum_j N_j S_j(k) F_j(k) e^{-2\sigma_j^2 k^2} e^{-2r_j/\lambda_j(k)} \frac{\sin(2kr_j + \phi_{ij}(k))}{kr_j^2},$$ (A.3)

where $N_j$ is the number of neighboring atoms of the $j$th type at the distance $r_j$ with the backscattering amplitude $F_j(k)$. Thermal and static disorders are described by the Debye–Waller factor $\sigma_j$, while the term $e^{-2r_j/\lambda_j(k)}$ is due to inelastic losses in the scattering process with the mean free path $\lambda_j$. $S_i(k)$ is the amplitude reduction factor due to many–body effects at the central atom and $\phi_{ij}(k)$ is the total phase shift experienced by the photoelectron which depends on both absorber and the backscatterer. Since $\phi_{ij}(k)$ depends on $k$ it also influences the frequency of the sinusoidal function.

In practice the results of EXAFS spectroscopy depend on the feasibility of resolving the measured spectra into individual waves that correspond to different type of neighbors of the absorbing atom. This is typically done by two methods, curve–fitting or Fourier transform technique. The first method is based on a best fitting of the data with a modeled sum of individual waves. The other method uses Fourier transformation of the data into a real space where the $k$–space wave behavior is localized into separated peaks. Positions of those peaks contain the information about the distances between the absorber and the neighbors while the sizes of the peaks are related to the numbers of and types of the neighboring atoms.
Appendix B

Character Tables of $D_{\infty h}$, $C_{2v}$, $D_{4h}$ and $T_d$ Point Groups

Character tables of $D_{\infty h}$, $C_{2v}$, $D_{4h}$ and $T_d$ groups which are important for this study are presented here. Labeling of their irreducible representations is done according to Ref.[135]. Each table is preceded by a figure where the group transformations are presented pictorially on a ball and stick models representing Ir$_4$ clusters.

In addition to the characters of the irreducible representations the characters of the representations in the subspaces that are spanned over the atomic orbitals are shown. These representations are formed by grouping the atomic orbitals sitting on each atom in the irreducible representations of the small group\(^1\) of a given atom. Orbitals belonging to the same irreducible representations of the small groups of equivalent atoms are also equivalent and form an invariant (but reducible) subspace. When these subspaces are reduced their basis vectors form the symmetry adapted basis set which is convenient for expanding molecular orbitals. Furthermore, it provides a natural way for the analysis of possible hybridizations. Reduction of these invariant subspaces to the irreducible representations of the corresponding group is also presented next to characters of the corresponding (reducible) representations.

In each figure a coordinate frame is assumed. Since all groups except $T_d$ are axial groups the principal axis is always set to be the $z$-axis while $x$ and $y$ lay in the mirror planes. The atomic orbitals of a tetrahedron are labeled according to the local coordinate frames. Each atom has its own frame with the $z$ axis pointing from the atom to the center of the tetrahedron. The $x$ and $y$ are chosen in such a way to be connected with other local $x$ and $y$ by a group transformations that connect the corresponding atoms.

\(^1\)The subgroup of the symmetry group that leaves the given atom invariant.
Upon reducing symmetry from $D_{\infty h}$ to $D_{4h}$:

- $A_{1g} \Rightarrow A_{1g}$
- $A_{1u} \Rightarrow A_{2u}$
- $E_{1g} \Rightarrow E_g$
- $E_{1u} \Rightarrow E_u$
- $E_{2g} \Rightarrow B_{1g}+B_{2g}$
- $E_{2u} \Rightarrow B_{1u}+B_{2u}$
Superscripts 1 and 2 of the symbols of atomic orbitals refer to the two types of nonequivalent (by symmetry) atoms in the butterfly structure, the wingtip and body atoms.

<table>
<thead>
<tr>
<th>$C_{2v}$</th>
<th>E</th>
<th>$C_2$</th>
<th>$\sigma_v$</th>
<th>$\sigma_{v'}$</th>
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<tbody>
<tr>
<td>$A_1$</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
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<tr>
<td>$B_1$</td>
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<td>1</td>
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<td>-1</td>
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<td>1</td>
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- $s^{1/2}$: 2 0 2/0 0/2 $\Rightarrow A_1 + B_{1/2}$
- $d_{x^2}^{1/2}$: 2 0 2/0 0/2 $\Rightarrow A_1 + B_{1/2}$
- $d_{x^2-y^2}^{1/2}$: 2 0 2/0 0/2 $\Rightarrow A_1 + B_{1/2}$
- $d_{xy}^{1/2}$: 2 0 -2/0 0/-2 $\Rightarrow A_2 + B_{2/1}$
- $d_{xz}^{1/2}$: 2 0 2/0 0/2 $\Rightarrow A_1 + B_{1/2}$
- $d_{yz}^{1/2}$: 2 0 -2/0 0/-2 $\Rightarrow A_2 + B_{2/1}$
- $p_z^{1/2}$: 2 0 2/0 0/2 $\Rightarrow A_1 + B_{1/2}$
- $p_x^{1/2}$: 2 0 2/0 0/2 $\Rightarrow A_1 + B_{1/2}$
- $p_y^{1/2}$: 2 0 -2/0 0/-2 $\Rightarrow A_2 + B_{2/1}$
\[ D_{4h} \begin{array}{c|cccccccc} \text{E} & 2C_4 & C_2 & 2C_2' & 2C_2'' & i & 2S_4 & \sigma_h & \sigma_v \\ \hline \text{A}_{1g} & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\ \text{A}_{2g} & 1 & 1 & 1 & -1 & -1 & 1 & 1 & 1 \\ \text{B}_{1g} & 1 & -1 & 1 & 1 & -1 & 1 & 1 & -1 \\ \text{B}_{2g} & 1 & -1 & 1 & -1 & 1 & -1 & 1 & 1 \\ \text{E}_g & 2 & 0 & -2 & 0 & 0 & 2 & 0 & -2 \\ \text{A}_{1u} & 1 & 1 & 1 & 1 & -1 & -1 & -1 & -1 \\ \text{A}_{2u} & 1 & 1 & 1 & -1 & -1 & -1 & -1 & 1 \\ \text{B}_{1u} & 1 & -1 & 1 & 1 & -1 & 1 & 1 & 1 \\ \text{B}_{2u} & 1 & -1 & 1 & -1 & 1 & 1 & 1 & -1 \\ \text{E}_u & 2 & 0 & -2 & 0 & 0 & -2 & 0 & 2 \\ \end{array} \]
\[
\begin{array}{c|cccc}
T_d & E & 8C_3 & 3C_2 & 6S_4 & 6\sigma_d \\
\hline
A_1 & 1 & 1 & 1 & 1 & 1 \\
A_2 & 1 & 1 & 1 & -1 & -1 \\
E & 2 & -1 & 2 & 0 & 0 \\
T_1 & 3 & 0 & -1 & 1 & -1 \\
T_2 & 3 & 0 & -1 & -1 & 1 \\
\end{array}
\]

\[
\begin{array}{c|cccc}
s & d_{z^2} & d_{xy}, d_{x^2-y^2} & d_{xz}, d_{yz} & p_z & p_x, p_y \\
4 & 1 & 0 & 0 & 2 & 0 & 0 & \Rightarrow A_1+T_2 \\
4 & 1 & 0 & 0 & 2 & 0 & 0 & \Rightarrow A_1+T_2 \\
8 & -1 & 0 & 0 & 0 & 0 & 0 & \Rightarrow E+T_1+T_2 \\
8 & -1 & 0 & 0 & 0 & 0 & 0 & \Rightarrow E+T_1+T_2 \\
4 & 1 & 0 & 0 & 2 & 0 & 0 & \Rightarrow A_1+T_2 \\
8 & -1 & 0 & 0 & 0 & 0 & 0 & \Rightarrow E+T_1+T_2 \\
\end{array}
\]
Bibliography


[35] Quantum-ESPRESSO is a community project for high-quality quantum-simulation software, based on density-functional theory, and coordinated by Paolo Giannozzi. See http://www.quantum-espresso.org


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# Curriculum Vitae

<table>
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<tr>
<th>Name and surname</th>
<th>Vladan Stevanović</th>
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<tbody>
<tr>
<td>Date of birth</td>
<td>October 28th, 1975</td>
</tr>
<tr>
<td>Place of birth</td>
<td>Bor, Serbia</td>
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<tr>
<td>Marital status</td>
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2005–2009  Doctoral Assistant at the *Institut Romand de Recherche Numérique en Physique des Matériaux* (IRRMA), EPFL.  
Supervisor: Prof. Alfonso Baldereschi

2007–2009  Supervision (with Prof. A. Baldereschi) of a master student Jean–René Moret during both practical work and the master project.

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