CHEMICAL ENVIRONMENT OF WATER MOLECULES ADSORBED ON THE ANATASE SURFACE: QUANTUM CHEMICAL STUDY

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Theoretical studies of the interactions of an oxygen molecule with the surface of a solid phase are of great importance for understanding the mechanisms of reactions involving oxygen on a solid surface. In this work, the spatial and electronic structure of oxygen defects and nitrogen impurity centers of the anatase surface and their manifestation in water adsorption are investigated. The anatase surface was simulated for by clusters of the composition Ti_{14}H_{22}O_{39} (defect-free face (001)), Ti_{14}H_{22}O_{38} (surface with oxygen vacancy), and Ti_{14}H_{22}N_{2}O_{36} (surface with both oxygen vacancy and incorporated nitrogen), terminated with boundary hydrogen atoms.

Calculations on the total energy of the optimized geometric structures of the model TiO_2 clusters and on the corresponding theoretical XPS spectra were performed using the density functional theory (DFT) method and the B3LYP hybrid functional with an extended valence-split basis set 6-31G (d, p).

In the XPS spectra of O1s, each peak can be attributed to a specific type of oxygen atoms in the initial structure, dependent on their coordination environment. The substitution of nitrogen atoms for oxygen ones leads to a complication of the spectrum. Simultaneous replacement of nitrogen atoms for oxygen ones and presence of an oxygen vacancy leads to further complication of the XPS spectrum.

The different structure of the adsorption complexes of water molecules on the anatase surface formed due to the HOH···O hydrogen bond (this displaces by 0.05 eV all the peaks in the XPS spectrum) or the coordination bond Ti···OH_2 has been also considered.

Based on the analysis of theoretical results, the role of various forms of defects in the adsorption of water on the anatase surface is considered.

The results of quantum chemical studies of molecular models simulated for the bulk and surface of titanium dioxide are compared with the literature data available.

Keywords: anatase, oxygen defect, nitrogen impurity, water adsorption, XPS spectra, quantum chemical density functional theory (DFT) method, cluster models

INTRODUCTION

Interest has grown significantly to the theoretical studies on the interaction of an oxygen molecule, its ground state being a triplet diradical, with a solid-phase surface. The results of such studies are of fundamental importance for both adsorption and catalysis, for the general theory of corrosion of metals (they let it possible at least to slow down the corresponding reactions). Of great importance are studies on the interaction of an O_2 molecule with a widely used semiconductor material, namely titanium dioxide; its forbidden band makes it possible to use only a relatively small range of ultraviolet radiation from the sun. The formation of nonstoichiometric areas on the TiO_2 surface, due to impurity centers, removal of oxygen atoms, and adsorption of atomic and molecular oxygen, makes it possible to reduce the width of the forbidden band of TiO_2 prior to its shift to the visible range of electromagnetic radiation.

The photochemistry of the ultraviolet absorption region of molecular oxygen O_2 is of interest for chemical physics, photobiology, and many other scientific fields [1].

Molecular oxygen very weakly absorbs radiation throughout the entire spectral range - from infrared to ultraviolet, but the presence and influence of the molecular environment make possible the existence of molecules of singlet oxygen O_2 (^1Σ_g), the formation of which is forbidden during photoexcitation of "isolated" O_2 molecules. O_2 molecules, due to their high reactivity, play a key role in natural photobiological and photochemical processes; therefore, a detailed study on the mechanisms of its formation is believed to be a fight-to-date scientific research task. Thus, examining the
formation of singlet oxygen upon photoexcitation of a weakly bound \( X \cdot O_2 \) (\( X \) - active surface sites) complex is not only fundamental, but also of practical interest, since this process is a new way of generating \( O_2 \) molecules. This process provides a new mechanism for photo-oxidation of various molecules in a condensed medium, and can also be used in UV air purification, in photocatalysis, photobiology, and medicine. Theoretical study on the processes of interaction between molecular oxygen and surface, for example, that of TiO\(_2\), is associated with the corresponding complications caused by the singlet state of both the surface itself and respective adsorption complex. There is a need to expand the generally accepted cluster model (CM) of adsorption, which involves a theoretical calculation of the cross section of only one potential energy surface (PES), which, as a rule, corresponds to the singlet ground electronic state. In the case, when the \( O_2 \) molecule reacts in the triplet state, it becomes necessary to calculate the cross section of another PES, which characterizes the triplet state of the cluster consisting of an \( O_2 \) molecule and a fragment of both surface and adjacent bulk phase of the TiO\(_2\) crystal.

Titanium dioxide belongs to the class of transition metal oxides, in nature it occurs with the structures of anatase (tetragonal), brookite (rhombic), rutile (tetragonal), less often with monoclinic TiO\(_2\) structure. Differences in the crystal lattices of titanium dioxide cause differences in density, mass and in the electronic band structure of titanium dioxide. The width of the band gap for the anatase structure is 3.2 eV, and for rutile – 3.0 eV [2].

The photocatalytic activity of titania nanoparticles was found to increase with a decrease in their diameter for both anatase and rutile modifications. When measuring their absorption spectra, an increase in their diameter was shown to shift the maximum of the absorption spectrum from the ultraviolet region to the long-wavelength part of the electromagnetic scale [3].

In previous article [4], the spatial structure and electronic structure of defect and impurity centers of polymorphous modifications of titanium dioxide and their manifestation in adsorption, catalytic and photocatalytic processes were analyzed. Here, we analyze the densities of states of oxygen atoms in titania matrices (parent and containing impurities of nitrogen and/or oxygen vacancies).

**METHODS**

The aim of the studies was to calculate the density of states of cluster models for the anatase surface (pure, nitrogen-substituted, and containing oxygen defects). Theoretical densities of states can be compared with the experimental spectra of XPS. As a result, we can determine the effect of the local environment of oxygen atoms on the chemical shift relative to their structurally nonequivalent crystallographic positions [4].

The H-terminated clusters that simulate the surface of an ideal anatase crystal, as well as the surface with an oxygen vacancy on the face (001) were considered. The defect-free face (001) of anatase was modeled by the cluster of the gross formula Ti\(_{14}H_{22}O_{39}\) which contains 7 surface five-fold coordinated titanium atoms. The presence of an oxygen vacancy was reproduced by the cluster Ti\(_{14}H_{22}O_{38}\), which is obtained from the former by removing of a two-fold coordinated surface oxygen atom as well as the surface with both oxygen vacancy and incorporated nitrogen was mimicked by Ti\(_{14}H_{22}N_{2}O_{36}\) cluster model.

Quantum chemical calculations were carried out within the frameworks of the so-called cluster approximation [5]. Computations on the total energy values of the optimized geometrical structures of TiO\(_2\) cluster models including 14 titanium atoms (Ti\(_{14}H_{22}O_{39}\)) and of relative models containing impurities of nitrogen and/or oxygen vacancies were performed using density functional theory (DFT) method [6] and the hybrid B3LYP [7] functional with extended valence-split basis set 6-31G(d,p). Calculations were carried out by means of the software package PC GAMESS [8] (version Firefly 8.1.0 by A. Granovsky, http://classic.chem.msu.su/gran/firefly/index.html). The densities of states were designed based on the quantum chemical results with use of the Spectra Builder 2.4 program package [9] earlier used by us in the article [10].

**RESULTS AND DISCUSSION**

The experimental XPS spectra of initial (pure) titania matrix and of related oxygen-vacancy and/or of nitrogen-impurity samples were examined and discussed in work [4]. Here,
let us make an attempt to evaluate interrelations between the model structures and their calculated XPS spectra (Figs. 1–4).

All the oxygen atoms in the parent structure (Fig. 1) can be divided into some groups according to their coordination environment (2- and 3-coordinated) and to chemical nature of the second and third neighbors. Five O1s peaks one can observe in the XPS spectrum calculated for this cluster model (near 521 eV). Every peak can be assigned to a definite type of O atoms depicted in the inset in Fig. 1.

Elimination of an oxygen atom from the titania matrix results in a divergence between 2-coordinated and 3-coordinated oxygen atoms (the split peak at 521.44 and 521.28 eV respectively, see Fig. 2).

Dinitrogen substitution for oxygen atoms results in the complication of the spectrum (at least 12 peaks can be observed). Nevertheless, 2 new peaks appear near 523 eV (522.87 and 523.28 eV) related to 3-coordinated oxygen atoms (Fig. 3).

Simultaneous dinitrogen substitution for oxygen atoms and oxygen vacancy leads to further complication of the XPS spectrum (15 peaks can be distinguished, including a pair near 523 eV, namely 522.60 and 522.74 eV, see Fig. 4).

The different structure of water molecule adsorption complexes have been considered, one of them being due to formation of a hydrogen bond HOH···O whereas another is characterized by a coordination bond Ti···OH₂ (Fig. 5).

In the first case, a shift for approximately 0.05 eV takes place of all the peaks in the XPS spectrum calculated for the adsorption complex though the spectrum is somewhat broadened and as a whole keeps its image.
**Fig. 2.** Cluster model and calculation XPS spectra for structure with oxygen vacancy Ti$_{14}$H$_{22}$O$_{38}$

**Fig. 3.** Cluster model and calculation XPS spectra for structure with dinitrogen substitution for oxygen atoms Ti$_{14}$N$_2$H$_{22}$O$_{37}$
Fig. 4. Cluster model and calculation XPS spectra for combined structure with simultaneous dinitrogen substitution for oxygen atoms and oxygen vacancy Ti$_4$N$_2$H$_2$O$_{36}$

Fig. 5. Cluster model and calculation XPS spectra for adsorption of water molecules on titania surfaces. $E_a = -36$ kJ/mol
CONCLUSIONS

An analysis of the results of quantum chemical calculations gives evidence that the chemical environment of water molecules adsorbed on anatase surface is reflected in experimental XPS spectra. The adsorption of water molecules leads to a bathochromic shift of all the peaks O1s in the spectra. The presence of oxygen defects and nitrogen impurities results in complication of the spectra and in appearance of O1s double peaks.

Идентификация химического окружения молекул воды, адсорбированных на поверхности анатаза: квантовохимические расчёты

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Теоретичні дослідження взаємодії молекул відповідних спектрах РФЕС виконані з використанням методу теорії функціоналу густини (DFT) і гібридного функціоналу B3LYP з розрахунком валентно-розділенням базисним набором 6-31G (d, p).

У РФЕС-спектрах O1s кожен пік може бути віднесенний до певного типу атомів відповідно до координаційного числа від окисних дефектів в адсорбції води. Поверхня анатаза моделювалась кластерами складу Ti2H2O39 (бездефектна грань (001)), Ti14H22O36 (поверхня з кисневою вакансією) та Ti14H22N2O36 (поверхня з кисневою вакансією і впровадженими атомами азоту), які обмежуються атомами води.

Розрахунки величин повної енергії оптимізованих просторових структур моделей кластерів TiO2 і відповідних теоретичних спектрів РФЕС виконані з використанням методу теорії функціоналу густини (DFT) і гібридного функціоналу B3LYP з розрахунком валентно-розділенням базисним набором 6-31G (d, p).

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Розглянутые также рёзультаты розглянутого метода адсорбционных комплексов молекул воды на поверхности анатаза, утвержденных заведующему водным звёздом NOH-O (при котором в спектре РФЕС вдубчаются звёзд на 0.05 eV всех пиков), або координационной звёзды Ti-OH2.

На підставі аналізу теоретичних результатів розглянуто роль різних видів дефектів в адсорбції води на поверхні анатаза. Теоретично одержані результати порівнюються з властивостями молекулярних моделей для обліку і поверхні доксії титану, які існують у літературі.

Ключові слова: анатаз, спектри РФЕС, діоксид титану, метод теорії функціоналу електронної густини, кластерні моделі
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моделировалась кластерами состава Ti₄H₂₂O₃₀ (бездефектная грань (001)), Ti₄H₂₂O₃₀ (поверхность с кислородной вакансией) и Ti₄H₂₂N₁₂O₃₀ (поверхность с кислородной вакансией и внедренными атомами азота), которые ограничены атомами водорода.

Расчеты величин полной энергии оптимизированных пространственных структур модельных кластеров TiO₂ и соответствующих теоретических спектров РФЭС выполнены с использованием метода теории функционала плотности (DFT) и гибридного функционала B3LYP с расширенным валентно-расщепленным базисным набором 6-31G (d, p).

В РФЭС-спектрах O1s каждый пик может быть отнесен к определенному типу атомов кислорода в зависимости от их координационного окружения. Замена атомами азота атомов кислорода приводит к усилению спектра. Одновременное замещение атомами азота атомов кислорода и наличие кислородной ваканси приводит к дальнейшему усилению спектра РФЭС.

Рассмотрено также различное строение адсорбционных комплексов молекул воды на поверхности анатаза, образованных благодаря водородной связи HOH···O (при этом в спектре РФЭС происходит смещение на 0,05 эВ всех пиков), или координационной связи Ti···OH₂.

На основании анализа теоретических результатов рассмотрена роль различных видов дефектов в адсорбции воды на поверхности анатаза. Теоретически полученные результаты сравниваются со свойствами молекулярных моделей для объема и поверхности диоксида титана, имеющихся в литературе.

Ключевые слова: анатаз, спектры РФЭС, диоксид титана, метод теории функционала плотности, кластерные модели

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