Stoichiometry and disorder influence over electronic structure in nanostructured VO\textsubscript{x} films

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Abstract We present and discuss the role of nanoparticles size and stoichiometry over the local atomic environment of nanostructured VO\textsubscript{x} films. The samples have been characterized in situ using X-ray absorption near-edge structure (XANES) spectroscopy identifying the stoichiometry-dependent fingerprints of disordered atomic arrangement. In vanadium oxides, the ligand atoms arrange according to a distorted octahedral geometry depending on the oxidation state, e.g. trigonal distortion in V\textsubscript{2}O\textsubscript{3} and tetragonal distortion in bulk VO\textsubscript{2}. We demonstrate, taking VO\textsubscript{2} as a case study, that as a consequence of the nanometric size of the nanoparticles, the original ligands symmetry of the bulk is broken resulting in the coexistence of a continuum of distorted atomic conformations. The resulting modulation of the electronic structure of the nanostructured VO\textsubscript{x} as a function of the oxygen content reveals a stoichiometry-dependent increase of disorder in the ligands matrix. This work shows the possibility to produce VO\textsubscript{x} nanostructured films accessing new disordered phases and provides a unique tool to investigate the complex matter.

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Introduction

Vanadium oxides have been studied extensively due to their various possible stoichiometric phases characterized by different oxidation states, from V$^{+2}$ to V$^{+5}$ as well as mixed-valence oxides (Magneli and Wadsley series) (Schwingenschlögl and Eyert 2004; Chen et al. 2019) which provides an equally large variety of electronic and physico-chemical properties. For instance, V$_2$O$_5$ and V$_6$O$_{13}$ have been investigated for possible applications as cathode material in Li-ion batteries (Chan et al. 2007; Xu et al. 2016; Shi et al. 2018; Xiao and Zhang 2018; Tan et al. 2018). Vanadium oxides also bear the seed of a strong electronic correlation. VO$_2$, V$_2$O$_3$ and the mixed-valence V$_n$O$_{2n-1}$ ($n = 3$–$6$, 8 and 9) systems all exhibit first-order metal-insulator transitions (MIT) characterized by a nanoscale phase separation (Marcelli et al. 2017; McLeod et al. 2017), which can be exploited in the emerging field of Mottronics (Brahlek et al. 2017). Nevertheless, tuning the properties of vanadium oxides during the early synthesis stages is a difficult task, due to the complex interplay among stoichiometry, lattice order and size of the sample.

The control of the lattice order is one of the most used tools to modulate the electronic structure of matter (Fan et al. 2014; Pinto et al. 2018a; Rezvani et al. 2016a; Rezvani et al. 2018; D’Elia et al. 2020a). However, while methods to slightly modify the inter-atomic distances, e.g. application of epitaxial strain, are widely used, the development of a technique able to produce disordered samples with controllable stoichiometry is in its early stage despite its potential.

It has been shown that the amorphous VO$_x$ exhibits superior intercalation properties with respect to the crystalline counterpart due to the large number of the dangling bond within the defective structure revealing their technological appeal (Uchaker et al. 2014; Chae et al. 2014; Wu et al. 2018).

Disordered VO$_x$ could benefit from reducing their size to the nanoscale (Jagiello et al. 2017; Pasqualini et al. 2017; Sartale et al. 2013) because of confinement effects over the electronic structure, e.g. quantum confinement enhancement of the work function (D’Elia et al. 2020b; Zhou and Zachariah 2012), and because of a large effective surface area (Rezvani et al. 2016b; Pinto et al. 2016; Pinto et al. 2018b).

The possibility to produce disordered nanostructured (NS), i.e. assembled by nanoparticles, VO$_x$ films and to monitor in situ their local atomic arrangement is therefore of great interest.

In the present work, we present an in situ X-ray absorption near-edge structure (XANES) spectroscopy investigation of NS VO$_x$ with tunable stoichiometry.

XANES is associated with the electronic transitions from core shells to unoccupied states following a photo-absorption process, possessing the unique capability to probe concurrently the stoichiometry and local structure of the sample (Bachrach 1992; Bianconi 1982). The fabricated films have been characterized at distinct oxygen concentration stages monitoring the evolution of the electronic structure as a function of stoichiometry.

The fingerprints of a disordered ligands arrangement have been observed and are discussed comparing our data with those of a crystalline samples and with several simulations linking the coexistence of multiple randomly distributed atomic dislocations, with surface area and nanoparticle size.

Experimental

The NS VO$_x$ films were deposited using a combination of the Pulsed Micro-plasma Cluster Source (PMCS) and the Supersonic Cluster Beam Deposition (SCBD) method. SCBD is a well-known synthesis method able to produce cluster assembled NS materials (D’Elia et al. 2020b; Borghi et al. 2018; Barborini et al. 2011). This method offers considerable advantages for neutral cluster manipulation based on effusive beams making the seeded supersonic beam approach very powerful for the deposition of NS films. PMCS is a cluster source characterized by high deposition rate, control over cluster growth and landing energy of the cluster beam (Wegner et al. 2006; Piseri et al. 2004; Barborini et al. 1999). Combining SCBD and PMCS is an original approach to the preparation of disordered NS vanadium oxide films, which facilitated their in situ characterization by XANES spectroscopy.

The PMCS is a pulsed-cluster source driven by a high-power pulsed electric discharge. A vanadium cathode (6 mm diam. rod, purity 99.9%, EvoChem GmbH) was used to generate a seeded supersonic beam of
vanadium oxide clusters. To oxidize the nanoparticles in the early synthesis stage, an Ar-O2 gas mixture (high purity Ar: 99.9995%, SIAD) with a controlled quantity of oxygen was used. In the experimental layout, the supersonic cluster beam produced by the PMCS is deposited in situ on a copper substrate, which is electrically isolated from the rest of the experimental chamber. The base pressure of the experimental chamber was 1 × 10^-9 mbar, while operating the PMCS, raised it to ~5 × 10^-9 mbar due to the carrier gas injected by the cluster source.

To synthesize samples with different stoichiometry, we fed the PMCS with an Ar-O2 gas mixture with an O2 molar percentage in the range of 0–1.25%. The working parameters of the PMCS have been adjusted to maximize the deposition rate. The delay between the gas injection and the discharge firing was 0.51–0.64 ms; the discharge operating voltage was 0.9 kV, and the discharge duration was 60 µs; the pulsed-valve aperture was between 196 and 220 µs; and the pulse repetition rate was 3 Hz, and the Ar-O2 pressure feeding the Parker valve was 60 bar. All the samples have been synthesized using Ar or Ar-O2 gas mixture as specified elsewhere (D’Elia et al. 2020b). The substrate was kept at ~300 K during the deposition, and the deposited films have been studied in situ using XANES spectroscopy. The UHV chamber has been coupled to the PMCS, available at the TASC laboratory (De Simone et al. 2012) and equipped with an XYZ manipulator as depicted in Fig. 1. The XANES measurements have been performed at the Circular Polarization beamline of the Eleftra synchrotron radiation facility using the CryoAlp UHV experimental chamber (Marcelli et al. 2018). Total electron yield (TEY) spectra have been acquired measuring the drain current, connecting the sample to ground through a Keithley 486 picoammeter (see Fig. 1). The sample was kept normal to the cluster beam, while the angle between the incident photons and the sample surface was 45°.

Furthermore, ex situ XANES measurements on a crystalline sample, synthesized by molecular beam epitaxy (Fan et al. 2014), were performed using the IRMA experimental chamber (Sacchi et al. 2003) and used as a reference in our experiment. All the XANES spectra have been normalized to the incident photon flux and have been normalized to the incident photon flux and the sample surface was 45° normal to the cluster beam, while the angle between the incident photons and the sample surface was 45°.

Results and discussion

The morphology and size of nanoparticles deposited by SCBD have been investigated using TEM. A deposition of 1 s has been performed on a TEM grid with an amorphous C supporting film. Figure 2 shows the TEM image of the VO2.5 sample. Although the individual nanoparticles form large agglomerates, they remain distinctly visible in the magnified image. Despite the aggregation that hampers the size distribution assessment, the size of the nanoparticles can be estimated to be in the range 3–6 nm with an average value of ~4.4 nm.

No clear differences in the particle size distribution have been observed as a function of carrier gas and oxidation state.

The different NS VOx films have been deposited and investigated in situ using XANES spectroscopy at V L_2,3 and O K edges. All the deposited films have a thickness of at least 15 nm in order to avoid probing the Cu substrate. We identify the different VO_x samples using the stoichiometric ratio defined as x=[No. O atoms/No. V atoms], in the range 0 ≤ x ≤ 2.5. The XANES spectra for different VO_x samples are shown in Fig. 3.

The vanadium L-edges spectra show two major components at ~515.2 eV (a) and ~522.3 eV (b) (see left panel of Fig. 3). We observe a chemical shift towards the higher photon energies with the introduction of the oxygen into the structure for values as low as x = 0.3.

The O K-edge spectra of the VO_x samples (Fig. 3 right panel), reveal a significant evolution of the electronic structure with stoichiometry which can be understood in terms of crystal field splitting and metal-ligand hybridization. In VO_x, O atoms arrange in an octahedral geometry generating a cubic crystal field (Hermann and Witko 2001; De Groot et al. 1989), which splits the degenerate 3d manifold into t_{2g} and e_{g} levels. As shown in Fig. 3, the O K edge XANES exhibits two main contributions at ~530 and ~532.5 eV, which can be associated with t^*_2g and e^*_g states, respectively (De Groot et al. 1989). Their relative intensity shows a marked dependence on the oxidation state.

Since XANES probes the empty density of states (DOS), t^*_2g and e^*_g signals can be observed at the O K edge only when t_{2g} and e_{g} electrons are bound with the O 2p electrons. The presence of t^*_2g in VO_0.6 samples (the weak component around 530 eV) can be attributed to the hybridization of V-3d and O-2p electrons in agreement with a previous Auger investigation, which
predicts a mixed $4s - 3d$ valence band for this suboxide (D’Elia et al. 2020b). $e^*_{2g}$ and $t^*_{2g}$ features are observable on the whole VO$_x$ set, even though they are broad and not well separated.

The origin of $e^*_{2g}$ and $t^*_{2g}$ width and energy separation can be interpreted by comparison with a reference single crystal (SC) sample. We investigated VO$_2$ as a case study. The comparison of the SC-VO$_2$ and NS-VO$_2$ is reported in Fig. 4.

In the SC sample spectra, the $e^*_{2g}$ and $t^*_{2g}$ features are well separated pointing to a strong and homogeneous crystal field and therefore to a locally ordered atomic environment. The $e^*_{2g}$ and $t^*_{2g}$ features of the NS film, on the other hand, are broadened and less resolved. The $t^*_{2g}$
energy position seems unaffected within our experimental resolution, while the e\textsuperscript{\textast}g component is red-shifted of about 0.4 eV. Coherently, a red-shift of the L\textsubscript{3} energy position is observed in the NS sample with respect to the SC since V L edges are particularly sensitive to the e\textsuperscript{\textast}g contribution to the empty DOS (Zaanen et al. 1985; Abbate et al. 1991). The observed e\textsuperscript{\textast}g red-shift in the NS-VO\textsubscript{2} is possibly due to the presence of Jahn-Teller-like distortions of the oxygen octahedron matrix which directly influences the electronic structure of the NS sample (Kasatikov et al. 2019; Ruzmeto et al. 2007). In the extreme case, a high number (ideally infinite) of different ligand distortions coexist within a disordered material. The disorder considered here is of geometric nature; around each metal site, the local symmetry of the ligands arrangement is broken because of random atomic dislocations, i.e. displacements from their symmetry site. In this picture, each vanadium site is surrounded by a different distorted ligand conformation which leads to an overall disordered structure.

In order to confirm this observation, and since the e\textsuperscript{\textast}g orbitals are the most involved, we investigated V L edges line shape of the NS VO\textsubscript{2} sample (Zaanen et al. 1985; Kasatikov et al. 2019). Several simulations were carried out within the charge transfer multiplet framework to investigate possible distortions effect on the VO\textsubscript{2} electronic structures. The calculations were performed by the configuration interaction cluster for octahedral and distorted VO\textsubscript{2} structures (see Fig. 5). The refinement of the analysis of the distorted structures was performed, employing the CTM4XAS (Stavitski and de Groot 2010) simulation code.

According to previously published work (Ikeno et al. 2011), the calculations were based on a crystal field 10Dq=2.35 eV with 80% reduction of the Slater integrals and an average Coulomb interaction $U_{dd} - U_{pd}$ of 0.88 eV. Finally, an overall broadening of 0.35 eV full width was applied.

In VO\textsubscript{2} the oxygen atoms are arranged according to a tetragonally distorted octahedral geometry (D’Elia et al. 2020a, c). The tetragonal distortion leads to a further splitting of the $t_{2g}$ and e\textsuperscript{\textast}g orbitals resulting in the four levels e\textsuperscript{\textast}g, b\textsubscript{2g}, a\textsubscript{1g} and b\textsubscript{1g} (see Fig. 5 right panel). The e\textsuperscript{\textast}g, b\textsubscript{2g}, separation is called Q\textsubscript{1}, while the a\textsubscript{1g} and b\textsubscript{1g} are splitted by Q\textsubscript{2}. To mimic the ligand atomic
dislocations, e.g. the bending of two opposite octahedron vertices which red-shift the $e^*_{\text{g}}$, the calculations were performed for increasing values of $Q_2 = 0.2, 0.5, 0.8$ and $1.0$ eV while keeping $Q_1 = 0.2$ eV. The choice to keep the $Q_1$ constant is justified by considering the prominent sensitivity of V L edges to the $e^*_{\text{g}}$ orbitals which hide small spectral changes in the $t^*_{\text{2g}}$ energy region (Zaanen et al. 1985). We compare the simulated spectra with the V L$_{2,3}$ spectrum of NS VO$_2$ sample in Fig. 5.

The results demonstrate that the observed $e^*_{\text{g}}$ red-shift in our sample is mainly due to the presence of distorted ligand environment and that simulations with $Q_2 = 0.2$ eV and $0.5$ eV match better the experimental data.

However, the experimental spectrum shows features of distinct distorted states. Since the geometrical distortion of the oxygen octahedron matrix reflects atomic dislocations, i.e. displacements, from the ordered atomic arrangement, the results displayed in Fig. 5 point to the coexistence of a multitude of distorted ligand conformations.

The cause of the disordered nature of our samples can be identified in the small size of the nanoparticles. The finite dimensions of the NS films constituents and their irregular shape (see Fig. 2) provide a higher surface area with respect to bulk materials, naturally leading to symmetry breaking and to the creations of random atomic dislocations.

The disorder-induced L$_3$ red-shift with respect to the bulk samples reported for VO$_2$ can be observed for the whole set of VO$_x$ films.

The L$_3$ maximum position as a function of the oxidation state is depicted in Fig. 6. The comparison with crystalline bulk samples found in literature (Chen et al. 1994) evidences a stoichiometry-dependent L$_3$ red-shift for the whole oxygen concentration range.

This implies that the degree of disorder increases with stoichiometry which can be understood considering that to a high number of oxygen atoms corresponds a high number of achievable distorted atomic conformations. As a consequence, the highest oxidation states are most likely to assume a severely distorted atomic arrangement with respect to the samples with less oxygen.

The evidence of the disordered ligand environment for all the oxygen concentration is linked to the synthesis method and to the nucleation conditions of the nanoparticles. During the supersonic expansion of the seeded cluster beam, the thermal energy acquired during the nucleation process is converted in kinetic energy quenching the atoms in disordered conformations. This process is particularly efficient in small nanoparticles whose high surface area favours the symmetry breaking of ligand conformations.

![Fig. 5](image1.png) Left: simulated V L edges spectra of VO$_2$ for different $Q_2$ energy splitting values and NS VO$_2$ measured spectrum (black). Right: schematic representation of the energy levels splitting passing from $O_h$ to $D_{4h}$ symmetry.

![Fig. 6](image2.png) Comparison between the V L$_3$ energy position as a function of stoichiometry for the NS VO$_x$ film (black) and crystalline samples data (blue) found in literature (Chen et al. 1994).
This demonstrates that there is a complex influence of stoichiometry and disorder in VO$_x$ NS films with respect to the bulk analogue that suggest a way to tailor the electronic structure of these materials.

**Conclusion**

We investigated in situ NS VO$_x$ NS films using XANES spectroscopy demonstrating that nanoparticle size and stoichiometry contribute to determine the symmetry of ligands arrangement.

We demonstrated that broad $t^*$ and $e^*$ features and reduced $t^*$ -e splitting with respect to crystalline samples represent the fingerprints of disordered ligand matrix. This is confirmed by XANES simulations that show the coexistence of multiple distorted local environments. Our results point towards a stoichiometry-dependent red-shift of the V L$_3$ edge with respect to the bulk counterpart. This transfer of spectral weight towards lower photon energy is induced by the disordered structure of our samples which can be directly connected to the reduced dimension of the nanoparticles composing the films. Our study shows the complex interplay among stoichiometry, sample size and disorder in vanadium oxides and provides a tool to monitor and control the disorder in NS VO$_x$ films. Our results are of potential interest in different research area, such as Li-ion insertion devices, where the use of a disordered and NS material may be beneficial.

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**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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