ABSTRACT: We review oxygen K-edge X-ray absorption spectra of both molecules and solids. We start with an overview of the main experimental aspects of oxygen K-edge X-ray absorption measurements including X-ray sources, monochromators, and detection schemes. Many recent oxygen K-edge studies combine X-ray absorption with time and spatially resolved measurements and/or operando conditions. The main theoretical and conceptual approximations for the simulation of oxygen K-edges are discussed in the Theory section. We subsequently discuss oxygen atoms and ions, binary molecules, water, and larger molecules containing oxygen, including biomolecular systems. The largest part of the review deals with the experimental results for solid oxides, starting from s- and p-electron oxides. Examples of theoretical simulations for these oxides are introduced in order to show how accurate a DFT description can be in the case of s and p electron overlap. We discuss the general analysis of the 3d transition metal oxides including discussions of the crystal field effect and the effects and trends in oxidation state and covalency. In addition to the general concepts, we give a systematic overview of the oxygen K-edges element by element, for the s-, p-, d-, and f-electron systems.

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Oxygen is the third most important element in the universe\(^1\) and is the most important element from the earth crust and oceans. From biological processes to industry, the reactivity of oxygen is at the basis of many key chemical reactions. Before dioxygen accumulated in the atmosphere (referred to as the Great Oxygenation Event\(^1\)), it first reacted with the earth crust minerals to oxidize them and form oxides such as iron oxides.\(^3\) The reactivity of oxygen has been the underlying process driving the evolution of early organisms from anaerobic species to aerobic ones via the evolution and development of a complex redox biology.\(^4\) In more recent time scales, and from a technological point of view, mankind has progressively developed knowledge and control on the oxidation reactions: from combustion reaction producing fire, which was the first source of energy for humankind,\(^5\) to modern fuel engines and advanced catalysis,\(^7\) oxidation is a source of energy; from bloomery to blast furnace,\(^8\) redox processing of metals, which thermodynamics is described in the Ellingham diagrams, led to the development of metallurgy and all its technological improvement across ages along with the ensuing oxidation-aging issues. In short, redox reactions are everywhere in our oxidized world.

Investigating matter from the point of view of the oxygen is thus particularly relevant to determine its role in the atomic and electronic structures and the chemical bonds and resulting reactivity. This is possible using the oxygen K-edge X-ray absorption spectroscopy as an element specific analytical tool.

Already in 1920, Kossel gave an attractive explanation of the X-ray absorption phenomenon: \(^11,12\) in Röntgen spectra, we see that the electron cannot make small jumps outside the atom all the neighbor trajectories are already occupied by electrons so it must make a large jump to find a free trajectory, and the absorption occurs only when the frequency is high enough to make it reach the surface of the atom. Further developments of theory of X-ray absorption in molecules were made by Petersen\(^13\) followed by the works of Smoluchowski\(^14\) and Kurylenko\(^15\) on the understanding of the absorption of crystals. Starting from the 1960s, a cascade of studies have been published on theory, applications, and techniques of X-ray methodologies, as outlined in the review by Stumm von Bordwehr.\(^16\)

In this review, we discuss the oxygen K-edge X-ray absorption spectra where the oxygen 1s core electron is excited to the lowest empty states. The 1s core electron can be excited by the absorption of a photon in X-ray absorption (XAS), by the scattering of an electron in electron energy loss spectroscopy (EELS) or by the inelastic X-ray scattering (IXS) of a high energy photon. We will systematically describe the oxygen K-edges of atoms, molecules, ions, adsorbates, liquids, and solids and the main experimental and theoretical aspects related to the measurements and the calculation of the oxygen K-edge spectra.

However, an exhaustive review of all published oxygen K-edge spectra is impossible and even beyond necessity. Our choices were dictated by the aim of covering the main families of oxygen based compounds: atoms, molecules, and solids and the whole periodic table of oxides. The review reveals that the application of oxygen K-edge spectroscopy is very variable depending on the research context: from the fingerprint approach to in-depth investigation of the origin of the spectral features based on theoretical calculations, the degree of understanding of all published spectra is very variable. Specifically, some systems have particularly been studied in great detail. For example, we decided to only very briefly discuss the oxygen K-edges of water and of cuprates, where several dedicated reviews exist. The relative size of each sections of this review reflects this heterogeneity. Nonetheless, we have tried to fill gaps in the understanding of O K-edge spectra from the different research fields and provide a unified understanding of the information that can be obtained from this technique.

1.1. Nomenclature

The names of many spectroscopy techniques show some historical variation, and also, some research fields have different names for X-ray absorption spectroscopy. Throughout this review, we will use the term, “Oxygen 1s x-ray-absorption edges of transition-metal oxides”. Instead of “K-edge”, one can use the 1s orbital notation. Some research areas use the term “X-ray absorption near edge structure” (XANES) or “near edge X-ray absorption fine structure” (NEXAFS). We indicate the electron energy loss spectra with EELS and alternative notations include...
near edge energy-loss spectroscopy" (ELNES). Finally, inelastic X-ray scattering (IXS) is alternatively indicated with "X-ray Raman scattering" (XRS) or "non-resonant inelastic X-ray scattering" (NIXS).

1.2. Published Reviews and Books

Oxygen K-edge XAS spectra have been reviewed in a number of papers and reviews. The section on 3d transition metal oxides can be seen as an updated version of the 1989 paper "Oxygen 1s x-ray absorption of transition metal oxides".16 Chen wrote a detailed review of X-ray absorption spectra of transition metal compounds, including a section on the oxygen K-edge.17 There have been a number of reviews on the X-ray spectroscopy of water, in particular, the recent reviews by Smith and Saykally18 and by Nilsson and Petterson.19 Books on X-ray absorption spectroscopy include NEXAFS spectroscopy by Stohr20 and Core level spectroscopy of Solids.21

2. EXPERIMENTAL SECTION

The oxygen K-edge can be measured with X-ray absorption spectroscopy (XAS), using an X-ray energy larger than the $30 \text{ eV}$ excitation energy of an oxygen 1s core electron to an empty state. There are two alternative techniques based on the inelastic scattering of electrons and the inelastic scattering of photons. Electron energy loss spectroscopy (EELS) measures the energy loss of an electron and inelastic X-ray scattering (IXS) the energy loss of an X-ray beam. EELS and IXS are discussed in sections 2.3 and 2.4. An X-ray absorption experiment needs (1) an X-ray source, (2) a monochromator, and (3) a detector, where the sample condition is also an important aspect. In the next section, we briefly introduce some aspects of the experimental conditions.

2.1. Sources for X-ray Absorption Spectroscopy

Historically, XAS was measured with X-ray tubes.12 X-ray tubes give the largest part of their X-ray emission in the form of intense monochromatic X-ray energies related to the specific core—core X-ray emission channel, for example, the aluminum Kα source at 1486 eV. A second source of X-rays from the tube is the so-called bremsstrahlung. The bremsstrahlung energy loss is due to the slowdown of electrons from the metal in the anode and gives rise to a continuous spectrum of X-rays. Bremsstrahlung is much lower in intensity than the specific X-ray emission lines, but with a source optimized for bremsstrahlung emission, one has enough photons for good X-ray absorption measurements. The 1920 paper from Kossel describes the analysis of the XAS spectral shape.11 Kossel already realized that the X-ray absorption spectral shape was influenced by the valence electrons and the other "external conditions" of the absorbing atom in the surroundings where it was embedded. Using an X-ray tube, Fischer measured a series of oxygen K-edges in 1971 with 0.9 eV resolution.22 To compare X-ray sources, one usually compares the spectral brightness or brilliance of the X-ray beam, which is defined as the number of photons (in a certain bandwidth) per second and per mm² and mrad², where the brilliance of an X-ray tube is $10^8$. Depending on the details of the XAS experiment (for example, whether one measures a gas or a solid), the divergence and the spot size are often not crucial and the X-ray flux is a more important property. The flux is defined as the number of photons (in a certain bandwidth) per second. In this review, we will define the term "intensity" as equivalent to flux.

Synchrotron radiation emerged in the 1960s, but a limiting factor for oxygen K-edge XAS measurements was the absence of X-ray monochromators for the 500—600 eV range with good resolution. Synchrotron radiation covers all X-ray energies with a brilliance between $10^{13}$ and $10^{18}$ photons/s/mm²/mrad², in other words, $10^5$ to $10^{10}$ brighter than an X-ray tube. The X-ray monochromators were improved in the 1970s, and in 1980, Stohr et al. measured the oxygen K-edge spectrum with 2 eV resolution.23 A big step in experimental resolution was set in the 1980s with the development of the SX700 monochromator by Petersen24 and the DRAGON monochromator by Chen and Sette.25 Both beamlines reached a resolution that was better than the oxygen K-edge lifetime broadening of 0.3 eV, which implies that from this time the experimental resolution of oxygen K-edges was limited not by the experimental resolution, but by the intrinsic lifetime broadening.

More than 95% of all oxygen K-edges are measured with synchrotron radiation sources, but next to the synchrotron and X-ray tubes, a number of additional X-ray sources exist, respectively: (a) plasma sources, (b) high-harmonic generation (HHG) lasers, and (c) X-ray free electron lasers (XFEL). With a plasma source, one is also able to measure soft X-ray spectra.26 Kuhl et al.27 measured an oxygen K-edge spectrum with 2 eV resolution. Soft X-rays can also be generated with HHG laser systems. Most present HHG laser systems operate up to 100 eV, and they are especially used for femtosecond pump–probe experiments. Extensions to the oxygen K-edge at 530 eV are in principle possible on some newly developed sources and/or they are foreseen in the near future.28–32 The use of HHG and XFEL lasers for time-resolved experiments is discussed in section 2.6.

The oxygen K-edge of cosmic sources is measured with X-ray satellites XMM Newton and Chandra, with a resolution of approximately 2.0 eV.33

2.2. Detection Techniques

Transmission Detection. X-ray transmission is the most direct technique to measure an oxygen K-edge XAS spectrum and it allows the quantitative detection of the X-ray absorption cross section. The attenuation of 530 eV X-rays is only 100 nm at the oxygen K-edge of solid oxides, implying that one needs thin samples. A transmission measurement is only quantitatively correct if the measured sample is homogeneous in thickness. If thickness variations occur, the spectral shape will appear distorted with essentially the high-intensity peaks appearing flattened. This so-called pinhole effect also plays a role for samples that are inhomogeneous, for example, samples containing nanoparticles or otherwise nanostructured objects that are measured with an area-averaged X-ray probe. Alternative detection methods are based on decay channels of the core—hole. Fluorescence yield (FY) measures the amount of X-rays that are emitted. The energy that is gained by the electron that fills the core—hole can also be used to emit another electron in the Auger process. In the further relaxation of the core—hole, more electrons can be emitted and electron yield (EY) also measures the total amount of emitted electrons.

Fluorescence Yield Detection. The core—hole is filled with an electron from another shell and the energy can be emitted as an X-ray. Detecting the fluorescent X-rays after radiative core—hole decay can yield a method that is proportional to the XAS spectral shape. In total fluorescence yield (TFY) detection, one detects all emitted X-ray photons, from all elements present in the sample. Assuming that all nonresonant photoionization provides a constant background, the TFY measures the XAS spectral shape. However, spectral distortions can occur due to saturation and self-absorption effects, because usually the X-ray emission (being off-resonance)
has a larger probing depth than the penetration depth of the X-ray at the edge. 34−36 While scanning through an absorption edge, the X-ray penetration depth varies due to the XAS spectrum, and if there is no background absorption, the observed spectral shape would be highly distorted. If the background X-ray absorption dominates, the X-ray penetration depth is constant over the edge and no saturation occurs. In other words, TFY can only be used for materials where the observed element is dilute. We note that if the concentration of the absorbing element is very low, the FY from the other elements might dominate the signal, and the specific FY from the dilute element will have low signal-to-noise. Bulk oxides are too concentrated and their TFY spectra appear highly saturated. In addition to saturation effects, there can be effects due to self-absorption, in other words, the reabsorption of the emitted X-ray emission. Again, this effect is only significant for concentrated samples. 37 Saturation and self-absorption effects are angle-dependent, which also offers a method to quantify the saturation effects. In the case of the oxygen K-edge, there are no spectral modifications due to state-dependent decay. 38−40

Instead of TFY, only a limited energy range of the X-ray emission can be detected. Such a partial fluorescence yield (PFY) method can have an advantage if the X-ray emission signal is dominated by another element. As such, PFY measurements allow the detection of XAS spectra of very low concentrations, down to the 100 ppm range. 41 A special type of PFY is inverse PFY (IPFY). In this method, one detects the FY of a different element from the element from which the XAS spectrum is measured, where one can prove that in IPFY measurements no saturation effects can occur. 42

Electron Yield Detection. In total electron yield (TEY), electrons that escape from the sample are detected in electron analyzers. This can include the integrated signal from energy-dispersive detectors or, in the case of conductive samples, the current measurement to the sample. If the energy of the electrons is selected, one speaks of partial electron yield (PEY) or with the detection of a specific decay channel also of Auger electron yield. 43 A special property of electron yield is that the electrons have an escape depth that is on the order of a few nanometers, much shorter than the X-ray penetration depth, implying that in most cases the TEY signal is not saturated. 44,45 Thus, electron yield detection turns XAS into a surface-sensitive probe. This can be used to enhance the signal from the surface and near-surface part of the sample. The surface sensitivity also implies that the measured signal is not exactly equal to the bulk signal. In the case of surface oxidation or surface modification, the electron yield signal is different from the XAS spectrum of the bulk system. Even for a perfect surface in a single crystal, the TEY signal will be different from the bulk because the atoms in the top layer at the surface have different surroundings and corresponding electronic structure. Due to the probing depth of only 4 nm, the top layer contributes significantly (10%) to the total spectral shape. In addition, most crystals show surface reconstructions that extend a few layers into the crystal. This makes the TEY measurements usually affected by surface-induced effects to some extent.

If TEY is measured under a gas atmosphere, conversion electron yield (CEY) appears as an additional option. 46 In a gas atmosphere, the electrons emitted from the sample create ions that are counted, and as such, the CEY method effectively measures the XAS spectral shape. Often, two detectors are used that measure respectively the gas phase and the sample surface plus gas phase, allowing both signals to be differentiated. 47 The conversion of electrons to ions can be dependent on the electron kinetic energy and the molecules in the gas phase, which implies that CEY should be well calibrated during the experiments. Another type of CEY is applied in liquids. One can measure the XAS spectrum of a surface that is in contact with a liquid, by detecting the ions that are generated in the liquid by the escaping electrons, also known as the ion-current detection. 48 In general, these CEY methods should always be well calibrated and tested to check their linearity with the XAS cross section.

Ion Yield Detection. Ion yield detection (IY) is different from the ions measured in CEY methods described above. In ion...
yield, one measures a specific ion with a mass spectrometer while scanning through the XAS spectrum. Himpel et al. used IY to measure the surface of a CaF$_2$ crystal. IY is a common technique in gas phase experiments, where it is linked to photofragmentation detection. The oxygen K-edge can be measured by detecting various possible ionic fragments that are produced by the XAS process. IY has been applied to the oxygen K-edge by Hayakawa et al., who measured a series of cerium oxide clusters. Figure 1 gives an overview of the detection techniques of X-ray absorption spectroscopy, including transmission, electron yield, fluorescence yield, and ion yield methods.

2.3. Inelastic X-ray Scattering

Inelastic X-ray scattering (IXS), also known as X-ray Raman Scattering (XRS), measures the inelastic scattering of hard X-rays, not resonant with a core level. IXS is a low-intensity experiment, implying that an intense synchrotron or XFEL beamline is required. The oxygen K-edge can be measured with IXS by detecting the 530 eV energy loss. By changing the scattering angle of the IXS experiment, one can modify the momentum transfer. An important application of q-dependent measurements is the change of the ratio between dipole and quadrupole transitions. Pyknanen et al. applied q-dependent IXS to the oxygen K-edge of a series of alcohols. The advantage of the hard X-rays used in IXS is that one can measure the oxygen K-edge under extreme conditions, for example, high pressure. High pressure studies include the study of the phases of supercritical and solid phases of H$_2$O and studies regarding the nature of the high pressure phases of silicates. A review of high pressure publications until 2016 is given by Sternemann and Wilke.

2.4. Electron Energy Loss Spectroscopy

Electron energy loss spectroscopy (EELS) is usually measured with transmission electron microscope (TEM), but historically some dedicated EELS machines have also been used, for example, the EELS experiments performed by the group of Fink. The resolution of EELS in dedicated electron microscopes is 0.3 eV or better since the 1990s. As a rule, EELS is performed with an electron beam above 100 keV. The high energy and low momentum transfer effectively turns the 1/e Coulomb operator in inelastic electron scattering into an effective optical operator. Studies using low-energy (100 eV) electrons, for example, resonant EELS at electron energies equal to core level binding energies, are mainly limited to metal edges and to valence excitations. Using a 100 keV electron beam at low scattering angle, one can approximate the EELS spectrum with the dipole approximation. Under these approximations (high primary energy, low scattering angle), EELS becomes essentially equivalent to X-ray absorption. In EELS, one can tune the dipole/quadrupole ratio by changing the scattering angle. In the remainder of the manuscript, we will treat EELS spectra conceptually equally to XAS spectra. TEM-EELS has the advantage that one can reach atomic resolution, allowing the measurement of an oxygen K-edge of one atom (or better one atomic column) through a sample.

2.5. In Situ Experiments

Oxygen K-edge XAS is usually performed in vacuum. Soft X-rays have a penetration depth through air on the order of 1 cm. In principle, this is long enough to perform transmission or X-ray fluorescence experiments. From 1988, Fischer et al. performed a series of in situ TFY experiments at pressures up to 100 mbar. Using nanoreactors in a scanning transmission X-ray microscopy (STXM), the oxygen K-edge of Fe$_3$O$_4$ supported on SiO$_2$ has been studied at 1 bar under Fischer–Tropsch conditions. More examples of STXM experiments are given in the section on X-ray microscopy.

A more popular approach is to use electron yield detection. Because TEY is surface sensitive and TFY is bulk sensitive, their comparison can be very useful. TFY and TEY were compared to study the changes of propylene on the surface of Ag/Y zeolites. The use of both TFY and TEY nicely separates the behavior of the surface and the bulk. Traditionally, in situ TEY measurements are performed in combination with X-ray photoemission and the XPS detector makes it difficult to go above 10 mbar pressure. A series of 1 to 10 mbar in situ oxygen K-edge studies have been performed on so-called near-ambient pressure photoemission beamlines. This allows the study of the surface of heterogeneous catalysts under working conditions in a flowthrough reactor. An example of an oxygen K-edge study that has been performed is the methanol oxidation to formaldehyde over copper metal. Figure 2 shows the oxygen K-edge of a mixture of methanol and oxygen at 0.52 mbar that reacts on a Cu foil at 520 K. The spectral shapes are a combination of O$_2$, CH$_3$OH, and the Cu$_2$O surface species. The system uses one detector that measures the gas phase and a second detector that measures the combination of the surface and the gas phase. Using the difference between the two detectors, the surface signal can be revealed. Experiments can be performed at different gas mixtures, pressures, and temperatures, as has been described in a number of applications. A series of detailed in situ oxygen K-edge studies has been applied to vanadium oxides, under different gas atmospheres and with different loadings of the vanadium. The analysis of the oxygen K-edge, in combination with DFT studies allows the distinction between separate vanadia, silica, and interface contributions. A more recent development is to use CEY detection up to pressures of 3 bar. One can combine oxygen K-edge XAS with time-resolved measurements. An overview of the recent developments for
time-resolved X-ray experiments with synchrotron sources, X-ray FELs, and high harmonic sources is given in the review by Kraus et al.\textsuperscript{71} Cavalleri et al. measured the time evolution of the oxygen K-edge of VO\textsubscript{2} using a laser-sliced synchrotron beam with a time resolution of 500 fs, from which new information on the photoinduced metal–insulator transition in VO\textsubscript{2} was found.\textsuperscript{72} The LCSL X-ray FEL was used to study the changes in the excited-state electronic structure of the molecule by using the changes in the oxygen K-edge.\textsuperscript{73} A number of fs oxygen K-edge XFEL studies have been performed on adsorbates, for example, CO on Ru(0001).\textsuperscript{74,75} Femtosecond XFEL studies focusing on the oxygen K-edge of small oxide material, and for example, Ward et al. studied the oxygen K-edge of uranium oxide with a STXM microscope.\textsuperscript{85} The STXM microscopes can measure the oxygen K-edge with spatial resolution. Such measurements can reach 0.1 nm, while STXM and TXM 10 nm. On the other hand, STXM allows for thicker samples and operando conditions.\textsuperscript{84} De Smit et al. used the oxygen K-edge STXM data to provide a thickness profile of their Fe\textsubscript{3}O\textsubscript{4}/SiO\textsubscript{2} catalyst sample, while the metal edges were used to probe their spatial variation and valence.\textsuperscript{82} The methods used to derive the atomic resolution images are explained by Wang et al.,\textsuperscript{83} who used the oxygen K-edge of SrTiO\textsubscript{3} as an example.

With scanning transmission X-ray microscopy (STXM) one can measure the oxygen K-edge with spatial resolution. Such STXM-XAS experiments are analogous to STEM-EELS, with the difference being that with TEM one can reach 0.1 nm resolution and with TXM 10 nm. On the other hand, STXM allows for thicker samples and operando conditions.\textsuperscript{84} De Smit et al. used the oxygen K-edge STXM data to provide a thickness profile of their Fe\textsubscript{3}O\textsubscript{4}/SiO\textsubscript{2} catalyst sample, while the metal edges were used to probe their spatial variation and valence.\textsuperscript{84} The literature is rich with oxygen K-edge studies performed with STXM microscopes, but a full overview of the performance of this technique goes beyond the scope of this review. Sharma et al. studied the oxygen K-edge spectra in graphene and graphene oxide with a STXM microscope.\textsuperscript{85} The STXM microscopes can also be used to select a specific small oxide material, and for example, Ward et al. studied the oxygen K-edge of uranium minerals,\textsuperscript{86} while a series of rare earth oxides have also been studied with a STXM.\textsuperscript{87} An alternative X-ray microscopic technique is X-ray photoelectron emission microscopy (PEEM). In X-ray PEEM, one measures the emitted electron with an electron microscopic lens system. X-ray PEEM is mainly used to study magnetic structure, using either magnetic linear dichroism (MLD) or magnetic circular dichroism (MCD). Kinoshita et al. studied the magnetic domain structure of a NiO(100), comparing the X-ray PEEM images measured at the oxygen K-edge with the nickel L\textsubscript{3} edge.\textsuperscript{88} Oxygen K-edge PEEM was also used to study polymer mixtures.\textsuperscript{89}

### 2.8. Using the Oxygen K-edge as Resonance

Resonance studies include the use of the oxygen K-edge in resonant photoemission spectroscopy (RPES) and resonant inelastic X-ray scattering (RIXS). In addition, one could perform X-ray scattering at the oxygen K-edge and resonant EELS with electron sources. The oxygen K-edge of solid samples does not contain excitonic states, which turns the two-dimensional RIXS plane essentially in a convolution of the oxygen K-edge and the nonresonant 1s2p X-ray emission spectrum.\textsuperscript{50} The oxygen K-edge is particularly used as a resonator for the iridates. Because the Ir 2p edge lies in the hard X-ray range, it has low resolution. The soft X-ray oxygen K-edge then adds improved resolution to study the electronic structure. This has been used for Ba\textsubscript{2}IrO\textsubscript{4}\textsuperscript{\textsuperscript{21} and SrIrO\textsubscript{3}\textsuperscript{92} where the magnon dispersion is measured. The oxygen K-edge RIXS spectra are also used to study the core–hole clock mechanism, in which it is reasoned that by changing the excitation energy, one change the ratio between resonant and normal decay channels. An example is the study on the oxygen K-edge RIXS of alcohols.\textsuperscript{93}

### 3. THEORY

#### 3.1. Density Functional Theory (DFT)

**Independent Particle Approximation.** The main point of the independent particle approximation is the decoupling of the dynamics of the many-body system in a single-particle description. The Hamiltonian can be rewritten with a term that mimics the interaction of the electrons with the average field that they feel and contains the kinetic energy of the electron and the potential energy of the electron in the field of the nuclei. With each independent electron is associated a spatial function and a corresponding energy eigenvalue, obtained by solving the Schrödinger equation with this Hamiltonian. The independent particle approximation is also called the single-particle approximation and the corresponding Hamiltonian the one-electron Hamiltonian.

Within the independent particle approximation, it is possible to replace the many-body quantities in the Fermi golden rule using the many-body initial and final state wave functions $\Psi$ and $\Psi_f:

$$\sigma(\omega) \approx \sum_f \langle \Psi | \hat{\epsilon} \cdot \hat{R} | \Psi_f \rangle^2 \delta(\omega - E_f)$$

$\hat{R}$ is the transition operator that we have approximated as the dipole operator. $\delta$ indicates a $\delta$-function including $E_f$ the energy difference between the states $\Psi$ and $\Psi_f$. The goal is the computation of the dipole matrix elements and the transition energies where different approximations can be adopted.

**3.2. Resonant Photoemission Spectroscopy (RPES) **

The many-body eigenstates and upper case operators on the left side of the equation are substituted by the lower case effective single-particle states.\textsuperscript{54} In the remainder of this manuscript, we will use the term “single-particle state” as equivalent to the term “orbital”. We note that this change from the many-body description to a single-particle description involves two coupled approximations: (1) the description of the ground state in the independent particle approximation and (2) the description of the final state in the independent particle
approximation. We note that in the case of 2p L_{2,3} X-ray absorption of transition metal systems, the 2p core−hole in the final state has large electron−electron interactions with the 3d valence electrons that create the so-called multiplet effects. In the case of oxygen K-edges, we are dealing with a 1s core−hole and (exchange) interaction with the valence electrons smaller than 1 meV which can be ignored.

Transition energies can be computed with the Delta Kohn−Sham (ΔKS) method. The idea behind the ΔKS method is that the transition energies can be calculated as the difference between a ground-state energy, computed with a self-consistent field calculation, and an ionized and excited-state energy, obtained through a restricted open shell calculation. The procedure introduces an occupation number constrained in the 1s oxygen KS orbital, justified by its localized character. The nonorthogonal matrix elements between the Kohn−Sham ground-state determinant and each excited-state determinant are used in order to obtain the oscillator strength. With the ΔKS scheme, two calculations are needed, one for the ground state and another one for the excited state. Good agreement with experimental results have been obtained in the calculation of small molecules and metal adsorbate cluster oxygen K-edges in terms of excitation energies and trends in oscillator strengths, where the energy underestimation from the TD-DFT scheme is improved.

Only one calculation is needed in an approximate procedure that is called the Slater transition state, even if different calculations are needed for each excited state. This procedure consists of the estimation of the excitation energy as orbital energy differences between two levels of variationally determined states, which are defined as the transition state, with only one-half electron excited. The Slater transition state provides an estimation of the excitation energy with a correction to second order in the occupation numbers. It is important to notice that the number of electrons is conserved.

Once we have the two energies, the transition energy can be computed as a difference between them. Viewed as a balanced compromise between initial and final states, the transition-state orbitals can be used to represent both states, which reduce the evaluation of transition moments to single-electron transitions without affecting the other occupied orbitals. The evaluation of the oscillator strengths becomes simplified since the same orbitals are used to describe the final and initial states. The weakness of the method is that one has to compute an X-ray absorption excitation for each state. An approximation to the Slater transition-state scheme that circumvents the state-by-state calculations required is the transition potential approach formulated by Triguer et al. The approximation consists of removing half an electron from the oxygen 1s core level and to obtain all the possible excited states with only one global diagonalization from the potential generated by the core with the half-occupied core-hole. The complete spectrum evaluation in transition potential density functional theory (TP-DFT) consists of the calculation of the matrix elements between the orbitals corresponding to the 1s and the unoccupied orbitals since the initial and final states have the same orthogonal basis set. It has been demonstrated that excitation energies computed in the TP-DFT framework require relaxation effects to be taken into account up to the second order. Since the TP-DFT calculations can be considered as approximations to the Slater transition-state scheme, the use of the half core−hole is theoretically justified. Extensions of the TP-DFT method, particularly suited when excitonic effects are strong, are the full core−hole (FCH) and the excited core−hole (XCH). A correct description of the core−hole is one of the most challenging tasks of XAS simulations.

In Figure 3, some of the approximation schemes for the core−hole are indicated. The Z+1 approximation, also called the empty core−hole (ECH) approximation, introduces the core−hole as an additional nuclear charge. The equivalent core−hole approximation is justified, since for electrons in the external shells, a deep core−hole is felt as a positive charge. On an oxygen 1s core excited spectrum, oxygen is replaced by fluorine, but because of the high electronegativity of the latter, a charge transfer could be induced in the neighboring atoms with a subsequent increase in the p-population of the excited core causing a distortion in the spectral shape. Its applicability can be questioned also if the system has shallow core−holes. In the exchange core−hole (XCH) approach, the potential of the first fully core excited state determines all the excited-state orbitals. A strong approximation is made in the XCH procedure since the orbitals computed for one specific final state are
supposed to describe the initial and final states, and no information about the initial ground states is present in the orbitals used. The FCH method is similar to the XCH: the core electron is ionized, which creates a hole in the core level, but in this case, the impact of the excited electron on the core—hole is neglected. The DFT-based methods cited above can, in principle, be applied to atoms, molecules, and condensed phases. All these one-electron calculations approximate the particle hole and the correlation effects.

3.2. DFT in Solids

If one deals with solids, the Fermi Golden Rule is usually rewritten as

\[ \sigma(\omega) \approx M^2 \rho \]

where \( M \) is the one-electron transition matrix element, and \( \rho \) is the density of states. All electronic structure schemes can in principle be used in order to generate the density of states (DOS) and within a conceptual DFT framework; many routes can be used to evaluate the empty DOS and the XAS spectral shape. The specific methods can be divided into the following:

- real space multiple scattering methods (FEFF)
- real space wave function method (ADF, ORCA)
- reciprocal space band structure methods (Wien2K, Quantumespresso)

We refer to the websites and original references of the methods indicated for details. The majority of solid-state X-ray absorption calculations have been performed with local density approximation of the density functional theory. A large number of specific procedures have been developed to solve the one-electron Schrödinger equation.

Using a linear method, different routines can be used to solve the band structure problem. The two main schemes both make use of the variational principle for the one-electron Hamiltonian, and their difference lies in the formulation of the trial functions: linear combinations of energy independent augmented plane waves (LAPW) or muffin-tin orbitals (LMTO). The strength of these procedures is the use of the variational principle to solve the Schrödinger equation with energy independent basis functions, so that the secular equations become linear in energy.

A plane wave basis set with a pseudopotential approximation has been introduced to model the interaction between ions and valence electrons. The pseudopotential neglects the inner core electrons and their strong potential due to the attraction to the nuclei. All electron descriptions of the system involve a strong orthogonal constraint between all the core and valence orbitals. The orthogonality relation presumes strong oscillations of the valence orbitals in the vicinity of the nuclei, as indicated in Figure 4. These oscillations, which require a large number of plane waves to be described, can be neglected in the pseudopotential framework, where the orthogonality relation is no longer valid, and only a reduced number of plane waves is needed to describe the correct behavior of the valence wave functions. The pseudopotential is then constructed for an initial DFT calculation for an isolated atom: a core radius \( r_c \) should then be chosen and the valence wave functions are adapted to it in order to remove the nodal structure; these are the pseudo-wavefunctions. The pseudopotential approximation has been widely employed to model the oxygen K-edge XAS in many different systems, and satisfactory agreement with experiment has been reached.

3.3. Core—Hole Effects in Solids

It has been demonstrated that the core—hole effects are stronger for spectra arising from elements in an ionic or covalent compound with lower electronegativity. In Figure 5, the theoretical results for the ionic compound BeO are compared for both the metal and the oxygen edge with and without the core—hole. The presence of the core—hole alters both spectra, but in the oxygen case, the modifications to the spectral shape are less significant. Although core—hole effects have to be taken into account in order to have a good agreement with experiment, omitting any treatment of the core—hole usually results in a shift of the threshold to higher energies and modified peak intensities in the low energy part of the spectra. In Figure 6, the oxygen K-edge spectra of GeO\(_2\) with and without the core—hole is compared to the experiment. In this case, the relative intensities of some peaks are better reproduced without the inclusion of the core—hole. This is evidence that the core—hole potential, in the adopted supercell approach, is too attractive. If one compares the spectra without the inclusion of the core—hole with the ground-state DOS calculations, one can show that in some systems a ground-state calculation can give features consistent with experiment and provide information to explain the features in terms of orbital hybridization.

3.4. Beyond the One Particle Approximation

We have seen that the one-body DFT calculations successfully describe the oxygen K-edges of delocalized systems. However, they lack accuracy in the description of partly localized and atomic edges, where many-body and multiplet effects are important. This encouraged the push to go beyond the single-particle picture with the inclusion of many-body effects. After the first attempts made by Zangwill and Soven and Zaanen et al. to overcome the single-particle description, nowadays the most widely used schemes are the Bethe Salpeter equations (BSE) and the time dependent density functional theory (TD-DFT) based methods. When Runge and Gross proposed the TD-DFT scheme to compute spectra, it became very popular for molecules, but its implementation for solid systems is more recent. The TD-DFT description of the X-ray absorption cross section enriched with the core—hole contribution is also called the adiabatic local-density approximation. In this framework, the exchange correlation kernel depends only on the density in the ground state (\( t = 0 \)). The
potential generated by the kernel is local in space and time, and it is exactly this feature that produces an unsatisfactory long-range behavior in solids.\textsuperscript{126,129} Another obstacle to the correct description of solids with TD-DFT is linked to the inability of the latter to reproduce bound excitations\textsuperscript{128} that leads to failure of the core–hole effect description. Many attempts\textsuperscript{130–132} have been made in order to make TD-DFT suitable for the description of the photoabsorption process in extended system, since it would be desirable in terms of simplicity,\textsuperscript{126} that is, two-point equations are required in TD-DFT instead of the four-point equations in BSE.

Figure 5. (Left) Beryllium K-edge of BeO calculated with and without core–hole. (Left) Oxygen K-edge of BeO calculated with and without core–hole.\textsuperscript{119}

Figure 6. (a) Theoretical GeO\textsubscript{2} oxygen K-edge with and without core–hole compared with experiment. (b) Theoretical oxygen K-edge XAS without core–hole compared with a ground-state oxygen DOS calculation.\textsuperscript{120}
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The Bethe-Salpeter equation (BSE)\(^\text{133}\) represents the equation of motion of a particle—hole state, as for example a photoexcited electron to the conduction band from the 1s oxygen orbital. The BSE description of X-ray absorption includes single-particle terms that describe the quasi-particle energies of the core–hole and the excited photoelectron, together with the interaction between them. The interaction between the electron–hole pair has two terms: the Coulomb interaction, which includes adiabatic screening of the core–hole, and an unscreened exchange term.\(^\text{134}\) BSE calculations are considerably heavy in terms of computational cost and usually they have been limited to systems of restricted dimension. A BSE calculation consists of the following steps: (i) determine the ground-state electronic structure, (ii) correct the quasiparticle energies by adding a GW self-energy, (iii) evaluate the screening response to the core–hole, and (iv) determine the excitation spectrum of the BSE Hamiltonian. Liang et al. reported the oxygen K-edge spectra of several transition metal oxides obtained through the BSE procedure and they showed the accuracy of the method in predicting the excitonic character of some peaks. The reproduction of the correct excitonic features of the spectra is ensured in BSE by the screened Coulomb interaction \(W\), present in the interaction kernel, that acts on the electron–hole matrices and introduces a long-range coupling between the electron and the hole wave functions.\(^\text{35}\)

An overview of the calculation procedures is given in Figure 7.

3.5. Charge Transfer Multiplet Calculations

Van Elp and Tanaka calculate the oxygen K-edge spectra of transition metal ions from a charge transfer multiplet model.\(^\text{136}\) The charge transfer multiplet model has been successfully applied to the metal L\(_{2,3}\)-edges \(\text{(2p XAS)}\) of transition metal ions,\(^\text{21}\) which makes it interesting to check its applicability to the oxygen K-edges of the same systems. There are however a number of differences between the metal L\(_{2,3}\)-edge and the oxygen K-edge; in particular, the metal L\(_{2,3}\)-edge creates excitonic states that are dominated by large \(2p\text{3d}\) interactions (multiplet effects) and a large \(2p\text{core–hole}\) spin orbit coupling. These interactions are absent for the oxygen K-edge, and only the dd-electron correlations remain. The exchange interaction of the dd-electrons is treated in spin-polarized DFT calculations, leaving out only the orbital polarization. In other words, the charge transfer multiplet calculations include the orbital polarization, but they do not include the band structure effects due to the crystal structure. Because the oxygen K-edge is broadened by the 300 meV lifetime broadening, we do not expect that the orbital polarization effects will be visible.

3.6. Post Hartree–Fock Methods

Excitations from core orbitals to empty molecular orbitals cannot be computed with Hartree–Fock because of some serious difficulties:\(^\text{137}\)

- convergence achievement
- variational collapse\(^\text{138}\)
- correlation effects

Post Hartree–Fock schemes have been investigated in order to overcome these difficulties. Approaches based on the coupled cluster hierarchy have been developed by Coriani\(^\text{139}\) and Nooijen\(^\text{140}\) using a linear response approach and an equation of motion, respectively. These methods yield a very good description of the initial and final state wave functions, but because of the high computational cost, their applicability is limited to small molecular systems.

Asmuruf and Besley proposed a second-order perturbative approximation to the coupled cluster with single and double excitation (CCSD) based on a single configuration interaction (CI). The methodology provides results equivalent to TD-DFT for the valence excitations, but because of the inclusion of the exact Hartree–Fock exchange, better performance is obtained in the core excitation treatment.\(^\text{141}\) Additional methods include the symmetry-adapter cluster configuration interaction (SAC–CI)\(^\text{142}\) and multiconfigurational self-consistent field schemes.\(^\text{143}\) The applicability of these methods is limited to small systems since the choice of a relevant active space including local and nonlocal orbitals yields large calculations.\(^\text{144}\)

4. ATOMS AND MOLECULES

Energy States of Open-Shell Systems

The description of the energy states of open-shell systems needs a description of the coupling of their angular momenta to describe the many electron states. The \(L\), \(S\), and \(J\) quantum numbers of the multielectron state are indicated with a so-called term symbol that is written as \(\text{2S+1}LJ\). The \(2p\text{1}\) configuration of an oxygen atom has the term symbols \(3P\), \(1D\), and \(3S\), where the orbital quantum is indicated with letters: \(L = 0\) is written as

![Figure 7. Oxygen K-edge spectra can in the first approximation be calculated from DFT codes, where molecules are usually calculated from molecular DFT codes and solids with band structure codes or multiple scattering. The oxygen p-contribution to the MOs or the oxygen p-projected DOS can be compared with the oxygen K-edge spectral shape, where matrix elements are also included. As a next step, the core–hole effect can be included, where many different procedures have been used both for molecules and for solids. In the case of solids, one has to perform a supercell calculation to prevent the core–hole from interacting with each other. Formally, the correct way to calculate the oxygen K-edges is to apply electron–hole excitation schemes such as TD-DFT or BSE, where the complexity of the calculations necessitates approximations. In systems where the dynamics of the system is important (including liquids), the calculations must be combined with molecular dynamics calculations. For example, a series of atomic positions can be determined and their X-ray absorption spectra added. Because of the 300 meV lifetime broadening, (in most cases) the effects of multiplets, orbital polarization, and magnetic exchange are not visible. Vibrations are visible in the case of (small) molecules.](https://dx.doi.org/10.1021/acs.chemrev.9b00439)
S(harp), L = 1 as P(incipal), L = 2 as D(iffuse), and L = 3 as F(undamental). The 1P state is split by the 2p spin–orbit coupling into, respectively, 3P, 3P, and 3P states, using the rule that J runs from L+1 to L−1 in steps of 1. The Hund’s rules determine the state with the lowest energy, respectively, the state with (1) the largest S, (2) the largest L, and (3) the largest J, where in case a shell is less than half-filled, the lowest J is the ground state due to the inverted effect of the spin–orbit coupling. This makes the 3P state the ground state. The 3P has an energy of 2.0 eV above the ground state and the excitation energy of the 1S state is 5.5 eV. The 2p spin–orbit splitting is small and the energy difference between the 3P ground state and the 3P state is only 25 meV.

4.1. Oxygen Atom

Atomic oxygen has a total of eight electrons with the electronic configuration 1s22s22p6. The 2p orbitals are partially filled and there are different ways to arrange the electrons in the 2p orbitals. Figure 7 shows the oxygen K-edge of atomic oxygen on a logarithmic scale. In the first peak at 528 eV, a 1s electron fills one of the two 2p holes and the final state configuration is 1s22s22p5. This configuration has two open shells, i.e., 1s22p5. The term symbols of such a configuration can be found from the multiplication of the term symbols from the individual shells, in this case, S times P, yielding respectively, P and P. In oxygen, these states are split by 2.5 eV. Including the 2p spin–orbit coupling yields the 3P, 3P, and 3P states. This implies that from the 3P Hund’s rule ground state, one can reach three final states, respectively, the 3P, 3P, and 3P states. The 3P and 3P states are separated by only 40 meV and appear as one peak in the experiment. The 3P state, which is 2.5 eV higher in energy, is not visible due to its low relative intensity of 10−5. The result is that the excitation from the 1s to the 2p state has effectively one visible peak as shown in Figure 7 at 528 eV.

The ionization energy of the oxygen atom relates to the excitation of the 1s electron into a free electron. In fact, there are two ionization energies depending on the relative orientations of the spins of the excited core electron and the two holes in the 2p states. This yields, respectively, states with S = 2 or S = 4, as indicated in Figure 8 at 545 and 548 eV.146 The peaks in between the edge and the ionization energy above 540 eV are assigned to transitions from the 2p ground state to the 1s2p4 configuration states, where n can take the values 3, 4, 5, etc., the so-called Rydberg states. The 1s2p4 configuration is split into multiplets by the 1s2p exchange, the 2p2p electron–electron interactions, and the small 2p spin–orbit coupling. The term symbols are, respectively, S, P, P, and P, from which only the and states are split by 3.5 eV, and this yields two series of Rydberg states as given numerically in Table 1.147

| configuration | energy (eV) | | | relative intensity |
|---------------|------------|---|---|------------------|
| 1s2p4         | 527.8      | 528.1 | 100 | 100              |
| 1s2p4         | 541.3      | 542.2 | 6.2 | 5.1              |
| 1s2p4         | 542.6      | 543.8 | 1.6 | 1.9              |
| 1s2p4         | 543.2      | 544.3 | 0.36 | 0.36            |
| 1s2p4         | 545.6      | 546.8 | 5.4 | 2.0              |
| 1s2p4         | 547.2      | 547.2 | 1.9 | 0.5              |
| 1s2p4         | 547.7      | 547.8 | 0.36 | 0.24            |

The 1s2p peak and the Rydberg peaks in the oxygen K-edge of the oxygen atom have been calculated by Petrini and Araujo.148

4.2. Oxygen Ions

When oxygen is ionized, the ground state changes from 2p4 to 2p3, yielding a new ground-state symmetry as indicated in Table 2. Because the positive charge of the O1+ ion, it costs more energy to excite its 1s core electron. The energy shift from a neutral oxygen atom to a O1+ ion is 4.5 eV. Table 2 makes use of data on oxygen ions from X-ray satellites such as the Chandra X-ray observatory and XMM Newton. These X-ray satellites are able to detect the oxygen K-edge XAS spectrum with around 200 meV resolution. Oxygen is often studied, as it plays a key role in the understanding of the chemical evolution of the Universe. If the oxygen atom is ionized to a O1+, the oxygen 1s binding energy increases due to the decrease of the valence electrons. The O2+ and O3+ ions have an XAS spectrum containing three peaks. Due to their less-than-half-filled configuration and related Hund’s ground state with minimum J, three final states have similar intensity.

| ion | configuration | symmetry | 1s2p (eV) |
|-----|---------------|---------|-----------|
| O   | 2s2p4         | p       | 528.8     |
| O+  | 2s2p3         | S       | 533.3     |
| O++ | 2s2p2         | p       | 537.2     |
| O+++| 2s2p1         | p       | 545.8     |
| O++++| 2s       | S       | 554.6     |
| O++++| 2s       | S       | 562.3     |

The Hunds rule ground state is given along with the symmetries of the main peaks in the final state.
Photoionization and photoabsorption cross sections used to model important astrophysical processes have been provided first by theory, as at that time, limited experimental reliable data were available. Early theoretical photoionization cross sections are computed by means of Hartree–Slater wave functions and Dirac–Slater wave functions. Pradhan et al. calculated the resonant transition energies and the photoionization cross sections are computed by means of Hartree–Slater wave functions and Dirac–Slater wave functions. McLaughlin et al. reported photoabsorption cross sections obtained with RMPS and high-resolution experimental measurements.

4.3. Binary Molecules CO, NO, and O₂

In CO, NO, and O₂ molecules, there are, respectively, 14, 15, and 16 electrons that can be placed into the molecular orbitals. The orbitals in molecules can be constructed from the combination of the atomic orbitals. Figure 9 shows the molecular orbitals of O₂, CO, and NO. The 1s orbitals of O₂ are not overlapping with the neighboring orbitals and retain their atomic character. The 2s and 2p orbitals of O₂ overlap with the neighboring 2s and 2p orbitals and have σ bonding and antibonding orbitals. The 2p orbitals of O₂ also have π overlap with the neighboring 2p orbitals. The antibonding π₋₂ orbitals of molecular orbitals contain two paired electrons, and the antibonding π₊₂ are empty. At higher energies, one can find the molecular orbitals due to 3s, 3p, and 3d atomic orbitals. The molecular orbitals of NO are equivalent with one electron in the antibonding orbital π₋₂, and CO has zero electrons in the antibonding orbital π₋₂. Because for CO, NO, and O₂ molecules there are two partly empty 2p states, one expects two peaks in the XAS spectrum corresponding, respectively, to transitions to the π⁺ and σ⁺ states.

O₂. The π₋₂ orbital in O₂ is doubly degenerate and half filled, implying that different electronic states can be obtained depending on the electronic configuration. There are three different states, respectively, ¹Σ₋, ¹Π, and ¹Σ⁺, corresponding to different arrangements of these electrons in the C₂ point group symmetry. The ground state of molecular oxygen is the triplet state ¹Σ₋. The oxygen K-edge spectrum of O₂ has been extensively studied. The strong first resonance in the oxygen K-edge absorption spectrum (E₂₃₈₅) at 531 eV (O 1s → π⁺) has been analyzed, and the vibrational structure has been resolved. The ionization energy E₁₈₅ of O₂ is found at approximately 547 eV, and like for the oxygen atom, it is split into a quartet and a doublet state depending on the coupling of the core electron spin to the valence spins, indicated in Figure 10.

The challenging part of the assignment of the peaks in the O₂ absorption spectrum is the region between 535 and 545 eV. In this region, we expect the 2pσ⁺ states but also the Rydberg states. The 1s → σ⁺ resonance is split into two features converging to the ⁴Σ⁻ and ⁴Σ⁺ ionization thresholds at 543.39 and 544.43 eV. Kosugi et al. reported symmetry-resolved oxygen K-edge spectra where two σ features with different intensities are assigned to the bound-state transitions 2pσ⁺. The solid-state oxygen K-edge spectra of O₂ show no strong quenching or broadening of these peaks. This suggests that the peaks cannot be assigned to Rydberg states, because the Rydberg states will behave differently in the solid state, due to strong intermolecular overlap of the extended 3p and 4p states. Wurth et al. reported an exchange splitting of 4 eV and assigned the σ⁺ resonance to the spin up and spin down transitions 1σ₋ → 3σ₂⁺. Ruckman et al. studied the O₂ gas-phase spectrum in comparison with the oxygen K-edge of the alkali metal superoxides, and they reported an experimental exchange splitting value of 0.4 eV with both components located in peak B. The exchange splitting value that can be found in the literature spans a broad range, and this can be ascribed to the theoretical difficulties in treating the Coulomb and exchange interaction in the core–hole excited states. These two different interpretations of the absorption interaction lead to different conclusions. By comparison of the adsorbed O₂ and gas phase, the changes in the peak positions and intensities are interpreted either as quenching of the exchange splitting or as the consequence of the Rydberg nature of peak C in free O₂. ⁴Σ⁻.
The analysis of the de-excitation spectra helps the separation between $\sigma^*$ and Rydberg core excited states. The de-excitation spectra excited at feature C can be understood from the decay of one or more Rydberg states, whereas the de-excitation spectra for the chosen excitation energies within feature B are associated with the decay of the 2p$^2\sigma^*$ state. Under pressure, O$_2$ condenses to a liquid, and at 5.5 GPa, it solidifies to a solid. As a function of pressure, several solid O$_2$ phases exist. The oxygen K-edges of the liquid and solid O$_2$ phases show that as a function of pressure the $\pi^*$ and the $\sigma^*$ peaks move to higher energy, consistent with a decreasing O–O interatomic distance. In addition, the $\pi^*$ loses intensity due to the increased interaction between the O$_2$ molecules leading to a singlet ground state for the $\epsilon$(O$_2$)$_4$ phase.

**NO.** The electronic configuration of NO is equivalent to that of O$_2$ with only one electron in the 2p$_\pi$ orbital, resulting in a ground-state symmetry with term symbol $^1\Sigma^+$. Figure 11 shows the oxygen K-edge spectrum of NO, where the ionization limits are indicated as $^3\Pi$ and $^1\Pi$. The three theoretical components ($^2\Sigma^-$, $^2\Delta$, and $^2\Sigma^+$) in the 1s2p peak at 532.7 eV are indicated.

The feature with a maximum around 540 eV has been assigned to Rydberg transitions with quantum number $n \approx 3$, while the $\sigma$ resonances are observed above the ionization threshold. Three series (ns, np, and nd) of Rydberg states are found.

**CO.** Carbon monoxide (CO) has a singlet ground state, where there are no electrons in antibonding orbitals. The molecule has a closed-shell configuration, and its ground state is a totally symmetric singlet $^1\Sigma^+$. In Figure 12, the CO oxygen K-edge spectra is reported. The first feature in the oxygen K-edge is the bound excitation into an antibonding $\pi^*$ orbital that can be vibrationally resolved. In addition, for CO the $\sigma$ shape resonance appears above the ionization threshold. The two higher energy peaks are assigned to 3s$^1\sigma$ and 3p$\pi$ Rydberg excitations. The features in the continuum have been interpreted as the $l = 3$ shape resonance enhancement in the $\sigma$ channel or as a transition to the quasi-bound $\sigma^*$. In Table 3, we compare the excitation energies to the 2p$\pi^*$ peak in CO, NO, and O$_2$. The excitation energy increases from O$_2$ to CO, indicating that it becomes more difficult to excite the 1s core electron to the 2p$\pi^*$ state. The nitrogen K-edge shifts down in going from N$_2$ to NO. An increase in the excitation energy was also seen for the change from oxygen atoms to ions.
Extrapolating this to NO indicates that the oxygen atom in NO is more positive than in O₂, with the N atom being more negative. In CO, the oxygen atom is even more positive.

**Ions of Two Atom Oxides.** Lindblad measured the oxygen K-edge of a O₂⁺ and CO⁺ molecules. The 2pπ* peak shifts to higher energy in O₂⁺, in agreement with its positive charge implying a higher excitation energy. Ruckman et al. measured the oxygen K-edge spectra of the alkali superoxide KO₂. These systems can be considered to contain K⁺ and O₂⁻ ions and the oxygen K-edge of KO₂ can be interpreted as that of O₂⁻. The 2pπ* peak shifts to lower energy in O₂⁻. The removal of a valence electron in the CO molecule will create a vacancy in the 5σ orbital, implying that the first peak corresponds to transitions into this orbital. The peak at 533.8 eV is the 2pπ* peak that has shifted to lower energy from 534.2 eV in CO, with the corresponding carbon 2pπ* peak shifting 2.6 eV to higher energy. This indicates that the positive charge of the CO⁺ ion is mainly at the carbon site.

### 4.4. Molecules with Three Atoms

**Renner Teller Effect.** The breakdown of the adiabatic approximation in triatomic molecules is called the Renner Teller effect. The experimental manifestation of this effect is an irregular vibrational structure of the spectra that reflects the coupling of two electronic states in the bent nuclear conformations which were degenerate at the linear molecular geometry. The Renner Teller effect is a consequence of the electrostatic interaction between two components of an electronic state with a nonzero angular momentum.

In this section, we discuss some molecular systems with three atoms. The general interpretation of the oxygen K-edge remains similar to the binary molecules, that is, one observes the lowest empty orbitals, and at higher excitation energies, they are mixed with Rydberg states. Figure 13 shows the molecular orbitals of O₃, CO₂, and NO₂.

O₃. Ozone, O₃, is a bent molecule with a central oxygen atom and two equivalent end oxygens. This creates a new situation compared to O₂ because it is now possible to excite a 1s core electron from two different oxygen atoms. The molecular orbitals of O₃ can be divided into π orbitals and σ orbitals. The π orbitals are for bonding, nonbonding, and antibonding combinations, with only the 2pπ* antibonding orbital empty. At higher energy, there are two 2pσ* antibonding orbitals. The O₃ oxygen K-edge spectrum shown in Figure 14 shows two peaks at 529 and 536 eV are excitations into the same 2pπ* state, from, respectively, the terminal O₇ and the center oxygens O₅. Note that the small peak at 531 eV is due to the presence of some O₂.

#### Table 3. Energy Position of the 2pπ* Peak of Binary Molecules and Ions

| Molecule | 2pπ* peak energy (eV) | Ref. |
|----------|-----------------------|------|
| O₂⁻      | 529.0                 | 176  |
| O₂       | 530.8                 | 158  |
| O₂⁺      | 533.2                 | 187  |
| NO       | 532.7                 | 158  |
| CO       | 534.2                 | 170  |
| CO⁺      | 533.8                 | 188  |

Figure 13. Molecular orbital diagrams of CO₂, NO₂, and O₃. The MO schemes are equivalent to the difference that the number of electrons in the 2pπ* orbital is respectively 0, 1, and 2.

Figure 14. Oxygen K-edge TIXY spectrum of ozone. The peaks at 529 and 536 eV are excitations into the same 2pπ* state, from, respectively, the terminal O₁ and the center oxygens O₅. Note that the small peak at 531 eV is due to the presence of some O₂.
sharp peaks appear at 529 and 536 eV. The first sharp peak has been ascribed to the transition from the 1s electron of the terminal oxygen to the 2p\(\pi^*\). The second sharp peak at 536 eV is ascribed to transitions from the 1s electron of the central oxygen to the 2p\(\pi^*\). Note that both the peaks at, respectively, 529 and 536 eV originate from the same molecular orbital, but still they appear at 7 eV difference in their excitation energy because the terminal oxygen atoms are much lower in energy. In other words, the central oxygen atom is more positively charged. The complete analysis shows that the second peak also has some intensity from transitions into the 2p\(\pi^*\) antibonding orbitals for the terminal oxygens. The peaks at 540 to 545 eV are due to transitions into the (mixed) 2p\(\sigma^*\) antibonding orbitals and to Rydberg transitions. To summarize, it is interesting to compare the assignments for the oxygen atom and the \(\text{O}_2\) and \(\text{O}_3\) molecules:

- The oxygen atom only has the main 2p peak and at higher energy the Rydberg states.
- The \(\text{O}_2\) molecule has different chemical states, respectively, the 2p\(\pi^*\) and 2p\(\sigma^*\) peaks.
- The \(\text{O}_3\) molecule has different oxygen atoms, respectively, at the center and at the edge of the \(\text{O}_3\) molecule, yielding different peaks related to the same 2p\(\pi^*\) orbital due to different core–hole potentials.

\(\text{NO}_2\). Nitrogen dioxide is a bent molecule with a central nitrogen atom and two equivalent end oxygens; in other words, it has a similar structure to \(\text{O}_3\). This implies also that the molecular orbitals of \(\text{NO}_2\) are similar to those of \(\text{O}_3\) and the \(\pi\) orbitals form bonding, nonbonding, and antibonding combinations. Because there is one electron less, the nonbonding \(\pi\) orbital contains only one electron. The oxygen K-edge of \(\text{NO}_2\) contains two sharp peaks, respectively, related to the nonbonding and antibonding \(\pi\) orbitals. At higher energy, we observe broad structures due to the \(\pi\) orbitals and Rydberg states. It is interesting to note that both \(\text{NO}_2\) and \(\text{O}_3\) have two \(\pi\) related peaks, but for \(\text{NO}_2\), they relate to two different orbitals, while for \(\text{O}_3\), they relate to the same orbital but a 1s core–hole on two different oxygen atoms, center, respectively, end.

\(\text{CO}_2\). Carbon dioxide is a linear molecule with the bonding and nonbonding 2p orbitals occupied. It is a closed shell system that has a ground-state term symbol \(^1\Sigma_g^+\). The lowest three empty states are, respectively, the double degenerate 2p\(\pi^*\) state, followed by two antibonding 2p\(\sigma^*\) states. The 2p\(\pi^*\) state is doubly degenerate, but upon the oxygen K shell excitation, the equilibrium bond angle is changed and the degeneracy is removed by the Renner Teller effect. Because of the energy lowering resulting from bending the structure, the 2p\(\pi\) splits in two components. The first peak in Figure 15 corresponds to the promotion of an oxygen core electron to the two component of the lowest empty molecular orbital 2p\(\pi\).

This peak presents a large fwhm compared to the elastic peak, which suggests that not only the excitation from the K shell to the 2p orbital contributes to this peak but also the lowest 1s\(\sigma_u\) → 3s\(\sigma_u\) Rydberg transition with antibonding valence character. This hypothesis is supported by the fact that this transition is optically allowed. The peaks in the higher energy part of the spectrum can be assigned to Rydberg transitions. Thus, all the \(\pi\), \(\sigma\), and \(\pi\) transitions are allowed. This can lead to the conclusion that a mixing of the valence state with the \(\sigma^*\) antibonding resonance occurs. Shieh et al. study the oxygen K-edge of \(\text{CO}_2\) under pressure. The \(\pi\) peak disappears at 37 GPa and it is replaced by two structures at, respectively, 532 and 540 eV. At pressures above 37 GPa, \(\text{CO}_2\) does not exist as separate molecules, but has polymerized. There are several options for the condensed phase of \(\text{CO}_2\), with the phase where carbon is fourfold coordinated can best explain the observed spectral changes.

### 4.5. Atomic Adsorption

Molecules adsorbed on a surface change their electronic structure, and the oxygen K-edge is a useful tool to study the adsorption processes of oxygen-containing molecules. The adsorption phenomena of small molecules are particularly important as the fundamental steps in catalytic processes.

The interaction of oxygen atoms with a surface is strong due to the bonding of the oxygen 2p orbitals with the metal orbitals near the Fermi level. Three 2p orbitals are involved in the surface chemical bond with an adsorbing atom. The interaction with the 2p\(\pi\) orbital has \(\pi\) symmetry with respect to the surface. The 2p\(\sigma\) and 2p\(\pi\) orbitals are parallel to the surface and can only interact with the surface s atomic orbitals in high coordination sites. This bonding interaction contributes significantly to the total surface bond energy, implying that bonding to high coordination sites is favored. The 2p orbitals form bonding and antibonding orbitals with the metal orbitals.

Figure 16 shows the oxygen K-edge of oxygen adsorbed on a \(\text{Cu}(100)\) surface (points) compared with the energy-shifted copper 2p XAS spectrum of copper metal. The atomic energy of the 2p orbital is below the Fermi level, implying that the antibonding combinations will have mainly metal character. The oxygen K-edge in a first approximation maps the empty metal states, slightly shifted to higher energy due to the antibonding character. From this analysis, one would expect that if an oxygen atom is adsorbed on a copper surface, in the first approximation...
and the diatomic plane. This yields two polarization dependent spectra, sample is turned with the electric vector perpendicular to the plane, bonds perpendicular to the plane are excited when the X-ray polarization vector lies in this plane. Assuming a planar molecule lying on a surface, X-ray excites the electric dipoles that are aligned in this plane. At a bending magnet beamline, the X-ray is linearly polarized in the plane of the synchrotron. This implies that the shape of a system as a function of its angle with respect to the X-ray will also be linear dichroism, which then is called magnetic linear dichroism (MLD).

## 4.6. Molecular Adsorption

X-ray absorption of the adsorption of molecules to surfaces has been discussed in detail in the book *Chemical bonding at surfaces and interfaces* by Nilsson, Pettersson, and Norskov. The adsorption of oxygen-containing molecules to a surface is weaker than that of oxygen atoms. In a molecule, the atom that binds to the surface is still bonded to other atoms from the molecule. The adsorbed atom divides its bonding strength between the molecule and the surface, with the consequence of weaker bonding than an isolated oxygen atom. One can distinguish physisorption and chemisorption. In chemisorption, there is a chemical bond between the molecule and the surface, while in physisorption, the molecule is attached to the surface without a chemical bond, for example, via van der Waals interactions.

**Linear Dichroism.** Linear dichroism or polarization dependent X-ray absorption concerns the difference in XAS spectral shape of a system as a function of its angle with respect to the X-rays. At a bending magnet beamline, the X-ray is linearly polarized in the plane of the synchrotron. This implies that the X-ray excites the electric dipoles that are aligned in this plane. Assuming a planar molecule lying on a flat surface, one can excite the planar bonds if the X-ray polarization vector lies in this plane. The bonds perpendicular to the plane are excited when the sample is turned with the electric vector perpendicular to the molecular plane. This yields two polarization dependent spectra, and the difference spectrum is the linear dichroism spectrum. If the symmetry of the system is broken by a magnetic field, there will also be linear dichroism, which then is called magnetic linear dichroism (MLD).

**CO on Metal Surfaces.** The bonding of CO on Ni(100) was studied in detail in 1982 by Stöhr and Jaeger. One observes the transitions to the $2\pi^*$ orbital at 534.0 eV and to the $\sigma^*$ orbital at 550.0 eV. In the free CO molecule, these excitation energies are, respectively, 534.2 and 550.9 eV (see Table 3), implying that the $2\pi^*$ excitation energy does not shift, while the $\sigma^*$ orbital shifts by 0.9 eV. Thus, the energy difference between the $2\pi^*$ and the $\sigma^*$ orbitals is reduced from 16.8 eV in free CO to 16.0 eV in CO/Ni(100). This energy reduction can be ascribed to the shift to higher energy by the antibonding combination of the $2\pi^*$ orbital and the Ni 3d orbitals. Figure 17 shows an angular dependence. At normal incidence, the $2\pi^*$ peak is strong because the electric field is aligned with the $2\pi^*$ resonance. At grazing incidence, the electric field is parallel to the CO axis, hence perpendicular to the $2\pi^*$ orbital, and the $2\pi^*$ resonance is small. The argument is reversed for the $\sigma^*$ orbital that has the highest intensity when the electric field is parallel to the CO axis. This angular dependent effect, also known as linear dichroism, indicates the bonding orientation of the adsorbed molecule.

Tillborg et al. studied the oxygen K-edge XAS spectra of CO adsorbed on three different sites on a Ni(100) surface, respectively, hollow sites, bridge sites, and on-top sites. The energy position and relative intensity of the oxygen K-edge indicates the difference in chemical bonding.

## 4.7. Analysis of Catalytic Reactions

The information on the adsorbed species can be used in the analysis of oxygen K-edges of catalytic reactions. One can study the oxygen K-edge XAS spectra under in situ conditions. The most common approach uses conversion electron yield; i.e., the excited electrons ionize the gas phase and the resulting ionized ions are detected. Often, these experiments are performed in so-called ambient-pressure XPS setups, implying that the gas pressure is in the millibar range, but one could also use 1 bar pressures in dedicated in situ XAS reactors. Knop-gericke et al. studied methanol oxidation to formaldehyde over copper metal. One detector tracks the oxygen components in the gas phase, and a second detector tracks the oxygen components of the surface and gas phase combined, allowing the determination of the surface species by subtracting the pure gas phase spectrum. The oxygen K-edge as a function of pressure being discussed in detail by Nilsson, Pettersson, and Norskov in their book *Chemical bonding at surfaces and interfaces*.
temperature indicates a transition from Cu$_2$O at 540 K to a species that is described as adsorbed oxygen atoms that mainly interact with the copper s states. The different oxygen species are then correlated with the catalytic reactions.

Pfiefer et al. performed a detailed study on the comparison between amorphous and crystalline rutile IrO$_2$, which is important regarding its behavior as electrocatalyst for the oxygen evolution reaction. The rutile phase is a pure Ir$^{IV}$/O$^{2−}$ oxide and the oxygen K-edge shows two peaks: a sharp peak at 530 eV related to the empty Ir 5d states and a broad peak at 533 eV due to the sp band. The amorphous IrO$_2$ shows additional fine structure in the 5d spectral region, which in the paper is assigned to partial reduction to Ir$^{III}$, due to the formation of Ir vacancies. In our opinion, a shift to lower energy in the oxygen K-edge should be related to an oxidation to Ir$^{IV}$; in other words, the Ir vacancy creates Ir$^{III}$ sites at neighboring positions, which causes a shift to lower binding energy.

Tesch et al. studied the behavior of MnO$_2$, electrocatalysts as a function of the potential. The oxygen K-edge of the MnO$_2$ system changes as a function of the potential, which is analyzed as a change in the Mn−O hybridization. In our opinion, what is visible in the oxygen K-edge is mainly the change of the ratio of MnO$_2$ and nafion/graphene background, and the oxygen K-edge related to the MnO$_2$ species remains the same for all potentials.

5. LIQUIDS AND SOLUTIONS

5.1. Water

H$_2$O is studied in the gas, liquid, and solid phases. In this section, we briefly discuss all the states of water. The oxygen K-edge of water is a very active research field and we do not intend to cover all developments but instead focus only on the basic components in the analysis. Readers are referred to recent reviews on the X-ray spectroscopy of water, in particular, the recent reviews by Smith and Saykally and by Nilsson and Petterson.

Gas-phase H$_2$O is a bent molecule with five occupied orbitals. The highest occupied molecular orbital is a nonbonding 1b$_1$ orbital derived from the oxygen 2p$_x$ orbital without contributions from the hydrogen 1s orbital; in other words, it is the “lone pair”. At lower energy, the orbitals are, respectively, from highest to lowest energy the 3a$_1$, 1b$_2$, and 2a$_1$ molecular orbitals with contributions from hydrogen 1s and oxygen 2s and 2p orbitals, i.e., participating in the O−H bond. The two lowest unoccupied molecular orbitals are the O−H 4a$_1$ and 2b$_2$ antibonding orbitals.

Water molecules in liquid and solid phases show two different types of O−H interactions: (1) covalent intramolecular O−H bonding and (2) weak intermolecular hydrogen bonds. The study of the oxygen K-edge allows the investigation of the unoccupied states; most importantly, the antibonding O−H orbitals and their character that is sensible to the H-bonding network around the probed oxygen atom. The oxygen K-edge spectrum of water is considered as being formed by three regions: the pre-edge region at 535 eV, the main edge region around 537 eV, and the post-edge region at 541 eV. These regions go through modifications depending on the water phase. In Figure 18, the oxygen K-edge spectra of ice-, liquid-, and gas-phase water are compared.

The gas-phase spectrum shows the antibonding orbitals in the molecular MO scheme. In the case of the liquid state, the pre-edge has been ascribed to the presence of broken H-bonds by computational and experimental studies and the main edge region has been ascribed to interstitial water molecules. The post-edge region contains features that consist of states delocalized through the H-network and results from a tetrahedral ice-like coordination. Cavalleri et al. investigated the spectral change for the pre- and main edge going from the free molecule to the pentamer. They highlighted the role of the H-bond directionality on the spectral shape: the unoccupied antibonding O−H orbitals, in gas phase, are polarized toward the hydrogen, so they have a strong sensitivity to additional H-bonds. The features that constitute the post-edge region have been related to states that are localized along the H-bond. These states give rise to the conduction band in the ice phase with a strong intensity due to the fact that both OH groups are involved in donor hydrogen bonds. The three different regions of the water spectra are affected by several factors such as a change in temperature, different solvents, and isotope effects. Heating the liquid will cause the post-edge region to lose intensity, while the pre-edge and the main edge will increase their intensity. The study of Wernet et al. gave rise to a huge debate on the local structure of liquid water at different temperatures. It has been concluded that the spectrum of liquid water is composed of two different species: a species with two hydrogen-bonded configurations with one strong donor and one strong acceptor bond donating H-bond and a second species that is more symmetric with a tetrahedral-like coordination close to the ice coordination. By increasing the temperature, the water structure diverges from the tetrahedral-like coordination.

In addition, different solvents such as benzene, acetonitrile, and chloroform can be seen as hydrogen bonding network perturbations that allow symmetrical bonding and promote ice-like structures with a consequent increase in the post-edge structure. Fuchs at al. observed a significant isotope influence on the spectra of liquid water; the pre-edge peak shows a blue shift in the D$_2$O spectra due to the different zero point energies of the ground states and a reduced intensity in the pre-
edge region of the deuterated species due to smaller distortion of the bonding network.

5.2. Bio-Organic Molecules

The general interpretation of the oxygen K-edges of large molecules is similar to the small molecules discussed above. The peaks in the oxygen K-edge are analyzed in terms of the lowest empty orbitals, where in the case of two or more nonequivalent oxygen atoms in a molecule, the individual spectra of the oxygen atoms are added, taking into account that there can be different excitation energies into the same orbital. A large number of oxygen K-edge spectra is given in the COREX database.209

XAS studies of bio-organic molecules are difficult due to the risk of beam damage.210 The main degradation processes that can occur when measuring the oxygen K-edge can be grouped as: (a) dehydrogenation due to the O−H group, (b) dehydration, i.e., the loss of water molecules because of the C−OH bond, (c) decarboxylation (loss of molecular CO2), and (d) decarboxylation, the loss of molecular CO. Zubachius et al.210 investigated the radiation induced modification in the oxygen K-edge spectra of cysteine (Figure 19). The spectral shape is less affected by the beam damage than the intensity; the spectra recorded after 17 min of exposure have an intensity of 1 order of magnitude lower than that of the pristine system. This implies that at least 90% of the oxygen present in the molecule has been detached as H2O or CO2. Radiation damage can manifest itself in the oxygen K-edge measurements also as perturbation of the supramolecular structure.211,212 Panajotov et al. investigated the radiation damage on phospholipidic membranes, and they tested the impact of different incidence modes on the spectra: many broken σ bonds, especially in normal mode, and new features appeared.213 Other common chemical modifications induced by ionizing radiation are the formation of free radicals,214,215 where the reduction product is usually formed by the addition of an electron to the carbonyl oxygen of the carboxyl group214,216−218 and mass loss phenomena,219,220 depending on factors such as the initial mass, the dose rate, and the area of irradiation.219

Amino Acids, Polypeptides, and Proteins. The measurement of the oxygen K-edge XAS in amino acids is a powerful tool, since oxygen can be in different bonding environments, and different chemical shifts for each oxygen in amino acids are observed.221

In Figure 20, the oxygen K-edge spectra of all 22 amino acids are reported.222 The spectra show a dominant peak that has been assigned to the π* resonance followed by the σ* resonance at higher energy. In the aspartic acid spectra, a peak around 534.0−534.5 eV, with an energy difference from the main peak of 1.5 eV has been reported and assigned to the 1s → π* excitation of the OH oxygen in the carboxy group.221−223 In addition, the glutamic acid shows a similar feature.222 From the hydroxyl group of serine, threonine, and hydroxyproline, a σ* (O−C) transition occurs in the range ~538−540 eV.222 A probable contribution from residual water is expected in the energy range 537−540 eV.222,224

Even if isolated amino acids in the gas phase are always neutral, in solution any change in pH can bear dramatic changes in their charge state, as they can exist as cations at low pH values, charge neutral zwitterions at intermediate pH values, and anions in basic solutions. The addition or removal of a proton affects the local symmetry of the electric field surrounding the terminal oxygen,224 so the energy levels of molecular orbitals at the carboxylic group would be modified by any pH change.

In Figure 21, the spectra in solution of glycine at different pH values is reported. The terminal carboxyl group of glycine is protonated in an acid environment, so a lift of the degeneracy of the methyl group.226 In addition, the uracil spectra has a splitting in the range ~538−540 eV.222,224

**DNA Components.** In Figure 22, the oxygen K-edge XAS spectra of uracil (U), thymine (T), cytosine (C), and guanine (G) are reported.225 The bases show a strong feature around 532 eV arising from the carbonyl group (π*C=O), and a σ* resonance above 535 eV, thymine shows a splitting of the π* main resonance due to an energy difference of the two C=O bonds.226−228

In addition, the uracil spectra has a splitting in the first π* peak, but less pronounced; the stronger blue shift of the π*C=O of thymine is related to the substitution effect of the methyl group. DNA is built up by nucleobases attached to ribose sugar and phosphate groups. The way the sugar and the phosphate group affect the spectra of the oxygen K-edge of thymine is shown in Figure 22.
With the addition of sugar and phosphate, the spectrum acquires features characteristic of the phosphate group as the enhancement of the $\sigma^*$ ($C-O$) structure. The small peak in the $\pi^*$ region comes only from the base. 

6. SOLID OXIDES OF S- AND P-ELEMENTS

6.1. Alkali Metal Oxides

The alkali metals lithium, sodium, potassium, rubidium, and cesium form a number of different oxides. Taking lithium as an example, lithium forms, respectively, the oxide $\text{Li}_2\text{O}$, the peroxide $\text{Li}_2\text{O}_2$, and the superoxide $\text{LiO}_2$. The superoxide $\text{MO}_2$ can be considered as a $\text{M}^+$ ion combined with a negatively charged oxygen molecule $\text{O}_2^-$. The molecular orbitals of $\text{O}_2^-$ are similar to those of $\text{O}_2$ with one more electron occupying the antibonding $\pi^*$ orbital leaving only one hole. The extra charge on the oxygen molecule makes it easier to excite a 1s core electron to the empty orbitals. Figure 23 shows the spectra of superoxide $\text{MO}_2$ ($\text{M} = \text{Li, K, Cs}$) compared to the molecular $\text{O}_2$. The spectra of the superoxides show a first peak assigned to the $\pi^*$ orbital and two peaks assigned the $\sigma^*$ orbital and the ionization potential. The three features shift toward lower energy for the different superoxides. Pedio et al. also measured the superoxide $\text{KO}_2$. The experimental spectrum is equivalent to that shown in Figure 23, though with a $\pi^*$ peak at 531.0 eV instead of 529.0 eV. Kang et al. reproduce the oxygen K-edge of $\text{KO}_2$ from band structure calculations using Wien2K.

The $\text{M}_2\text{O}$ oxides ($\text{M} = \text{Li, Na, K, Rb}$) crystallize in the cubic antifluorite structure; $\text{Cs}_2\text{O}$ crystallizes in hexagonal anti-$\text{CdCl}_2$ structure. Alkaline ions occur mainly as cations $\text{M}^+$ and have rather weak bonds. Lithium oxides are an important compounds of many battery systems. This includes lithium—air batteries, where the reactions between lithium and oxygen play an important role. The reversible oxidation of lithium at the anode and reduction of oxygen at the cathode are important.

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**Table: Oxygen K-edge assignments for amino acids**

| Energy position (eV) | Assignment | Occurrence |
|----------------------|------------|------------|
| 532.3-532.5          | $\pi^*$ (COO)/(COOH)/(CONH$_2$) | All         |
| 534.0                | $\pi^*$ (C-OH) | Asp, Glu   |
| 542-549              | $\sigma^*$ (COO)/(COOH)/(CONH$_2$) | All         |
| 538-539.4            | $\sigma^*$ (C-OH) | Tyr, Thr, Hyp, Ser |

Figure 20. Experimental oxygen K-edge spectra of the 22 amino acids and their molecular formulas. In the table, the peak positions together with their assignments are reported.

Figure 22. With the addition of sugar and phosphate, the spectrum acquires features characteristic of the phosphate group as the enhancement of the $\sigma^*$ ($C-O$) structure. The small peak in the $\pi^*$ region comes only from the base.
parameters that influence the energy capacity of the battery system. Li$_2$O and Li$_2$O$_2$ have been studied as key components forming at the solid electrolyte interphase in lithium batteries, and their evolution was studied with the oxygen K-edge by Qiao et al.\textsuperscript{231} Figure 23 shows the spectra of Li$_2$O and Li$_2$O$_2$. Li$_2$O$_2$ shows a first peak assigned to the $\sigma^*$ orbital of the O—O bond. It is found that both Li$_2$O$_2$ and Li$_2$CO$_3$ evolve toward Li$_2$O under the soft X-ray irradiation, where Li$_2$CO$_3$ is very sensitive.\textsuperscript{231}

Figure 21. Oxygen K-edge spectra of 0.6 M glycine (aq) as a function of pH. A small red shift is observed for the acidic solution, which has been illustrated by a dotted line through the center of each peak. This shift is caused by the protonation of the carboxylate subgroup at low pH, and the resultant breaking of the degeneracy in the O 1s $\rightarrow \pi^*$ transition.\textsuperscript{224}

Figure 22. (a) Oxygen K-edge spectra of uracil, thymine, cytosine, and guanine together with their molecular structure and the peak assignment.\textsuperscript{225} (b) Effect on the oxygen K-edge spectra from adding sugar and phosphate groups of DNA to thymine.\textsuperscript{226}

Figure 23. Oxygen K-edge spectra of the oxide Li$_2$O, the peroxide Li$_2$O$_2$,\textsuperscript{231} and the superoxides LiO$_2$, CsO$_2$, and KO$_2$\textsuperscript{176} compared to O$_2$. The $\sigma$ and $\pi$ character of the antibonding states are indicated.
6.2. Alkaline Earth Oxides

Alkaline earth metals combined with oxygen form binary oxides with a rock salt structure and a well ordered surface. Figure 24 shows the oxygen K-edge of the alkaline earth oxides MgO, CaO, SrO, and BaO, from refs 232–236.

Figure 24 shows some discrepancies between the experimental data182,234,235 for CaO, SrO, and BaO. The discrepancies between the different sets of data can be linked to the high reactivity of the alkaline earth metal oxide surfaces. The surfaces of these materials, when exposed to air, are covered with CO2, water, and, in some cases, dioxygen.238 No details on how the materials were handled prior to the measurements have been reported;232,234 hence, possible surface contamination cannot be ruled out. In the experimental spectra reported by Nakai232 and McLeod,234 the absorption threshold from CaO to BaO does not change significantly, while the MgO spectrum shows a red shift of about 4 eV with respect to the oxides with the heavier metals. Significant changes can be observed in the near-edge structure going from MgO to BaO: the intensity of the first peak increases from MgO to SrO and the shape of the first peak changes from CaO to BaO.

Mizoguchi et al.239 analyzed the oxygen K-edge of MgO trough band structure calculations. The first feature at 532 eV is assigned to the Mg 3s band and the main structure 537–540 eV is assigned to the magnesium 3p band. In the MgO experimental spectra by Lindner,235 a small peak around 532 eV appears in the MgO oxygen K-edge, and a theoretical study from Pascual et al.240 has shown that a small peak around 532 eV appears in the MgO oxygen K-edge. While the participation of 3d orbitals from magnesium in the bonding with oxygen has been ruled out because of their high energy position,241,242 the participation of 3d orbitals is important for the heavier alkaline earth oxides.243

The electronic structure of solid oxides differs from that of molecules, which have molecular orbitals with fixed energy. In a solid, the electronic states form bands that continuously span a certain energy range, which is due to the presence of translational symmetry in crystalline solids. Because of the translation symmetry, it is conceptually and computationally easier to describe solids in reciprocal space. In band structure calculations, the electronic states are described as a series of bands that are given as a function of their (q) wave-vector in the Brillouin zone. The oxygen 1s core state is localized, implying that it has the same energy in all q-vectors. Because the X-ray photon has negligible momentum, X-ray absorption is effectively a vertical transition in the band structure. This implies that the oxygen K-edge spectrum integrates the band structure to the density-of-states. Because the oxygen 1s core state is localized at oxygen, the local density-of-states is visible, and because of the dipole selection rule, only oxygen projected states of p-symmetry have nonzero matrix elements. In other words, an oxygen K-edge of a solid maps the oxygen p-projected density-of-states from that of molecules, which have molecular orbitals with fixed energy. In a solid, the electronic states form bands that continuously span a certain energy range, which is due to the presence of translational symmetry in crystalline solids. Because of the translation symmetry, it is conceptually and computationally easier to describe solids in reciprocal space. In band structure calculations, the electronic states are described as a series of bands that are given as a function of their (q) wave-vector in the Brillouin zone. The oxygen 1s core state is localized, implying that it has the same energy in all q-vectors. Because the X-ray photon has negligible momentum, X-ray absorption is effectively a vertical transition in the band structure. This implies that the oxygen K-edge spectrum integrates the band structure to the density-of-states. Because the oxygen 1s core state is localized at oxygen, the local density-of-states is visible, and because of the dipole selection rule, only oxygen projected states of p-symmetry have nonzero matrix elements. In other words, an oxygen K-edge of a solid maps the oxygen p-projected density-of-states from that of molecules, which have molecular orbitals with fixed energy. In a solid, the electronic states form bands that continuously span a certain energy range, which is due to the presence of translational symmetry in crystalline solids. Because of the translation symmetry, it is conceptually and computationally easier to describe solids in reciprocal space. In band structure calculations, the electronic states are described as a series of bands that are given as a function of their (q) wave-vector in the Brillouin zone. The oxygen 1s core state is localized, implying that it has the same energy in all q-vectors. Because the X-ray photon has negligible momentum, X-ray absorption is effectively a vertical transition in the band structure. This implies that the oxygen K-edge spectrum integrates the band structure to the density-of-states. Because the oxygen 1s core state is localized at oxygen, the local density-of-states is visible, and because of the dipole selection rule, only oxygen projected states of p-symmetry have nonzero matrix elements. In other words, an oxygen K-edge of a solid maps the oxygen p-projected density-of-states from that of molecules, which have molecular orbitals with fixed energy. In a solid, the electronic states form bands that continuously span a certain energy range, which is due to the presence of translational symmetry in crystalline solids. Because of the translation symmetry, it is conceptually and computationally easier to describe solids in reciprocal space. In band structure calculations, the electronic states are described as a series of bands that are given as a function of their (q) wave-vector in the Brillouin zone. The oxygen 1s core state is localized, implying that it has the same energy in all q-vectors. Because the X-ray photon has negligible momentum, X-ray absorption is effectively a vertical transition in the band structure. This implies that the oxygen K-edge spectrum integrates the band structure to the density-of-states. Because the oxygen 1s core state is localized at oxygen, the local density-of-states is visible, and because of the dipole selection rule, only oxygen projected states of p-symmetry have nonzero matrix elements. In other words, an oxygen K-edge of a solid maps the oxygen p-projected density-of-states from that of molecules, which have molecular orbitals with fixed energy.
the solid can be studied with X-ray absorption using EXAFS, where the EXAFS experiments are usually performed on the hard X-ray cation XAS spectra. Furthermore, amorphous materials do not have a periodic structure and XAS is a key method to determine the local structure of the atoms. The high covalency of the cation-oxygen bond has been a motivation for the investigation of the geometric and electronic structures from the oxygen point of view.

The oxygen 2p orbitals form bonding and antibonding bands with the cation ns and np orbitals. The oxygen K-edge maps the empty states and as such shows the (oxygen 2p character of the) cation combined sp bands. Figure 25 compares the spectra of quartz and rutile polymorphs of SiO2 and GeO2, and rutile SnO2. The spectra show the dependence of the oxygen K-edge on the crystalline structure, the bond angles, and the resulting consequences for the mixing of the oxygen 2p orbitals with the ns, np, and nd orbitals. In the quartz structure, the cation is in fourfold tetrahedral coordination and the oxygen is twofold coordinated. In the rutile structure, the cation is in sixfold elongated octahedral coordination and the oxygen is threefold coordinated.

The multiple scattering calculations247,248 later confirmed by band structure calculations,249 have allowed the assignment of the oxygen K-edge features in α-quartz SiO2: the main intense peak at the edge is assigned to the oxygen 2sp hybridized with silicon 3s followed by features assigned to transitions to oxygen 2p hybridized with silicon 3pd states split by the crystal field. We note that the cation tetrahedral coordination allows the cation pd state mixing. The spectral shape of quartz-GeO2 is drastically different as a consequence of the different cation-oxygen bond angle (from 143.7° in SiO2 to 130° in GeO2) on the orbital overlap.250 However, in both cases, the general final state assignment is similar for SiO2 and GeO2.

Contrary to the quartz compounds, the oxygen K-edge spectra of SiO2, GeO2, and SnO2 in rutile structure, Figure 25c shows similar shapes: a first sharp peak followed by a broader structured feature. In the case of SiO2, these features are not as well separated as in GeO2 and SnO2. According to the calculated p-DOS,250,251 the first sharp feature corresponds to the oxygen 2sp antibonding σ* states hybridized with the empty cation ns states (Si: n = 3, Ge: n = 4, and Sn: n = 5). The strong intensity of this feature is related to the strong bonding of the oxygen in a threefold planar coordination. This can be described with a orbital view: in this geometry, the hybrid sp2 orbitals of oxygen point at the cation’s orbitals in a strong covalent σ bonding. The following broad feature is assigned to oxygen 2p states and oxygen 2p states hybridized with the cation np states. For germanium and Sn, the contribution at higher energy from (n − 1)d states is also found. The p-DOS calculations shows that the mixing of the cation p and d states as not allowed in the octahedral symmetry. The spectra of the rutile compounds shift to low energy along the series in agreement with the decrease of their respective band gaps.

In the following sections, we first discuss the most studied p-element compounds: borate, carbonate, aluminate, silicate, and germanate crystals and glasses. Then, we review briefly the other p-element compounds, including p-electron heavy metals.

6.4. Overview of p-Element Oxides

6.4.1. Boron Oxides. Boron and borate oxides are formed by two different structural units depending on the nature and content of alkali cations and/or the temperature and pressure conditions: planar BO3 triangles or BO4 tetrahedrons. The coordination of oxygen can be twofold bridging oxygens or onefold coordinated nonbridging oxygens. In glasses, the addition of alkalis induces the transformation from BO3 to BO4 and a higher connectivity between the boron units. Further increase of the alkali content induces the depolymerization of the glass or crystal network.252

The determination of the boron and oxygen coordination is thus a critical parameter to understand the properties of borate crystals and glasses in various temperature and pressure conditions. For these systems, the low penetration depth of soft-X-rays and surface sensitivity have been a serious limitation for their investigation, especially because of their particular hygroscopicity. IXS has been used as a good solution to circumvent these limitations. Leelong et al. have investigated of the oxygen K-edge XAS via IXS in a series of crystalline lithium borates253 and in comparison with glasses.252 Figure 26 shows the oxygen K-edge of selected boron oxides. The spectra show two main features: the first 5 eV of the oxygen K-edge spectra is
assigned to $\pi^*$ overlap between oxygen 2p and boron 2p, while the next 10 eV range is assigned to $\sigma^*$ O–B overlap. DFT calculations have further distinguished the contribution of bridging and nonbridging oxygens according to the energy position of the $\pi^*$-peak, which shifts to lower energy in the case of nonbridging oxygens as observed when comparing amorphous B$_2$O$_3$ and Li$_3$BO$_3$. Comparison between crystals and glasses has allowed a further understanding of the structural variations in lithium borate glasses with the chemical composition. Lee et al. have studied v-B$_2$O$_3$ glass at high pressure using IXS up to 8.4 GPa. Figure 26 shows the decrease of the $\pi^*$-peak related to boroxol rings, i.e., planar structure formed by three BO$_3$ units, upon increasing the pressure together with the increase of the boron coordination from three- to fourfold suggesting that the unpaired Op orbital gets involved in forming a $\sigma^*$ bond with another boron resulting in a threefold oxygen.

6.4.2. Carbonates. Carbonates are important compounds for the carbon cycle. Carbonate rocks result from the mineralization of the CO$_2$ from the atmosphere. Figure 27 shows the oxygen K-edge of BaCO$_3$, Li$_2$CO$_3$, CaCO$_3$, and Na$_2$CO$_3$. The oxygen K-edge XAS spectra of carbonate compounds are dominated by the oxygen antibonding states from the CO$_3^{2-}$ units, similarly to the BO$_3$ units for boron (see above). These planar triangular units favor the formation of in-plane $\sigma$ covalent bonds involving sp$^2$ mixing. The first intense and sharp peak is assigned to the $\pi^*$ antibonding state from the C=O bonds resulting from out-of-plane $\pi$ bonds involving only p states. The second part of the spectra corresponds to the $\sigma^*$ antibonding state.

Calcium carbonate, CaCO$_3$, is an important biomineral occurring in shells and corals. CaCO$_3$ occurs in three different structures, calcite, aragonite, and vaterite; in addition, there are hydrated and amorphous structures. DeVol et al. studied the oxygen K-edge of these forms of CaCO$_3$, using the polarization dependence of single crystals. The experiments are simulated with FEFF calculations. Despite similar isotropic spectral shapes, the oxygen K-edge polarization dependence varies with the crystal structure of the polymorph. The polarization dependent contrast is used to study natural species (shells and tunic spicules) with X-ray spectral microscopy. De Clermont Gallerande et al. analyzed the monopole and dipole contributions to the oxygen K-edge spectra of Li$_2$CO$_3$ measured with IXS, separating the $\pi$ and $\sigma$ antibonding states. The oxygen K-edge of Na$_2$Si$_2$O$_5$ shows a sharp peak at 533 eV that is assigned to Na$_2$CO$_3$ adsorbed at the surface. Na$_2$CO$_3$ and NaHCO$_3$ have been measured by Espinal et al. They are used as references to study the interaction of CO$_2$ with amine groups. Recently, the oxygen K-edge of barium carbonates and nitrates have been studied, in comparison with the binary oxide. Karshoglu et al. measured the oxygen K-edge of BaO, Ba(OH)$_2$, BaCO$_3$, and Ba(NO$_3$)$_2$ prepared in situ as thin films.

6.4.3. Aluminum Oxides. Aluminum is the third most abundant element of the earth crust after oxygen and silicon and is present as trivalent cations in various biological and mineralogical forms. The oxygen K-edge values of $\alpha$-Al$_2$O$_3$ (corundum structure), MgO, and MgAl$_2$O$_4$ (spinel structure) are compared with their calculated projected DOS by Kostmeier et al. Kaneko et al. study the difference between the oxygen K-edge of the center of a Al$_2$O$_3$ grain with the grain boundaries. The features correspond to the antibonding oxygen states hybridized with aluminum 3s, p, and sp states. Henderson et al. studied the oxygen K-edge of a series of crystalline calcium aluminates followed by the study of a series of CaO–Al$_2$O$_3$–SiO$_2$ mixed glasses. The oxygen K-edge spectra of the glasses reveal the contribution of both calcium and aluminum hybridized states and the coordination change of calcium as well as the degree of polymerization of AlO$_4$ tetrahedral units. The oxygen K-edges of Cu–Al–O thin films have been compared with copper oxide and Al$_2$O$_3$ references. The oxygen K-edge of crystalline $\alpha$-Al$_2$O$_3$ has been compared for different shapes.

![Figure 26. Oxygen K-edge spectra of amorphous B$_2$O$_3$ at 1 bar and 8.4 GPa, LiBO$_2$, Li$_2$B$_4$O$_7$, and Li$_3$BO$_3$ with the fraction of BO$_3$ units and nonbridging oxygens indicated. Nonbridging orbitals induce a shift of the $\pi^*$ feature to lower energy and the conversion from BO$_3$ to BO$_4$ induces the decrease of its intensity.](https://dx.doi.org/10.1021/acs.chemrev.9b00439)

![Figure 27. Oxygen K-edge spectra of carbonates: BaCO$_3$, Li$_2$CO$_3$, CaCO$_3$, and Na$_2$CO$_3$, all showing the $\pi^*$ antibonding state from the C=O bonds of the CO$_3^{2-}$ units.](https://dx.doi.org/10.1021/acs.chemrev.9b00439)
6.4.4. Silicon Oxides. SiO$_2$ is the first most abundant oxide of the earth crust$^{208}$ mainly as part of the mineral species like silica and silicates. Oxygen K-edge has been used to investigate the atomic structure of crystalline and amorphous silica and silicates. Marcelli et al. showed the sensitivity of the oxygen K-edge on the Si—O—Si bond angle in glasses, which stimulated intense efforts to the development of accurate modeling of the oxygen K-edge using multiple scattering approaches$^{247,266,267}$ and later followed by band structure calculations.$^{244,249}$ The experimental oxygen K-edge spectra of silica crystal polymorphs were investigated by Wu et al.$^{247}$ The influence of the chemical composition of silicate glasses on the structure was investigated.$^{245,268-270}$ The interpretation of the spectra is more complex when considering silicate glasses with also calcium, aluminum, and titanium. The oxygen K-edge spectra reveal the bonding with the different cations but is also influenced by the coordination geometry changes. Another important aspect of glass structure is the degree of polymerization of the glass network. The addition of alkali cations to amorphous SiO$_2$ induces the depolymerization of the SiO$_4$ network and formation of nonbridging oxygens (O bond to a single Si). It has been shown with the support of full multiple scattering calculations that the threshold of the oxygen K-edge is dominated by the presence of the nonbridging oxygens in silicates, similarly to the case of borates. The multiple scattering calculations,$^{247,248}$ and later DFT calculations,$^{249}$ have allowed reproducing the experimental data. Multiple scattering calculations on clusters of different sizes have allowed the contribution of the multiple scattering to the features due to medium range order. The effect of the core—hole has been discussed in these studies. While it is shown to be very important for the silicon K-edge,$^{251}$ the contribution to the oxygen K-edge has been described as small by Wu et al.$^{248}$ but it was shown to be non-negligible by Taillefumier et al.$^{249}$ These authors highlighted that some cautions should be used to compare the calculated oxygen K-edge and the oxygen 2p pDOS, in reference to the work of Mo et al.$^{244}$ which presented strong disagreement between the oxygen K-edge spectrum and pDOS calculations. While Mo et al. assigned this mismatch to the energy dependence of the dipole matrix element, Taillefumier et al. have pointed out the influence of the region used for the angular momentum projection.$^{249}$

6.4.5. Germanium Oxides. The oxygen K-edge of crystalline and amorphous germanates have been investigated to obtain structural information. Based on the study of crystalline references, oxygen K-edge has been used to determine the coordination of germanium in amorphous systems. The similarity of the oxygen K-edge of amorphous and crystalline (quartz) GeO$_2$ confirmed the twofold coordination of oxygen and the fourfold coordination of germanium in both cases as well as similar angles between the GeO$_4$ tetrahedrons.

Fivefold germanium is observed in the alkali germanate glasses but only in compositions at the germanate anomaly maximum and beyond. It was confirmed that there is no formation of sixfold coordinated germanium within the glass before the germanate anomaly maximum.$^{245}$ No study of oxygen K-edges in germanates has reported the contribution from nonbridging oxygens, even for high-alkali content glasses, although these are expected. Such an interpretation is indeed subject to great caution because of the surface sensitivity of XAS, but here, IXS could provide an alternative interesting probe. By comparison with the crystal references, Lelong et al. demonstrated with IXS measured oxygen K-edge the formation of the fivefold coordinated germanium at the intermediate high pressure (5 GPa) from amorphous germania and the formation of sixfold coordinated germanium at the highest pressure (18 GPa).$^{272}$ McLeod et al. used oxygen K-edge in light of calculations with Wien2k to study the oxygen vacancies in GeO$_2$ nanoparticles as electrodes for lithium storage.$^{273}$

6.4.6. Other p-Element Oxides. Zinc Oxides. ZnO is a common semiconductor oxide. The oxygen K-edge of the hexagonal wurtzite polymorph has been measured by McLeod et al.$^{234}$ Figure 28 shows the oxygen K-edge spectrum of ZnO.

![Figure 28. Oxygen K-edge spectra of ZnO (wurtzite) compared to Fe:ZnO (red), CdO (rock salt), and HgO (orthorhombic, montonydite).]  

Because all 3d orbitals are full, the spectrum reveals only the oxygen 2p states hybridized with the 4sp states of the Zn. Guo et al.$^{252}$ compare the bulk spectrum with a thin film and nanorods. ZnO doped with transition metal ions (Fe, Co) yields so-called dilute magnetic semiconductors. These systems have ferromagnetic order above room temperature and the combination of charge and spin degrees of freedom makes them useful for devices. Figure 28 shows that the doping with iron induces the presence of additional empty 3d states at the onset of the oxygen K-edge. Furthermore, the features observed for pure ZnO are broadened, probably due to the distortion of the crystal lattice. The oxygen K-edge of ZnO doped with iron and with cobalt mainly show the same spectral shape as pure ZnO with an additional peak due to the Fe—O and Co—O interaction.$^{275-277}$ The oxygen K-edges of the [Cd,Zn]O mixed oxides are discussed in the section on cadmium oxides.

Phosphates. Calcium phosphates are the main component of human bone. The materials include tricalcium phosphate Ca$_3$(PO$_4$)$_2$ and hydroxyapatite Ca$_5$(PO$_4$)$_3$OH. Rajendran et al. compare the calcium phosphate materials plus CaCO$_3$ and CaO with the oxygen K-edges from the bones of sheep, deer, chicken, etc.$^{279}$ They use TEY and FY to differentiate between surface and bulk; they find that the surface of dried bone has a larger proportion of carbonate than the interior that is mainly made up of phosphates. The phosphate systems with transition metal ions, (LiFePO$_4$, LiNiPO$_4$, etc.) are discussed in the sections of the respective transition metal ions.

Gallium Oxides. The gallium oxide Ga$_2$O$_3$ exists in five polymorphs (α to ε). The β-Ga$_2$O$_3$ trigonal structure has recently got much attention in relation to possible optoelec-
Ga2O3 has been calculated with the FDMNES code by Hegde et al. between 62% and 75% CdO. Figure 28 compares the spectra of wurtzite ZnO and rock salt CdO. The phase transition between the Sn p and pd bands. The monoxide yields and with thin films that are measured both with electron yield and with fluorescence yield. Detert et al. measured the oxygen K-edges of the [Cd,Zn]O mixed oxides for different ratios of cadmium and zinc. The phase transition between wurtzite ZnO and rock salt CdO is visible in the oxygen K-edge between 62% and 75% CdO. Figure 28 compares the spectra of ZnO and CdO.

Cadmium Oxides. CdO is a rock salt structured semiconductor. The oxygen K-edge spectrum is measured by Demchenko et al. McLeod et al. compare the oxygen K-edge with Wien2K calculations. The bulk spectrum of CdO is compared with thin films that are measured both with electron yield and with fluorescence yield. Detert et al. measured the oxygen K-edges of the [Cd,Zn]O mixed oxides for different ratios of cadmium and zinc. The phase transition between wurtzite ZnO and rock salt CdO is visible in the oxygen K-edge between 62% and 75% CdO. Figure 28 compares the spectra of ZnO and CdO.

Indium Oxides. Indium oxide In2O3 (cubic polymorph) is compared to the other 5s0 oxides by McLeod et al. as reproduced in Figure 29. The oxygen K-edges of the cubic, rhombohedral, and orthorhombic In2O3 polymorphs have been measured by de Boer et al. All polymorphs give similar oxygen K-edge spectra with a first intense sharp peak assigned to the oxygen 2p states hybridized to the cation empty 5s states and then a variable second broad feature corresponding to the metal 5p states. They calculated the spectra with Wien2K with and without core—hole potential. The final state calculations accurately reproduce the observed spectral shapes.

Tin Oxides. Cassiterite SnO2 has a rutile structure and is the common ore for tin. The oxygen K-edge is shown in Figure 29. It is shifted to lower energies but shows a similar shape to that of GeO2 and SiO2. This allows transposing the feature assignment from the germanium and silicon well studied systems: the first intense peak corresponds to the oxygen 2p hybridized with the σ* of the Sn 5s states and the higher energy features correspond to the Sn p and pd bands. The monoxide α-SnO has a PbO structure. Upon heating, it converts to an intermediate tin oxide SnOα before converting to SnO2. The EELS oxygen K-edges of the three different oxides SnO, SnOα, and SnO2 have been measured by Moreno et al. The oxygen K-edge of the intermediate SnOα phase has a shape that can be constructed from a linear combination of SnO and SnO2. Kronawitter et al. measured the oxygen K-edge of SnO2 and then calculated the spectrum with Wien2K. A Wien2K calculation was performed with a core—hole potential, but the agreement with experiment is not very good. We note that the oxygen K-edge of SnO2 is different as measured with EELS and with XAS. The EELS spectrum has two leading sharp peaks and the XAS spectrum only one. A possible explanation is that the EELS spectrum contains SnO that has its first peak at exactly the same energy. Kronawitter et al. studied the interface between SnO2 and TiO2. Their SnO2 spectra are measured with TEO XAS and agree with the spectra from Kronawitter.

Antimony Oxides. The oxygen K-edge of antimony oxides, shown in Figure 29, were studied as part of a broader investigation of chemical bonding and hybridization in 5p binary oxides by McLeod et al. The authors further compare 5s0 oxides (In2O3, SnO2, Sb2O5) with 5s2 oxides (SnO, Sb2O3, TeO2). Comparison of DOS and spectra calculations performed with Wien2k with experimental spectra reveals that in general the hybridization between oxygen 2p states and cations states decreases with increasing cation atomic number. In the 5s0 oxides, the first relatively broad feature of the oxygen K-edge is due to the antibonding oxygen 2p bands mixed with the cation 5p, while in the 5s2 oxides, the cation 5p band is shifted to high energy and the first sharp feature is due to the empty cation 5s band mixed with the oxygen 2p.

Tellurium Oxides. TeO2 is an important building block for telluride glasses used for nonlinear optics such as in optical fiber amplification. The oxygen K-edge of TeO2 measured by Jiang et al. with EELS is shown in Figure 29. The authors compare the spectrum with a number of different FEFF calculations. The core—hole calculation gives equivalent results to the Z+1 calculation. To improve the correlation with experiment, a Z+2

Figure 29. Oxygen K-edge spectra of (a) 5s0 oxides (In2O3, SnO2, Sb2O5) and (b) 5s2 oxides (SnO, Sb2O3, TeO2). Data are reproduced from ref 234 for all but TeO2. The cation contributions to the oxygen 2sp antibonding states are given according to the p-DOS calculations from McLeod et al. 246

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approximation is used, but the agreement obtained is still not very good. McLeod et al. have compared the Wien2k calculated oxygen 2p DOS with the experimental spectra. The projected DOS reveal that the first peak in the TeO₂ oxygen K-edge spectrum is dominated by oxygen 2p antibonding states hybridized with Te 5p, and the second peak is dominated by oxygen 2p states hybridized with tellurium 4d.

Mercury Oxides. Mercury is the third element of the group. Mercury oxide is a II–VI-type semiconductor. Divalent mercury ions have full 5d orbitals and 6s² and 6p⁰ valence electronic configuration. The oxygen K-edge of mercury oxide HgO, whose structure consists of chains of Hg–O in the montmorillonite form (orthorhombic), has been compared with Wien2K calculations. Figure 28 compares the spectral shape of HgO with CdO and ZnO. As the cation increases in mass, the core–hole has a decreasing effect on the local conduction band probed by XAS measurements. There is close agreement between the measured XAS and the calculated DOS for HgO.

Lead Oxides. Qamar studied four different types of PbO samples: α-PbO or litharge, the red tetragonal polymorph; β-PbO or massicot, the yellow orthorhombic polymorph; amorphous PbO; and polycrystalline PbO. They compare the experiments with Wien2K DFT calculations with and without the core–hole potential. Oxygen K-edges of lead oxides for lead acid batteries were published by Moseley et al. The oxygen K-edge of BaPbO₃ has been measured in comparison with calculations with and without core–hole. The whole series of Ba(Pb,Bi)O₃ perovskites has been measured as a function of the bismuth doping. A gradual change from BaPbO₃ to BaBiO₃ is observed experimentally (Figure 30). Mastelaro et al. used the oxygen K-edge to determine the hybridization between lead 6p and oxygen 2p orbitals in Pb₁₋ₓLaₓTiO₃ compounds. The lead 6p peak appears in between the two titanium 3d bands. A similar approach with the support of FEFF calculations was used in to study the ferroelectric Pb₁₋ₓBaₓZr₀.₆₅Ti₀.₃₅O₃ compounds.

Bismuth Oxides. Torruella et al. studied monoclinic α-Bi₂O₃ with EELS. They compared the spectral changes of a pellet with powder and nanowires, in particular, the effect of defects on the spectral shape. BaBiO₃ and the Ba(Pb,Bi)O₃ perovskite systems have been discussed in the section on lead. BiFeO₃ and BiVO₄ are discussed in the sections on the transition metal ions.

### 7. Solid Oxides of d- and f-Elements

Solid oxides from d- and f-elements are different from the s- and p-elements because of the presence of d- and/or f-electrons that have a stronger localization. One consequence of this local-
of the covalency on the trend of the 3d series and also on the
effect of the covalency on the spectral shape. Next, we discuss
binary oxides and their trends in the spectra as a function of
oxidation state.

**Crystal Field Splitting.** There is a close relation between the
energy difference separating the two peaks of the 3d band and
the ligand field splitting. In the case of 3d$^0$ systems and high-spin
3d$^0$ systems, there is no additional effect due to exchange, and
the relation between the peak distance and 10Dq should hold
exactly. In binary oxides, the crystal field splitting is
approximately 0.6 eV times the formal valence, which
would imply a crystal field of 1.2 eV in MnO and 2.4 eV in TiO$_2$, where
the actual values can be different by $\sim$0.2 eV. Comparing the
peak distance to the 10Dq value as determined from optical
excitations, one finds that the oxygen K-edge spectra yield a 10% shorter
energy distance.\(^{296}\) The final state core−hole localizes the
oxygen 2p states, thereby decreasing the overlap with the
metal 3d states. We note that this reduction is less than for the
metal 2p XAS spectra that have a finite overlap with the
metal 3d states.\(^{16}\) We note that this reduction is less than for the
metal 2p XAS spectra that have a final state 10Dq value that is
reduced by $\sim$20% via a similar core−hole localization effect.\(^{297}\)

**Exchange Splitting.** In systems with a finite spin, there is a
splitting between the spin-up and spin-down sub-bands. For
example, high-spin octahedral 3d$^0$ systems have three spin-up t$_{2g}$
electrons in their ground state. This creates an exchange splitting
between the empty high-spin and low-spin states. The exchange
splitting can be approximated as the number of unpaired spins
times 0.6 eV. This implies that in the case of Cr$_2$O$_3$ both the
exchange splitting and the crystal field are approximately 1.8 eV.

**Covalency and the Trend of the 3d Series.** Looking at the
oxygen K-edge of binary 3d oxides where the metal ions have an
octahedral surroundings, one finds a similar shape of the empty
states due to interaction of the metal 4s and 4p and oxygen 2p
states. The lowest energy states are the oxygen 2p states that are
hybridized with the empty 3d states. Assuming an equal
hybridization for all systems, the intensity of the 3d band yields
the number of empty 3d states. Analysis shows that one finds
approximately a linear relation between the intensity of the 3d
band and the number of empty 3d states,\(^{16}\) where the 3d band
intensity is normalized to the intensity of the 4sp band. Suntivich
et al. found for a series of perovskites that the covalency is not
constant but becomes larger through the 3D series.\(^{398}\) While
filling the 3d band, the energy position of the 3d states with
respect to the oxygen 2p states decreases, thereby increasing
their overlap. In addition, the average metal−oxygen distance
gets shorter, implying that both the O 2p−M 4sp and the O 2p−
M 3d interactions will increase. The average metal−oxygen
distance varies between different oxide systems; for example,
M$_2$O$_3$ corundum has larger metal−oxygen distances than
LaMO$_3$ perovskites. This variation can cause differences in the
intensity of the 3d part of the spectrum even for the same
element, valence, and spin-state.

**Covalency and Spectral Shape.** The covalency can be
further analyzed by including the difference between the
hybridization of the t$_{2g}$ and the e$_g$ states in octahedral symmetry.
The e$_g$ states point directly to the oxygen atoms and have $\sigma$
overlap. The t$_{2g}$ states point in between the oxygen atoms and
have $\pi$ overlap with the oxygen 2p states. Effectively, this yields a
difference in the hopping terms of 2:1, which implies a difference in
hybridization of 4:1 for each O 2p−M 3d overlap. Additional
factors for the intensity ratio are the fact that the $\pi$ bonding is
possible with two O 2p orbitals and $\sigma$ overlap with one O 2p
orbital. In the case of 3d$^0$ systems (TiO$_2$), there are six t$_{2g}$
and four e$_g$ states. The total (relative) intensity ratio for the
t$_{2g}$/e$_g$ band is $1/4 \times 2/1 \times 6/4 = 12/16$. This implies that one expects a
3:4 integrated intensity ratio for the t$_{2g}$ to e$_g$ band, which is close
to the observed ratio for TiO$_2$ and MnO. Figure 32 shows the
effects of degeneracy and overlap on the t$_{2g}$ and e$_g$ peaks of a 3d$^0$
-system. In the case of 3d$^1$ systems, one can assume that the
exchange splitting is smaller than the crystal field splitting, and
with this assumption, the t$_{2g}$/e$_g$ band ratio is 10:16.

The 3d$^1$ system has an exchange splitting that is 1.5−2.0 eV,
which implies that the spin-down t$_{2g}$ band overlaps with the spin-
up e$_g$ band, which complicates the two-band t$_{2g}$/e$_g$ analysis.
Figure 33 shows the effects of exchange splitting, degeneracy,
and overlap on the t$_{2g}$ and e$_g$ peaks of a 3d$^1$ system.

7.1.2. Binary Oxides and Oxidation State Trends.

We first discuss the trends of the oxygen K-edge through the 3d
series on three series of binary oxides, respectively, the rock salt
MO oxides, the corundum M$_2$O$_3$ oxides, and the rutile MO$_2$
oxides.

Figure 34 shows the divalent 3d transition metal oxides with a
rock salt structure. Respectively given are 3d$^5$ VO, 3d$^3$ MnO, 3d$^4$
FeO, 3d$^7$ CoO, and 3d$^8$ NiO, where these divalent 3d transition
metal oxides are always high-spin. They all have a crystal field
splitting of approximately 1.2 eV. The lowest energy peaks
relates to the empty 3d states, and the peaks around 540 eV
relate the metal 4s and 4p band. Rock salt has a typical sharp
band structure peak around 545 eV. In the case of MnO, the
empty 3d states are at relatively high energy at 534 eV and they
overlap with the 4s and 4p structures. MnO has a 3d$^5$
configuration, and in principle, one would expect two 3d peaks split by the crystal field. Because of the small crystal field splitting of 1.2 eV, compared with the band widths, the splitting is not visible in the spectral shape. FeO has a 3d⁶ configuration, and similarly to MnO, the crystal field splitting is too small to be visible. In the case of CoO, one could see the fine structure in the 3d peak, and the first small feature could be assigned to the empty t₂g state. NiO has a 3d⁸ configuration spin-down eg band is empty, which yields a sharp 3d peak.

Figure 35 shows trivalent 3d transition metal oxides with a corundum structure. Respectively given are 3d¹ Ti₂O₃, 3d² V₂O₃, 3d³ Cr₂O₃, 3d⁴ Mn₂O₃, 3d⁵ Fe₂O₃, and 3d⁶ Co₂O₃. All trivalent 3d transition metal oxides have a crystal field splitting of 1.8 eV and are high-spin, except Co₂O₃ that is low-spin. The 3d¹ system Ti₂O₃ has two peaks in the 3d part split by the crystal field. Trivalent oxides have a crystal field splitting of 1.8 to 2.0 eV, clearly visible in the oxygen K-edges. At 540−545 eV, the structure related to the metal 4s and 4p bands is visible. The 3d² system V₂O₃ has a similar spectral shape. In the case of the 3d³ system Cr₂O₃, the exchange splitting between spin-up and spin-down states is similar in size to the crystal, yielding a first intense peak related to a combination of the spin-down t₂g and spin-up e₈g band. The 3d⁴ system Mn₂O₃ is a Jahn—Teller system due to its half-filled spin-up e₈g state. The combination of (i) the octahedral crystal field, (ii) the exchange splitting, and (iii) the Jahn—Teller distortion yields a broad 3d-related feature. The 3d⁵ system Fe₂O₃ has a half-filled 3d band and shows two peaks split by the crystal field splitting. The 3d⁶ system Co₂O₃ is low-spin and has a filled t₂g band, hence a single sharp peak related to the empty e₈g band.

Figure 36 shows the tetravalent 3d transition metal oxides with a rutile structure. Respectively given are 3d⁰ TiO₂, 3d¹ VO₂, 3d² CrO₂, and 3d³ MnO₂, where these tetravalent 3d transition metal oxides have a crystal field splitting of 2.4 eV. TiO₂ shows two peaks split by the crystal field splitting, and the VO₂ spectrum is similar, but due to the presence of the vanadium L₂3-edge at 720 eV, the oxygen K-edge of VO₂ sits on a
background of the tail of this spectrum. The 3d² system CrO₂ is a well-known half-metal. The relatively large exchange splitting yields a first spin-up t₂g band, while the other sub-bands are all part of the second structure. The 3d³ system MnO₂ has a similar spectrum to Cr₂O₃, with the first peak a mixture of the spin-down t₂g and spin-up eg band.

The comparison of the three common valences 2+, 3+, and 4+ gives rise to the following observations:

Intensity of the 3d Part. The 3d part of the spectrum has more intensity for high-valence oxides. The reason is that the covalency increases when the valence is increased from 2+ to 3+ and 4+. Increased covalency implies more oxygen character in the metal 3d band and, as such, more intensity in the 3d part of the oxygen K-edge. Due to the increased covalency, the 4sp part of the spectrum will also gain intensity, but this effect is smaller than for the 3d part, because the 4s and 4p orbitals are already delocalized and gain less overlap than the 3d orbitals.

Excitation Energy. The excitation energy of the metal L₂,3- edges shifts to higher energy with increased valence. The simplest approximation is that it is due to the increased (effective) charge on the metal site, implying a larger energy needed to excite a core electron. In contrast, the energy position of the oxygen K-edge does not always shift in the same direction by changing the formal charge of the metal ion. Depending on the specific metal ion, the oxygen K-edge shifts to lower energy, to higher energy, or remains at the same energy.

Looking at the experimental data, there is no shift observed for the valence changes Ti⁴+/Ti³+, V⁴+/V³+/V²+, and Mn⁴+/Mn³+. In some cases, the increase in valence opens a sub-band that was occupied with the lower valence, with the result that the oxygen K-edge shifts to lower excitation energy. This happens for the octahedral 3d³, 3d⁴, and 3d⁵ systems, including Ni²⁺/Ni³⁺ and Fe²⁺/Fe³⁺. These two examples also contain a negative Δ ion (i.e., a system where the charge transfer energy (Δ) is smaller than zero, implying a significant amount of holes on the oxygen ions). A negative Δ is a second reason for a shift to lower energy, and the couple Cu²⁺/Cu³⁺ shows a shift to lower energy, although no new sub-band becomes available in the low-spin negative ΔCu³⁺ oxides. Finally, a change of spin-state from high-spin to low-spin can cause a positive shift as visible for Co²⁺/Co³⁺ (in cases where the Co³⁺ system is low-spin). In conclusion, we state that there is no uniform rule regarding the shift of the excitation energy with valence. A last point of warning is that the energy calibration of many oxygen K-edges is not uniform throughout the literature.

Crystal Field. As mentioned above, the crystal field in octahedral high-spin oxides follows the approximate rule that it is equal to 0.6 eV times the formal valence, implying a crystal field of 1.2 eV for 2+ oxides, 1.8 eV for 3+ oxides, and 2.4 eV for 4+ oxides. The exchange splitting for 3d oxides is approximately 0.6 eV per paired spin-couple. Taken together, these numbers imply that divalent oxides are always high-spin, tetravalent oxides are always low-spin, and trivalent oxides have both options, where Mn³⁺, Co³⁺, and Ni³⁺ are in many oxides close to a high-spin—low-spin transition point.

7.1.3. Perovskites. Perovskites have a general formula ABO₃, where A is a larger cation than B, and they are arranged in the ideal cubic structure illustrated in Figure 37a. Almost all metallic elements from the periodic table are stable in this structure. Depending on the chemical formulation, distortions from the ideal cubic structure might happen. Another related family of materials are the layered perovskites, which contain layers of perovskite-like arrangement of MO₆ metal cation units.

Figure 37. (a) Crystal structure of cubic perovskite with the ABO₃ formula. (b) Crystal structure of layered perovskite with the A₃B₂O₉ formula. A: green; B: orange, O: red.

One simple example is A₂BO₄, the structure of which is illustrated in Figure 37b. Depending on the chemical composition, substitution of various elements allows tuning of the electron or hole doping in the structure and the resulting properties. Perovskite and layered perovskite materials are intensively studied from their chemical (heterogeneous catalysis, surface chemistry) and physical properties (electric, magnetic, and optical properties). The layered structure results in enhanced electron mobility and can give rise to specific properties such as superconductivity.

Oxygen K-edge spectra have been largely used to probe the M–O chemical bonding or the metal valence and spin state, although it often requires complementary measurements such as the metal L₂,3-edges. Suntivich et al. performed a systematic analysis of the chemical bonding of LaTMO₃ systems based on the oxygen K-edge, for the transition metals Cr, Mn, Fe, Co, and Ni. The oxygen K-edge spectra of the series extended with the other 3d metals from complementary references is shown in Figure 38. The perovskite spectra show a similar general shape with a first single or double peak corresponding to the oxygen 2p states hybridized with the metal 3d states. The shape of this features is correlated to the valency and spin state of the transition metal ion. At 535 eV, one finds the oxygen 2p states hybridized with the lanthanide 5d states followed by the oxygen 2p states hybridized with the metal 4sp states.

Covalency in d⁶ Tetrahedral Oxyanions. The isoelectronic d⁶ metal cations from groups 6 and 7 (Cr, Mo, W, Mn, Tc, Re) form tetrahedral oxyanions MO₄⁻. The investigation of this series allows probing the metal—oxygen bond covalency trends. A first comparative study of the oxyanions MO₄⁻ (M = Ti, V, Cr, Mn) was done by Brydson et al. who proposed a rationalization of the observed trends in both metal L₂,3-edges and oxygen K-edge. Minasian et al. performed this comparative study, including the radioactive Tc compound, using oxygen K-edges with three different detection methods, respectively, transmission XAS in a STXM, TFY XAS, and IXS, and compared the experiments with TD-DFT calculations. In an earlier study, Bradley et al. had also compared the oxygen K-edge spectra obtained from different methods. Figure 39 shows the oxygen K-edge spectra of the series. The first two peaks in the spectra are, respectively, the transition to the empty c and t₂g states, split by approximately 2.0 eV in all six systems. The excitation energy of the first peak is the lowest in the 3d systems, followed by 4d and 5d. In addition, the Mn, Tc, and Re oxyanions have lower excitation energies than their Cr, Mo, and W analogs. There is a linear relation between the first peak in the oxygen K-edge and the first optical excitation energy, i.e., from the highest occupied state to the empty c states. Minasian et al.
demonstrated that the \( \epsilon (\sigma \text{ overlap}) \) mixing increases from group 6 to 7 and down the periods, while the \( t_2 \) symmetry \( (\pi + \sigma \text{ overlap}) \) mixing is rather constant. TD-DFT simulations of spectra are in good agreement with the experimental spectra.

### 7.2. Overview of the 3d Oxides

In the following sections, the 3d transition metal oxides are discussed by element.

#### 7.2.1. Titanium Oxides

Most titanium oxides have a valency of 4+, implying an empty 3d band. They are insulators with a band gap between the oxygen 2p band and the titanium 3d band. The binary oxide TiO\(_2\) occurs in three different crystal structures, respectively, rutile, anatase, and brookite. Its oxygen K-edge has been measured many times \(^{16,232,296,313}\). The "perovskite" structures, respectively, rutile, anatase, and brookite. Its oxygen K-edge has been measured many times \(^{16,232,296,313}\). Because the coordination in the perovskites is close to octahedral, the eg band is relatively wide.

The published oxygen K-edges of SrTiO\(_3\) show some variation in the intensity of the \( e_g \) peak. \(^{296,317,320,321}\) The oxygen K-edges of CaTiO\(_3\) \(^{316}\) and BaTiO\(_3\) \(^{318}\) are equivalent to that of SrTiO\(_3\). The best resolution oxygen K-edge of SrTiO\(_3\) is given by Palina et al. \(^{307}\) Palina et al. compared pure SrTiO\(_3\) with TiO\(_2\)-terminated SrTiO\(_3\) and thin films/interfaces with LaAlO\(_3\). They claim that for the conducting samples, a new peak is visible at 540.5 eV due to "molecular-like oxygen", related to the concentration of electronic defects.

The trivalent titanium oxides Ti\(_3\)O\(_4\) \(^{16,316}\) and LaTiO\(_3\) \(^{120}\) have a 3d\(^1\) ground state. Havercort et al. performed a detailed analysis of the ground state \(^{325}\) based on the titanium L-edge. They found that the ground state is affected by the crystal field distortion and the 3d spin-orbit coupling. Such details cannot be found from the oxygen K-edge due to the lack of final state interactions. Fujimori et al. studied the effects of doping from the oxygen K-edge spectra of a series of (La,Sr)TiO\(_3\) systems. \(^{319}\) While electron doping creates a pre-peak for the late transition metal ions such as in the cuprates, this is not the case for the Ti\(^{3+}/Ti^{4+}\) system. The first \( t_{2g} \) peak in the oxygen K-edge has approximately the same energy for LaTiO\(_3\) and SrTiO\(_3\), which is related to the large covalency of the titanium systems and the fact that the Hubbard U is smaller than the charge transfer energy.

#### 7.2.2. Vanadium Oxides

The vanadium 2p XAS edge sits at 515 eV, which implies that the oxygen K-edge of vanadium oxides is found in the tail of the vanadium L\(_{2,3}\)-edge. This complicates the analysis, as the background of the vanadium L\(_{2,3}\)-edge must be subtracted. In the 1989 paper on transition metal oxides, the spectrum of V\(_2\)O\(_5\) and VO\(_2\) has been included. \(^{16}\) A systematic study of the oxygen K-edges of vanadium oxides has been published by Abbate et al. \(^{323}\) They compared the spectra of V\(_2\)O\(_3\), VO\(_2\), and V\(_2\)O\(_5\) with the projected DOS calculated with the LSW method. Goering et al. measured V\(_2\)O\(_3\), VO\(_2\), and V\(_2\)O\(_5\) with the mixed valent oxide V\(_6\)O\(_{13}\). They measured a strong angular dependence in the oxygen K-edge of V\(_2\)O\(_3\) \(^{324}\) Hebert et al. measured the VO, V\(_2\)O\(_3\), VO\(_2\), and V\(_2\)O\(_5\) oxides with EELS and performed Wien97 calculations of the vanadium oxides. \(^{329}\) In the calculations, the core-hole was not included.

V\(_2\)O\(_3\) has a 3d\(^2\) ground state with a partly filled spin-up \( t_{2g} \) shell; it has two phase transitions at approximately 150 and 250 K. Below 150 kV\(_2\)O\(_3\) is an antiferromagnetic insulator; above 250 K an antiferromagnetic metal and above 250 K an insulator again. Park et al. performed detailed vanadium L\(_{2,3}\) XAS and linear dichroism for three different phases of V\(_2\)O\(_3\) as a function of temperature. \(^{325}\) This revealed the details of the electronic ground states. The oxygen K-edges measured by Park can be found in a paper by Huang et al. \(^{326}\) The first peak related to the transition to the \( t_{2g}^- \)up state is relatively low in intensity. \(^{299,324,326}\) much lower than the 3d\(^2\) system CrO\(_2\) (see below). The increased valence of Cr\(^{4+}\) drastically increases the overlap of the \( t_{2g}^- \)states with the oxygen states.

VO\(_2\) has a 3d\(^0\) ground state with a partly filled spin-up \( t_{2g}^- \) shell; it has a metal–insulator transition, and the differences in the electronic structure below and above the phase transition have been studied with the oxygen K-edge. \(^{327}\) In the insulating phase, the \( t_{2g}^- \) band is split into two sub-bands. Abe et al. measured the VO\(_2\) phase transition with EELS. \(^{328}\) Room temperature spectra of VO\(_2\) have also been published by van Aken et al. \(^{316}\) and Liang et al. performed detailed calculations that are discussed in the theory section. \(^{321}\)

V\(_2\)O\(_5\) has a 3d\(^0\) ground state and is a semiconductor with a band gap between the oxygen 2p band and the vanadium 3d band. \(^{329}\)\(^{329}\)\(^{329}\)\(^{329}\)\(^{329}\)\(^{329}\) V\(_2\)O\(_5\) has two different vanadium surroundings and five...
differently coordinated oxygen atoms. The spectrum is different from $\alpha$-V$_2$O$_5$ but well reproduced from DFT calculations. Maganas et al. calculated the oxygen K-edge of V$_2$O$_5$ using a cluster-based calculation. The important feature in their calculation is that they also calculated the vanadium L-edge using a restricted open shell configuration interaction with singles (ROCIS) method.

Lithium-doped vanadium oxides are potential battery materials. The oxygen K-edges of Li$_{1+x}$V$_3$O$_8$ oxides have been measured by Choi et al. The spectra revealed that the insertion of lithium affects the V=O bonds and induces a change in bonding geometry. Willinger et al. measured the oxygen K-edge of the vanadium phosphate VOPO$_4$. The V$^{4+}$ ions sit in a distorted octahedral site. The 3d-part of the spectrum looks similar to V$_2$O$_5$, and at higher energy, the oxygen K-edge is dominated by phosphor–oxygen bonds. The oxygen K-edge of vanadium bronzes has been measured with XAS and with EELS. The Na$_{0.33}$V$_2$O$_5$ spectrum is calculated with Wien2K calculations without core–hole. The Al$_2$V$_4$O$_8$ spinel system is measured with EELS and compared with Wien2K calculations. In contrast to the binary oxides, no separate 3d-related features are observed. Bismuth vanadate, BiVO$_4$, is used as an anode in photoelectrochemical water splitting. The oxygen K-edge of BiVO$_4$ is different from that of V$_2$O$_5$. Mixed oxides with other transition metal ions include the (V,Cr)O$_2$ systems. For a full overview of all systems, we refer to the XASEELS database, where all oxygen K-edge spectra are collected.

### 7.2.3. Chromium Oxides

Chromium exists in a large number of valences, from 3d$^4$ 2+ to 3d$^6$ 6+. In 1971, Fischer measured the oxygen K-edge spectra of Cr$_2$O$_3$, CrO$_3$, Na$_2$CrO$_4$, and K$_2$Cr$_2$O$_7$ using a laboratory X-ray source. We compare them below, system by system, with more recent data.

The divalent binary oxide CrO does not exist as bulk material. It has been made as a thin film, and its Cr L$_{2,3}$ XAS spectrum has been measured but not its oxygen K-edge spectrum. Cr$_2$O$_3$ has a 3d$^3$ ground state forming an antiferromagnetic ground state. For every Cr$^{3+}$ ion, the t$_{2g}$ shell is half-filled. The crystal field splitting is equivalent to the exchange splitting yielding a two-peak 3d-part of the spectrum with the first peak being a combination of spin-up e$_g$ and spin-down t$_{2g}$ states. The oxygen K-edge of Cr$_2$O$_3$ has been published in a number of papers. Figure 41 reproduces four published Cr$_2$O$_3$ spectra. Qualitatively, the spectral shapes look equivalent, and some variation is caused by the different experimental resolutions. The second peak/shoulder of the 3d
part of the spectrum shows intensity variations, and the energy calibration of the Cr$_2$O$_3$ spectrum is different in every publication, depending on the specific calibration method used.

Docherty measured a series of spinel spectra, including MgCr$_2$O$_4$ and ZnCr$_2$O$_4$, that all contain Cr$^{3+}$. The 3d part of the measured spectra looks essentially like Cr$_2$O$_3$. At higher energies, the peaks resemble the delocalized electrons of the spinel crystal structure. Remarkably, LaCr$_2$O$_4$ containing Cr$^{3+}$ and 3d$^3$ looks completely different from Cr$_2$O$_3$. From local considerations, one would expect every distorted, octahedral Cr$^{3+}$ to look similar to the two peaks observed for Cr$_2$O$_3$. What is special in MnO is that the excitation energy is high because of the high ionicity of MnO, i.e., a large charge transfer energy $\Delta$. In addition, MnO is a rock salt octahedral system that undergoes condensation reaction, as is the case for chromate ions forming dichromate ions. The binuclear [Cr$_2$O$_7$]$^{2-}$ complex has been measured by Diaz et al., and the spectrum is compared to the spectrum of the simple oxyanion in Figure 39. The shape is strongly altered with the broadening and shifting to lower energies of the two first peaks. This can be related to the strong distortion of the CrO$_4$ tetrahedra in the dichromate species and increase of three of the four Cr–O bond lengths should explain the decrease of the ligand field.

Applications of chromium include catalysts, paints, and steel compositions. Cr$_2$O$_3$ and Cr$_3$O$_4$ supported on titania are used for the selective catalytic reduction of nitric oxide by ammonia. The shapes of the oxygen K-edge of Cr$_2$O$_3$ and Cr$_3$O$_4$ films on TiO$_2$ were compared with bulk references, suggesting that new chromium oxide phases are not present in the bulk. Lopez et al. studied a series of stainless steels, comparing their chromium, iron, and nickel L$_{2,3}$-edges and oxygen K-edges. The spectra confirmed that in all cases the passive film was formed by Cr$_2$O$_3$.

Mixed oxides with other transition metal ions include the (V,Cr)$_2$O$_3$ systems, CuCr$_2$O$_4$ spinel, (Li,Ti)CrO$_7$, CuCr$_2$O$_4$ spinel systems, (Cu,Ni,Co)Cr$_2$O$_4$. For a full overview of all systems, we refer to the XASEELS database.

**Figure 41.** Oxygen K-edge of Cr$_2$O$_3$. (a) ref 339; (b) ref 302; (c) ref 340; (d) ref 341. There is variation visible in the spectral shape and especially in the energy calibration.

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to $e_g$-up, $t_{2g}$-down, and $e_g$-down. Mn$_3$O$_4$ has two octahedral Mn$^{3+}$ sites and one tetrahedral Mn$^{2+}$ site. The oxygen K-edge can in the first approximation be described as the Mn$_3$O$_4$ spectrum where the second peak is related to the tetrahedral Mn$^{3+}$ site. We note that there is some variation in the observed spectral shapes of Mn$_3$O$_4$.303,351,353,357

Mn$_2$O$_3$ has a rutile structure where Mn$^{3+}$ occupies distorted octahedral sites and has a 3$d^1$ antiferromagnetic ground state. For every Mn$^{4+}$ ion, the spin-up $t_{2g}$ states are occupied. The Mn$_4$O$_5$ spectrum has two peaks related to the 3$d$ part of the spectrum.16,300,303,351,355−358 The first peak is a combination of the spin-up $e_g$-states and the spin-down $t_{2g}$-states. The second peak is related to the spin-down $e_g$-states. The oxygen K-edge spectra of MnO, Mn$_2$O$_4$, Mn$_2$O$_3$, and MnO$_2$ have been calculated with FEFF7, yielding not very accurate simulations of the general shape.233 More accurate FEFF calculations were performed by Gilbert et al.303 They also remeasured the manganese oxides and carefully corrected for beam damage effects. The calculations essentially confirmed the exchange plus crystal field picture for the 3$d$-part of the spectrum and further explain the higher-energy spectral shape.

The tetrahedral Mn$^{3+}$ permanganate MnO$_4^−$ ion has a 3$d^0$ ground state and has been measured as KMnO$_4$.303,311 Kurata et al. have measured the 3$d^1$ equivalent BaMnO$_4$.353 However, the spectral shape is very close to that of MnO$_2$, and although the authors did not mention it, it is possible that the data correspond to a photoreduced compound. Minasian et al.311 measured the oxygen K-edge spectrum of KMnO$_4$ with three different detection methods, respectively, transmission XAS in a STXM, TFY XAS, and IXS and compared the calculations with TD-DFT calculations. They compared a series of MO$_4^−$ anions and performed a detailed analysis on the trends in their chemical bonding.

Suntivich et al. performed a detailed analysis of the chemical bonding of LaTMO$_3$ systems based on the oxygen K-edge,298 for the transition metals Cr, Mn, Fe, Co, and Ni. The LaMnO$_3$ contains the 3$d^4$ Mn$^{3+}$ cation. The oxygen K-edge spectrum has three distinct features, related mainly to, respectively, the manganese 3$d$ states, the lanthanum 5$d$ states, and the manganese 4$p$ states. The equivalent oxygen K-edge (Figure 42) of LaMnO$_3$ has also been measured by Abbate et al.359 and Lafuerra et al.,360 and the PrMnO$_3$ by Toulemonde et al.361 The (La, Sr)MnO$_3$ perovskites show a transition from Mn$^{3+}$ in LaMnO$_3$ to Mn$^{4+}$ in SrMnO$_3$.359 Changing lanthanum for Sr replaces the lanthanum 5$d$ states for the strontium 4$d$ states, while the manganese 3$d$ states change from Mn$^{3+}$ 3$d^4$ to Mn$^{4+}$ 3$d^3$, thereby increasing the intensity of the first structure in the oxygen K-edge. Lafuerra et al.360 studied the doping of the manganese sites in LaMnO$_3$ with gallium. The La(Mn,Ga)O$_3$ system shows in the first approximation a gradual decrease of the Mn$^{3+}$ signal, where both features in the 3$d$ band decrease and the manganese 4$p$ band is replaced by a gallium band. The (Pr,Ca)MnO$_3$ system studied by Toulemonde et al.361 shows a similar change as the (La,Sr)MnO$_3$ perovskites; the 3$d$ part of the PrMnO$_3$ spectrum is similar to that of LaMnO$_3$ and CaMnO$_3$ is similar to SrMnO$_3$. Galdi et al. performed a detailed oxygen K-edge study of LaMnO$_3$/SrMnO$_3$ multilayers.362 Their spectra for superlattices (SrMnO$_3$)$_n$/(LaMnO$_3$)$_m$ mainly showed the lanthanum 5$d$ states and no strontium 4$d$ states. Mixed oxides with other transition metal ions include the La(Mn,Co)O$_3$ perovskites363 and Ba$_3$Ti$_2$MnO$_9$.364

Figure 43 compares the oxygen K-edge XAS spectra of MnO with a series of (Li,Mn)$_2$O$_4$ oxides, respectively, 3$d^4$ Li$_2$MnO$_4$,

Figure 42. Oxygen K-edge of LaMnO$_3$.298 and SrMnO$_3$.359 The peaks are caused by the Mn 3$d$ states, the La and Sr 5$d$/4$d$ states, and the Mn 4$p$ states, as indicated.

Figure 43. Oxygen K-edge of MnO, LiMnO$_2$, Li$_2$MnO$_4$, Li$_3$MnO$_4$, and Li$_4$MnO$_4$. The 3$d^4$ oxide MnO contains impurities from Mn$^{3+}$ visible in the intensity before 529 eV. The 3$d^4$ oxide LiMnO$_2$ has a broad 3$d$ part due to a combination of the octahedral crystal field, exchange, and the Jahn–Teller distortion. The 3$d^4$ oxide Li$_2$MnO$_4$ has a first peak related to a combination of spin-up $e_g$ and spin-down $t_{2g}$ states. LiMnO$_2$ is a mixed valence Mn$^{3+}$/Mn$^{4+}$ oxide, and in the first approximation its 3$d$ part is a combination of the spectra of Mn$^{3+}$ and Mn$^{4+}$ sites.

7.2.5. Iron Oxides. Iron oxides are among the most abundant materials on earth, in particular, Fe$_2$O$_3$ and Fe$_3$O$_4$. Iron oxides have valences 3$d^6$ 2$^+$ and 3$d^5$ 3+. Fe$^{4+}$ and higher valences are uncommon for iron oxide materials. FeO has a 3$d^6$ ground state; it is known to be intrinsically unstable with regard to its stoichiometry. The oxygen K-edge of FeO has been often measured since the first measurement by Grunes in 1982.57,232,301,315,516 All spectra show a first peak related to mixed-valent 3$d^6$/3$d^5$ Li$_2$MnO$_4$, and 3$d^4$ LiMnO$_2$.352 The Li$_2$MnO$_4$ spectrum is similar to that of MnO$_2$, and the spectra of the two 3$d$ systems Mn$_2$O$_4$ and LiMnO$_2$ are also similar. The Li$_2$MnO$_4$ oxide was also measured by Risch et al.357 Mixed oxides with other transition metal ions include the Li(Mn,Ni)$_2$O$_4$ battery systems,358 La(Mn,Co)$_2$O$_3$ perovskites363 and Ba$_3$Ti$_2$MnO$_9$.364 For a full overview of all systems, we refer to the XAS database.337
the iron 3d states, that depending on the experimental resolution, have some structure.

Fe2O3 has a 3d6 ground state and appears in several crystal structures, the most common being the corundum \(\alpha\)-Fe2O3 structure.\textsuperscript{16,57,301,302,316,321,351,366} Fe3O4 is a half-filled anti-ferromagnet, and the first two peaks directly relate to, respectively, the spin-down \(t_{2g}\) and \(e_g\) states. The oxygen K-edge of \(\gamma\)-Fe2O3 looks similar, with some differences at higher energy due to the different crystal structure.\textsuperscript{57}

The \((\text{La},\text{Sr})\text{FeO}_3\) perovskites show a transition from \(\text{Fe}^{3+}\) in \(\text{LaFeO}_3\) to \(\text{Fe}^{4+}\) in \(\text{SrFeO}_3\).\textsuperscript{359} The oxygen K-edge of \(\text{LaFeO}_3\) is compared to that of \(\text{SrFeO}_3\) in Figure 44. We observed the \(t_{2g}\) and \(e_g\) peaks, followed by the lanthanum 5d states, the La and Sr 5d/4d states, and the Fe 4sp states, as indicated.

![Figure 44. Oxygen K-edge of LaFeO3 and SrFeO3 perovskite. The peaks are, respectively, caused by the Fe 3d states, the La and Sr 5d/4d states, and the Fe 4sp states.](Image)

7.2.6. Cobalt Oxides. CoO has a 3d\(^{7}\) ground state that is always high-spin in oxides. Octahedral Co\(^{3+}\) has a \(t_{1g}\) ground state that is split by the 3d spin–orbit coupling. The oxygen K-edge of CoO has two peaks split by the crystal field, respectively, related to the empty \(t_{2g}\) state and the empty \(e_g\) state.\textsuperscript{302,303,346,367,373–375} At higher energy is found the typical double structure of the rock salt crystal.

Co\(^{3+}\) has a 3d\(^{8}\) ground state; in oxides, the crystal field is similar in magnitude to the exchange splitting, implying that Co\(^{3+}\) occurs as \((t_{2g})^2\) low-spin and as \((t_{2g})^2\) high-spin oxides. The oxygen K-edge of CoO\(_2\) has a sharp single peak at the edge indicating a low-spin Co\(^{3+}\) ground state.\textsuperscript{304} CoO\(_2\) is a spinel with tetrahedral Co\(^{2+}\) and octahedral Co\(^{3+}\) sites.\textsuperscript{373} Because the Co\(^{3+}\) site is more covariant, it has more oxygen character, and the oxygen K-edge of CoO\(_2\) is dominated by the low-spin Co\(^{3+}\) features. The peaks of the tetrahedral Co\(^{3+}\) site are superimposed.

The oxygen K-edge of lithium-doped CoO and its end member LiCoO\(_2\) have been often studied in relation to charge doping in CoO and in particular related to battery research.\textsuperscript{373,374,376–378} LiCoO\(_2\) is a low-spin system with a similar spectral shape to that of CoO\(_2\), i.e., a sharp peak related to the empty \(e_g\) states, followed by the cobalt 4sp band. Doping CoO with lithium gives oxygen K-edges that can be approximated as a mixture of CoO and LiCoO\(_2\).\textsuperscript{373}

In the perovskite LaCoO\(_3\), the 3d\(^{4}\) Co\(^{3+}\) undergoes a spin transition from low-spin at low temperature to mainly high-spin above 500 K. The \(t_{2g}\) low-spin system EuCoO\(_3\) has the 3d peak related to the empty \(e_g\) states at 259.5 eV with a tail to higher energy.\textsuperscript{379} The \(t_{2g}\) high-spin system Sr\(_2\)CoO\(_3\)Cl contains Co\(_3\) pyramids and has its first \(t_{2g}\) peak at 258.5 eV and a second smaller \(e_g\) peak at 530.5 eV.\textsuperscript{380} The phase transition in LaCoO\(_3\) was studied by Abbate et al.\textsuperscript{380} The transition from the low-spin one-peak spectrum to a broader spectrum is observed, but there is little peak shift and the spectral change is not very clear. Similar LaCoO\(_3\) spectra have been measured\textsuperscript{298,375} but as these data are obtained at room temperature, they are a mixture between low-spin and high-spin. Simbok and co-workers studied the coexistence of low-spin and high-spin ions in LaCoO\(_3\) systems of a series of catalytically active systems. They correlate the oxygen K-edge derived electronic parameters with reaction rates.

The low-spin \((t_{2g})^2\) Co\(^{4+}\) system SrCoO\(_2\) shifts to lower energy with respect to low-spin Co\(^{3+}\) oxides, and a small shoulder is visible at low energy related to the empty \(t_{2g}\) state.\textsuperscript{375} The \((\text{La},\text{Sr})(\text{Li},\text{Co})\text{O}_3\) system studied by Hu et al.\textsuperscript{369} and the spectral shape is a combination of the low-spin Co\(^{3+}\) and Co\(^{4+}\) peaks.\textsuperscript{367,371} Harvey et al. also measured the Co\(^{4+}\) oxides SrCoO\(_2\) and BaCoO\(_2\)\textsuperscript{369} but their observed spectral shapes look different, with significant Co\(^{3+}\) contribution, in addition to large carbonate peaks that make the spectrum complex to analyze above 532 eV. The \((\text{La},\text{Sr})\text{CoO}_3\) system was studied by Moodenaugh et al. and by Hu et al.\textsuperscript{367,382} They concluded that already at 10% strontium doping the Co\(^{4+}\) switches to high-spin, modifying the whole spectral shape. The (partial) change in spin state of Co\(^{3+}\) and Co\(^{4+}\) sites makes the comparison of the oxygen K-edge spectra of Co\(^{3+}\) and Co\(^{4+}\) systems very complex to analyze on their own, without other information such as the cobalt L\(_{2,3}\)-edge.

Mixed cobalt oxides with other transition metal ions include the \((\text{La},\text{Sr})(\text{Co},\text{Fe})\text{O}_3\)\textsuperscript{371} \((\text{Ba},\text{Sr})(\text{Co},\text{Fe})\text{O}_3\)\textsuperscript{369} and \((\text{Co},\text{Ni})\text{O}_2\)\textsuperscript{364}. For a full overview of all systems we refer to the XASEELS database.\textsuperscript{337}

7.2.7. Nickel Oxides. \(\text{NiO}\) has a 3d\(^{8}\) ground state with two \(e_g\) holes. The oxygen K-edge of \(\text{NiO}\) is one of the most published oxygen K-edge spectra, whereas in this review, we limit the citations to the papers that focus on the spectral shape.\textsuperscript{16,232,302,313,316,354,375,373,375,383–388} A full overview of all published \(\text{NiO}\) spectra is given in the XASEELS database.\textsuperscript{337} The \(\text{NiO}\) spectrum has been measured over a long energy range by Zschech et al.,\textsuperscript{385} where they focused on the self-absorption effects in \(\text{NiO}\) detection. We refer to the experimental section for further discussion of \(\text{NiO}\) detection.

The 3d\(^{2}\) \(\text{Ni}^{3+}\) system \(\text{Ni}_2\text{Li}_3\text{Li}_3\text{Ni}_3\text{O}_4\) is considered to be a low-spin system based on its \(\text{Ni}^{3+}\) ground state. The oxygen K-edge shows a shift of 2.6 eV with respect to \(\text{NiO}\) with a sharp peak at the edge related to the three empty \(e_g\) states.\textsuperscript{371,373,375} The Ni L\(_{2,3}\)-edge is a complex to analyze above 532 eV. The \((\text{La},\text{Sr})\text{CoO}_3\) system was studied by Moodenaugh et al. and by Hu et al.\textsuperscript{367,382} They concluded that already at 10% strontium doping the Co\(^{4+}\) switches to high-spin, modifying the whole spectral shape. The (partial) change in spin state of Co\(^{3+}\) and Co\(^{4+}\) sites makes the comparison of the oxygen K-edge spectra of Co\(^{3+}\) and Co\(^{4+}\) systems very complex to analyze on their own, without other information such as the cobalt L\(_{2,3}\)-edge.
EuNiO$_3$. The layered perovskite (Nd,Sr)$_2$NiO$_4$ shows a different oxygen K-edge compared with the Nd$_3$Li$_5$Ni$_{36}$O$_{44}$ system, likely due to a partial high-spin ground state. Similar oxygen K-edge spectra are measured for La$_2$NiO$_4$ containing 3d$^4$ Ni$^{2+}$. Hu et al. measured the polarization dependent oxygen K-edge of the Ni$^{2+}$ oxide Y$_2$BaNiO$_4$. Upon doping, the yttrium sites with calcium and Ni$^{3+}$ state are found.

### 7.2.8. Copper Oxides

CuO has a 3d$^9$ ground state with one e$_g$ hole. CuO has a rather complex crystal structure with low symmetry. The oxygen K-edge of CuO has a sharp peak at the edge related to the empty e$_g$ state. At higher energy, the spectrum appears quite different from all other rock salt divalent transition metal oxides, related to its monoclinic crystal structure. The CuO spectral shape has been calculated by Jiang et al.

The oxygen K-edge of Cu$_2$O was measured by Stohr et al. in 1980. They measured the spectrum over an energy range of 400 eV. Jiang et al. calculated the oxygen K-edge of Cu$_2$O in good agreement with experiments. Hu et al. measured the oxygen K-edge RIXS of Cu$_2$O. The oxygen K-edge spectrum of the formally Cu$^{3+}$ oxide La$_2$CuO$_4$ has been studied by Hu et al.

There are many papers on the cuprate systems, and full coverage falls outside the scope of this review. We focus on the main papers that show the experimental data. The oxygen K-edges measured with XAS and EELS up to 1994 have been reviewed by Pink et al. The study on the effect of Sr-doping in the layered perovskite La$_2$CuO$_4$ has been studied with oxygen K-edges using EELS and XAS. La$_2$CuO$_4$ has a Cu$^{2+}$ related peak at 530.5 eV. Upon doping with Sr, part of the copper is changed to Cu$^{3+}$ with a peak at 528.5 eV, where the 528.5 eV peak shifts slightly up. Recent contributions on the oxygen K-edge of cuprates include RIXS studies and the measurement of layer specific spectra using a STEM EELS study.

### 7.3. General Considerations of the 4d and 5d Oxides

The crystal field splitting in 4d oxides is larger than for 3d oxides, typically around 3.5 eV for octahedral sites. This implies that 4d oxides always are in a low-spin ground state. Figure 45 shows a selection of oxygen K-edge spectra for 4d oxides. Most oxides are not magnetic, implying that the assignment of the peaks is essentially the same for all 4d oxides in octahedral symmetry: respectively (a) the 4d $t_{2g}$ states, (b) the 4d $e_g$ states, and (c) the empty sp bands of the metal ion. Mixed oxides have the empty states of the other cation superimposed.

### 7.4. Overview of the 4d and 5d Oxides

#### 7.4.1. 4d Transition Metal Oxides

**Yttrium Oxides.** Yttria, Y$_2$O$_3$, is a wide-gap semiconductor with a band gap of 5.6 eV. It has a cubic crystal structure where Y cations are sixfold coordinated in distorted octahedrons. The oxygen K-edge of Y$_2$O$_3$ (Figure 45) shows two main features corresponding to the crystal field split 4d states with the $t_{2g}$ peak at 533 eV and the $e_g$ peak at 538 eV, implying a crystal field of 5.0 eV. We note that there is an energy spread of 3.0 eV in the energy calibration of the first peak in the publications.

**Zirconium Oxides.** The most common oxide of zirconium is ZrO$_2$, which is also known as zirconia. ZrO$_2$ occurs in three phases, cubic, tetragonal, and monoclinic, and one can use an impurity element to stabilize one of those phases. The oxygen K-edge of ZrO$_2$ (Figure 45) shows subtle differences between the different polymorphs. It mainly contains the 4d $t_{2g}$ peak at 532.4 eV and the 4d $e_g$ peak at 535.5 eV, implying a crystal field of 3.1 eV. Several investigations of the oxygen K-edges in ZrO$_2$ and yttrium stabilized zirconia (YSZ) samples with varying compositions have been reported demonstrating that the spectra are very sensitive to local crystalline structure and dopant content. For example, ytrria stabilized zirconia ZrO$_2$[Y] is an important ceramic material used in dental crowns and nonmetal knife blades. Recently, a number of other dopants were studied in relation to the photocatalytic properties of ZrO$_2$, including ZrO$_2$[Ce]$^{4+}$ and ZrO$_2$[Cu]$^{4+}$.

**Niobium Oxides.** Niobium has three oxides, respectively, divalent 4d$^1$ in the cubic NbO$_2$, tetravalent 4d$^1$ in the tetragonal NbO$_2$, and pentavalent 4d$^3$ Nb$_2$O$_5$. Nb$_2$O$_5$ and also Nb$_2$O$_7$(OH) nanostructures are promising materials in the field of photoelectrodes in dye-sensitized solar cells and photoelectrochemical cells. Olszta et al. compared the oxygen K-edges of the three oxides. The crystal field splitting of the first two peaks corresponding to the 4d states is further reduced compared to Y and Zr oxides. The first peak of the oxygen K-edges shifts from 533.5 eV (NbO$_2$) to 534.3 eV (Nb$_2$O$_5$) and 535.1 eV (Nb$_2$O$_7$), as determined by our measurements on the published spectra. The shifts are inverted at the Nb M$_3$ edge, i.e., respectively, 365.9 eV (NbO$_2$) to 364.6 eV (Nb$_2$O$_5$) and 362.8 eV (Nb$_2$O$_7$). Note that the relative shifts are larger for the Nb M$_3$ edge than for the oxygen K-edge, related to the larger core−hole effect for the Nb 3p core−holes. Soriano et al. compared the Nb$_2$O$_5$ spectrum to...
the other 3d, 4d, and 5d oxides.\textsuperscript{402} The 5d oxide KNbO₃ has its first peak at 531.1 eV,\textsuperscript{419} at lower energy than NbO₂ (531.6 eV\textsuperscript{402} respectively, 533.3 eV\textsuperscript{404}). The crystal field is also 0.5 eV larger for KNbO₃.

**Molybdenum Oxides.** Molybdenum has two binary oxides, 4d\textsuperscript{0} MoO₃ and 4d\textsuperscript{2} MoO₂. MoO₃ has a distorted rutile structure. The oxygen K-edge of MoO₂, shown in Figure 45, has peaks related to the empty 4d states, but there is significant variation in the published spectra.\textsuperscript{406,420–422} Thakur et al. compared TEY and TFY spectra with high resolution,\textsuperscript{421} and they observed three peaks within the 4d band that they assigned to a distortion of the octahedral site. The three features were also observed by Khyzhun et al.\textsuperscript{426} Wang et al. studied the reduction of MoO₃ to MoO₂ in the electron beam during an EELS experiment.\textsuperscript{406} Possibly all MoO₂ spectra contain some admixture of MoO₃, but as the peaks essentially overlap, it is difficult to determine such mixing quantitatively.

The 4d\textsuperscript{4} system MoO₄ is used as an oxidation catalyst. Its oxygen K-edge spectrum is compared to MoO₂ in Figure 45, Schlögl et al. measured the oxygen K-edge of a single crystal MoO₃, which they compared with a partly reduced sample MoO₂.\textsuperscript{423} The other published MoO₃ spectra are all similar to this reduced spectrum.\textsuperscript{405,406,422,424} Wu et al. measured their MoO₃ sample both with electron yield and with ion yield. In ion yield, one detects the O⁺ ions that escape from the sample upon X-ray irradiation. Ion yield is an extremely surface sensitive technique and measures only oxygen atoms in the surface layer of the system.\textsuperscript{405} Several papers discuss oxygen K-edge spectra of mixed oxides, for example, the (Mo,Fe) mixed oxides in relation to magnetism\textsuperscript{425} and the (Mo,V) mixed oxides in relation to catalysis.\textsuperscript{426}

**Technetium Oxides.** Minasian et al. studied a series of d\textsuperscript{0} tetrahedral oxyanions of 3d, 4d, and 5d metal ions, including the pertechnetate TcO₄⁻ anion.\textsuperscript{311} They simulated all oxygen K-edge spectra with TD-DFT and obtained good agreement with the experimental spectra. The first two peaks in the spectra are, respectively, the transition to the empty e and t₂g states, split by approximately 2.0 eV in all six systems. The excitation energy of the first peak is lowest in the 3d systems, followed by 4d and 5d. In addition, the Mn, Tc, and Re oxyanions have lower excitation energies than their Cr, Mo, and W analogs. There is a linear resemblance to the other rutile oxides such as TiO₂ and IrO₂.

**Ruthenium Oxides.** The 4d\textsuperscript{4} system RuO₂ has a rutile structure. Its oxygen K-edge, Figure 45, has a sharp peak related to the t₂g states followed by a broader peak related to the e_g states.\textsuperscript{407,425,426} The oxygen K-edge of RuO₂ bears much resemblance to the other rutile oxides such as TiO₂ and IrO₂ (see 5d element section). Tsai et al. studied RuO₂ nanorods for field emission applications.\textsuperscript{407}

Ruthenates are systems that contain the RuO₆ octahedrons combined with other cations, mainly Sr and Ca. The most studied material is Sr₂RuO₄ and is a layered perovskite with a structure similar to that of (La,Sr)₂CuO₄, which also shares the superconducting properties with the copper analogs, though with critical temperature of less than 1 K. In 1996, Schmidt et al. measured the angular dependence of Sr₂RuO₄ single crystals.\textsuperscript{427}

Figure 46 shows the angular dependence of the oxygen K-edge spectrum of Sr₂RuO₄. The first two peaks at 528.5 and 529.5 are assigned to empty 4d t₂g states, where the first peak is related to the apical oxygen and the second to the in-plane oxygen. The following two peaks at 531 and 533.5 eV are related to 4d e_g states and have higher intensity in the out-of-plane direction.

This assignment is essentially reproduced in later experiments on Sr₂RuO₄ and Ca₃RuO₄.\textsuperscript{425,428–431} Fatuzzi et al. performed an elegant experiment to distinguish the three 2p orbitals x, y, and z, from a combination of angular and polarization dependence. They performed a RIXS experiment on Sr₂RuO₄ and Ca₃RuO₄ and analyzed the low-energy excitations as a combination of the crystal field splitting of the t₂g states and the effect of the 4d spin–orbit coupling.\textsuperscript{430} One can make more complex layered oxides from combinations of Sr and Ru. By keeping the valence of Ru at +4, the series includes several combinations of SrO with RuO₂, i.e., (SrO)₃(RuO₂)ₙ, including Sr₄RuO₁₀, Sr₃Ru₂O₇, and Sr₂Ru₃O₁₀, where a range of magnetic and superconducting properties are found in these systems. The oxygen K-edges of the Sr₂RuO₄, Sr₃Ru₂O₆, and Sr₄Ru₃O₁₀ ruthenate have been measured by Malvestuto et al.\textsuperscript{429} In addition, one can change the valence of Ru and make the Ru⁴⁺ system Sr₃RuO₄ for which Hu et al. measured the oxygen K-edge.\textsuperscript{425} Chikamatsu et al. compared the Sr₂RuO₄ system with the oxyfluoride system Sr₂RuO₃F₂. The oxygen K-edge spectra, in relation to other core and valence band spectroscopies, indicate that the Ru⁴⁺ in Sr₂RuO₃F₂ is in a high-spin ground state, in contrast to the low-spin Sr₂RuO₃ systems.\textsuperscript{431}

**Palladium Oxides.** Palladium is an important catalyst, for which the reactivity is enhanced by fabricating nanosized metallic particles. The oxidation of the surface of the nanoparticles motivates their study with microscopy and spectroscopy. Sun et al. studied a Pd/Al₂O₃ system.\textsuperscript{432} They measured the oxygen K-edge of the Pd nanoparticles with EELS and observed an oxygen K-edge that is different from the reference PdO. To interpret the experimental data obtained on nanoparticles, the oxygen K-edge of PdO was calculated for three crystal structures by Mogi et al.\textsuperscript{433} They used Wien2K calculations and reproduced the main features of the experimental spectrum.

Figure 46. Oxygen K-edge of Sr₂RuO₄ layered perovskite\textsuperscript{427} with the incident electrical field parallel to the layer ($e_{ab}$) and perpendicular to the layer ($e_g$). The contributions from the apical and in-plane oxygens are highlighted.
Silver Oxides. Metallic silver is importantly used for its surface reactivity, and several studies investigated the absorbed oxygen reactivity on the metal surface, in particular, for ethylene epoxidation. The oxygen K-edge of Ag₂O shows a first intense peak corresponding to the O 2p states hybridized with the Ag 4d states followed by the O 2p/Ag 5sp bands. Comparison with absorbed oxygen atoms reveal that the nucleophilic species bind differently from the electrophilic species at the silver surface. Ruzankin et al. studied the ethylene epoxidation on the silver surface. They observed several features in the oxygen K-edge that were simulated with an Ag₂O molecule. Another oxygen K-edge spectrum has been published for silver foil that was treated with C₂H₄ and O₂ at 470 K.⁴³⁴

7.4.2. 5d Transition Metal Oxides. Hafnium Oxides. Hafnium oxide HfO₂ is an important material in the semiconductor industry due to its high dielectric constant k. These so-called high k materials are used to replace a silicon dioxide layer of a device, which is important for further miniaturization of microelectronic components. Soriano et al. compared HfO₂ with its 3d and 4d analogs TiO₂ and ZrO₂. Unlike TiO₂, HfO₂ presents the same crystal structures and polymorphs as zirconia. The oxygen K-edge spectrum of HfO₂ is shown in Figure 47 where it is compared to TiO₂ rutile and HfO₂. The oxygen K-edge of Ag₂O shows a first peak as WO₃ related to the t₂g states overlaps with the structure related to the empty s and p empty states. Oxygen K-edge spectra have been published for films and nanorods, and γ-WO₃ is doped with Na to form sodium–tungsten bronze, where the sodium atoms occupy a fraction of the 12-coordinate interstitial A-sites in the WO₆ octahedra framework. The oxygen K-edge of Na₂WO₄ shows a similar first peak as WO₃ related to the t₂g states. The region between 535 and 545 eV shows a different spectral shape, which is related to the differences in crystal structure despite the same octahedral environment for W ions. The Na₂WO₄ system has a quite different structure from the bronzes. This system contains tetrahedral complexes of the tungsten oxyanion WO₄⁻. Its oxygen K-edge has two sharp peaks related to, respectively, the t₂ and the e empty states.²⁴²

Figure 47. Oxygen K-edge of HfO₂ compared to its 3d and 4d analogs TiO₂ (rutile) and ZrO₂ (rutile) and to IrO₂ (rutile).

Tantalum Oxides. The tantalum oxides Ta₂O₅ are important materials for memory resistors in memory devices. Kumar et al. studied the oxygen K-edge spectrum of Ta₂O₅ in such devices with a 30 nm resolved STXM experiment. The oxygen K-edge of the tantalum oxide Ta₂O₅ has two peaks at the edge, related to, respectively, the Ta 5d t₂g and e₃g states.⁴⁰²,⁴⁴³,⁴⁴⁴

Figure 48. Oxygen K-edge spectra of monoclinic WO₃ and cubic ReO₃ compared to MoO₃ (orthorhombic) with the assignment of the spectral features.⁴²⁴

Monoclinic WO₃. All spectra have a sharp first peak related to the empty t₂g states. The second structure related to the empty e₃g states overlaps with the structure related to the tungsten s and p empty states.¹¹⁹,⁴²⁴,⁴⁴⁵ Oxygen K-edge spectra have been published for WO₃ and WO₁₋ₓΓₓ. The oxygen K-edge of the series of oxyanion is similar to that of the other tetrahedral oxyanions. The Re⁷⁺ system Na₃ReO₄ contains Re in tetrahedral surroundings, and its Na₂WO₄ system has a quite different structure from the bronzes. This system contains tetrahedral complexes of the tungsten oxyanion WO₄⁻. Its oxygen K-edge has two sharp peaks related to, respectively, the t₂ and the e empty states.²⁴²

Rhenium Oxides. Rhenium oxide ReO₃ has a similar structure to that of perovskites and contains Re⁶⁺ 5d⁴ ions. Purans et al. measured its oxygen K-edge spectrum and investigated the effect of the distortion of the ReO₃ octahedron on the spectrum. The spectrum is shown in Figure 48 together with monoclinic WO₃. The ReO₃ spectrum bears much resemblance to the WO₃ and MoO₃ oxygen K-edge. The Re⁷⁺ system NaReO₄ contains Re in tetrahedral surroundings, and its oxygen K-edge is similar to that of the other tetrahedral oxynian complexes.²⁴¹ The oxygen K-edge of the series of oxynian is discussed and compared in section 7.2 on 3d metal oxides.

Iridium Oxides. The iridium oxide IrO₂ contains Ir⁴⁺ 5d⁵ ions and has a rutile structure. The oxygen K-edge spectrum is shown in Figure 47 where it is compared to TiO₂ rutile and HfO₂. The absorption threshold shifts to higher energies in the series in agreement with the larger band gaps for the oxides of the heavier elements. Since 2003, there have been a series of publications that studied the interface of HfO₂ with Si.⁴³⁷–⁴⁴⁴ In addition to HfO₂, the oxygen K-edge has been measured to be of amorphous HfSiO₄ and HfSiON.⁴⁴²
shape is very similar to the spectrum of TiO₂, revealing the effect of the rutile crystal structure. It contains a small first peak related to the single hole in the t₂g state followed by a large peak related to the e_g states. Kim et al. measured an epitaxial IrO₂ film on TiO₂. They observe that the sharp t₂g peak has a very strong angular dependence with almost all intensity being related to the σ direction. This observation implies that this empty t₂g state lacks involvement of an Ir d orbital, which in turn implies that despite the importance of the 5d spin–orbit coupling in IrO₂, the J_{eff} = 1/2 state (as observed in the iridates, see below) does not form in IrO₂. The IrO₂ oxide-based systems are often studied in relation to the oxygen evolution reaction. Pfeifer et al. showed the difference between the crystalline rutile IrO₂ and amorphous IrO₂. In amorphous IrO₂, the sharp t₂g peak at 529 eV is broadened into a double structure with a new feature at 528 eV. These two features are in two later papers assigned to, respectively, O²⁻ and O⁻ character.

Within the context on transition metal ions, a more usual assignment would be to assign the new peak at lower energy to a higher valence of the iridium, similar to the assignments in the iridates, and others. When Ir in IrO₂ is partly oxidized to Ir⁵⁺, the peaks in the oxygen K-edge will shift to lower energy. The ground state of high valent ions will have strong covalence, and part of the hole character will be on oxygen, especially for so-called negative charge transfer ions such as Cu³⁺ and Fe⁴⁺, but to relate this to O⁻ in our view not the best model. In conclusion, we follow a different assignment for the operando oxygen K-edge spectra of IrO₂ systems and assign the first peak to the creation of Ir⁵⁺ ions.

The iridates are an important group of strongly correlated iridium oxides with electron correlation properties that are governed by the combination of a large 5d spin–orbit coupling and the on-site electron correlation U. The oxygen K-edges of the SrIrO₃, Sr₂IrO₄, and Sr₃Ir₂O₇ have been measured on single crystal systems. Figure 49 compares the angular dependence of the t₂g peak in SrIrO₃ versus Sr₂IrO₄ and indicates the difference between the 3D system and the layered system: similarly to the ruthenium layered perovskites, the oxygens are all equivalent in the 3D perovskite; the out-of-plane oxygens pointing at the layer can be distinguished from the in-plane oxygens.

**Platinum Oxides.** As another noble metal, platinum is largely used as a metal for its surface reactivity. Oxygen K-edge spectroscopy has been used to investigate oxygen interaction with platinum metal surface or interface or the oxides formed. Platinum occurs in different oxide forms: tetragonal and cubic PtO, Pt₃O₄, α-PtO₂, and β-PtO₂, α-PtO₂ has been measured by Srot et al. in the context of Pt/yttrium-stabilized zirconia for oxygen electrode reaction. They compare the spectrum with a number of calculations for the other oxides. Guinel et al. studied Pt nanoparticles on carbon nanotubes and they concluded that platinum occurs as PtO₂ oxide nanoparticles.

### 7.5. General Considerations of the 4f Oxides

The oxygen K-edge spectra of lanthanide (Ln) and actinide (An) oxides are characterized by numerous features corresponding to the oxygen 2p orbitals hybridized with the f and d orbitals. Understanding these spectral features aims to determine the 4f⁻⁵ and 5d⁻⁶ orbital relative positions, electronic correlations, and their role in metal–oxygen bonding in relation with the chemical and physical properties. Determination of the nature and changes in the band gap of such materials has also been a motivation. Most of the lanthanides are nonradioactive and can be handled easily. Hence, they have often been considered as models for transuranic elements, while experimental studies of actinides are severely limited due to the toxic and radioactive nature of the samples. However, lanthanides and actinides show profound differences in their physicochemical properties due to the nature of differences between the localization of the 4f and 5f orbitals. The differences and similarities in the electronic structures and bonding properties of these f-elements as revealed by oxygen K-edge XAS are reviewed.

The systems with an open 4f shell are known under the names rare earths and lanthanides. Lanthanide atoms have 3 electrons in the 5d, 6s, and 6p mixed states and their configuration can be written as 4f⁰(5d⁶s⁶p)⁵. The 4f orbitals of the lanthanides are localized, while the 5d, 6s, and 6p orbitals form bonds with oxygen. This results in their main trivalent oxidation state, where the delocalized electrons fill the oxygen 2p valence band. The localized 4f-states are positioned in between the occupied oxygen 2p valence and the empty mixed 5d, 6s, and 6p bands.

Altman et al. studied the oxygen K-edge of the complete series of lanthanide sesquioxides Ln₂O₃ (Ln = La to Lu) with STXM and attempted to rationalize the observed spectroscopic variations in relation with the electronic structure of the Ln³⁺ ions. Two different structures were studied: the hexagonal phases where the Ln³⁺ ion is sevenfold coordinated and the...
cubic phase where the Ln$^{3+}$ ion is sixfold coordinated. The oxygen K-edges of six compounds of this series are shown in Figure 50. Starting from the assignment of the transitions observed for Lu$^{3+}$ in which the 4f is full, the authors assigned the transitions between 532 and 540 eV to oxygen 2p mixed with lanthanide 5d and 6p states. The 5d states are split by the crystal field and the authors distinguished the 5dσ contribution around 532 eV and the 5dτ contribution around 538 eV. The authors analyzed the variation of the intensity and position of the 5d and 6p states to deduce the contribution of 4f states in the assignment of the oxygen K-edge XAS peaks in the other lanthanide compounds using a configuration interaction approach. They discussed the observed abrupt changes in the spectra in terms of orbital hybridization between oxygen 2p and lanthanide 5d in relation to the chemical reactivity of these oxides.

A following study of the Ce, Pr, and Tb dioxides (cubic fluorite structure, eightfold coordinated Ln$^{4+}$ ions) by Minasian et al. revealed drastic changes in the oxygen K-edge spectra (Figure 51) as compared to the trivalent oxides with the corresponding schematic DOS adapted from ref 457.

![Figure 51](image_url)

**Figure 51.** (a) Oxygen K-edge spectra of the cubic CeO$_2$, PrO$_2$, and TbO$_2$ with the corresponding band assignment in (b) cubic fluorite structure (oxygen atoms in red and lanthanide atoms in yellow) and (c) the corresponding schematic DOS adapted from ref 457.

The contribution of an intense peak around 530 eV. This feature is assigned to transitions to the oxygen 2p states mixed with the 4f states in light of DOS calculated with VASP. It is followed by the crystal field split 5d states. Minasian et al. showed that the orbital mixing between lanthanide 4f and oxygen 2p increases in the order TbO$_2$ to CeO$_2$ to PrO$_2$.

Below, the oxygen K-edge studies of lanthanum and cerium oxides are specifically reviewed.

**7.5.1. Lanthanum Oxides.** La$_2$O$_3$ forms cubic and hexagonal crystal structures, where the hexagonal La$_2$O$_3$ is often studied with respect to its high-k dielectric constant. Calmels et al. measured the oxygen K-edge of cubic and hexagonal La$_2$O$_3$. They compared the experiments with both FEFF and Wien2K calculations and reached good agreement. Figure 52 shows the oxygen K-edge spectra of both polymorphs. Cubic La$_2$O$_3$ (sixfold coordinated La) shows two distinct peaks at 532.4 and 536.8 eV, that relate to the oxygen character of the La $e_g$ and $t_{2g}$ symmetry 5d states, but also containing 6s and 6p character. At higher energies, no clear peaks are visible. Hexagonal La$_2$O$_3$ shows a similar spectrum characterized by two features although broader and less distinct than in the cubic case. In both cases, the contribution of the hybridized 4f states was ruled out by the authors. Lu$_2$O$_3$ 4f oxides are hygroscopic and tend to form hydroxides in air. The oxygen K-edge of La(OH)$_3$ was measured and calculated by Calmels et al.

Lanthanum is present in a series of perovskites, including LaMnO$_3$ and LaCoO$_3$, and the layered perovskites including La$_4$Cu$_3$O$_8$. These 3d transition metal oxide systems have been discussed in the 3d section, as the leading peaks at the edge are the partly empty 3d states of the transition metal ion. The lanthanum 5d band is always visible at in the 530−540 eV energy range. Jeong et al. studied the decomposition of La$_2$CuO$_4$ into La$_2$O$_3$ and other oxides with TEM. By comparison to the reference spectra, their spectra showed that the decomposition product was hexagonal La$_2$O$_3$.

**7.5.2. Cerium Oxides.** Ce$_2$O$_3$ is the cerium analog of La$_2$O$_3$ and it has a 4f$^3$ configuration. The oxygen K-edge of Ce$_2$O$_3$ is shown in Figure 50 and is similar to that of La$_2$O$_3$. It is dominated by the 5d, 6s, and 6p states. Xu et al. studied mixtures of CeO$_2$ and Ce$_2$O$_3$ and the occurrence of a small peak before the Ce$_2$O$_3$ peaks was assigned to CeO$_2$ impurities.

In CeO$_2$, Ce$^{4+}$ has 4f$^3$ electronic configuration. The oxygen K-edge of cubic CeO$_2$ has been often measured both with XAS and with EELS. The spectrum is shown in Figure 51. The presence of a strong 4f-related peak highlights the non-negligible hybridization between oxygen 2p and Ce 4f orbitals. The crystal field in cerium dioxide splits the 5d states, but the splitting of the 4f states is not observed in agreement with the small overlap between the f orbitals and the ligands in lanthanides. The crystal field splitting of the 5d states is 4.3 eV for CeO$_2$. Mullins et al. reported spectra for cerium oxides thin films used as an important component in automotive exhaust catalysts. Wang et al. studied the mixture of cerium and terbium oxides. Garvie and Buseck showed that CeO$_2$ changes to Ce$_2$O$_3$ in the electron beam.

**7.6. General Considerations of the 5f Oxides.** With their relatively delocalized 5f valence electrons, actinides show a rich molecular chemistry, with oxidation states spanning from 3+ to 7+. In particular, actinides in high oxidation states 5+...
and 6+ can occur as a linear trans-dioxo actinyl cation, with two short highly covalent An=O bonds. These bonds are often considered as chemically inert. The actinyl cation forms bipyramidal polyhedra by bonding various number of equatorial ligands. Oxygen K-edge XAS is used to provide insights into the fundamental understanding of the participation of 5f and 6d orbitals to chemical bonding. Figure 53 presents the spectra of the light actinide dioxides: UO₂, NpO₂, and PuO₂, all in the cubic fluorite structure.

![Figure 53. Oxygen K-edge spectra of UO₂, NpO₂, and PuO₂.](https://dx.doi.org/10.1021/acs.chemrev.9b00439)

In light of DFT+U calculations, Modin et al. used oxygen K-edge XAS to compare the nature of the conduction band of these compounds. The oxygen K-edge XAS spectra of the three actinides show a first peak assigned to oxygen 2p states hybridized with An 5f and two peaks assigned to hybridized An 6d–O 2p bands split by crystal field. Differences in the XAS spectra arise from the relative position of the empty 5f and 6d bands. While in UO₂, the 5f and first 6d(e_g) band overlap, it appears that when changing from U to Np to Pu, the 5f band is lowered in energy and separates from the first 6d(e_g) band, in agreement with the increase of the number of 5f electrons from 2 in U⁴⁺ to 4 in Pu⁴⁺. Oxygen K-edge spectroscopy was used to probe the oxidation state of Pu in the PuO₂ single crystal. Oxygen K-edge XANES measured with STXM was applied to the characterization of sonochemical Pu colloids. The spatial resolution offered by the STXM technique enabled discovery of the contribution of the hydrolyzed species in hydrolytic colloids compared with sonochemical colloids.

7.6.1. Uranium Oxides. Thanks to its early discovery and high relative abundance, in addition to the relatively low activity of its radioisotopes, uranium is the most studied and best known actinide element. Understanding fully its chemical reactivity is of relevance for industrial, environmental, and health purposes. Efforts have been dedicated to determine the contribution of the 5f and 6p valence electrons to the chemical bonds. We discuss first the binary oxide UO₂ and subsequently the U⁶⁺ uranyl molecular complexes.

UO₂ is an important compound for nuclear fuel processing as well as a reference compound for the study of the fundamental physics of f electrons. The determination of the nature of the bottom of the conduction band is another important step in the physics of UO₂. Although known to be a Mott-Hubbard insulator, several studies aimed at discriminating between the f-f or f-d type. This information is relevant to understand the poor thermal conductivity of UO₂ as well as test the first-principle theoretical band structure predictions. Both are critical for the management of nuclear power plants. However, experimental proof is required to validate the first-principle calculations.

UO₂ formally has 5f⁶6d⁰ electronic configuration where U⁴⁺ is in a cubic site with eight equivalent U–O bonds. The crystal field in uranium splits the 6d states by 4.8 eV. The oxygen K-edge of UO₂ (Figure 53) analyzed by multiple-scattering and LSDA+U calculations provided proof of the f-f type Mott-Hubbard insulator. The oxygen K-edge XAS spectra have been calculated with LDA+U calculations.

The oxygen K-edge XAS has been used to study the evolution of UO₂-based nuclear fuel under irradiation. An early study of UO₂ pellets implanted with Mo ions used the low penetration depth of oxygen K-edge XAS to probe the oxidation state of uranium at the implanted surface. The observation of a low energy resonance after annealing was assigned to the presence of interstitial oxygens and the hyperstoichiometric nature of the surface. The thesis of Pizzi on the influence of the partial pressure of oxygen on the oxidation state of uranium in UO₂ used the oxygen K-edge to investigate the contribution of oxidized phases. Pizzi highlighted the difference between oxygen K-edge data obtained in TEY vs TFY modes: the TEY data from Wu et al. contain the contribution of an oxidized layer, which is absent in the data measured in TFY.

In ambient conditions, uranium is most stable in its oxidized U⁶⁺ uranyl state and tends to form strongly covalent linear trans-dioxo uranyl species. Understanding the electronic structure of the uranyl species has been the subject of many investigations to determine its reactivity. Denning et al. used linear polarized X-rays to distinguish between σ (parallel to the uranyl bond) and π (perpendicular to the uranyl bond) orbital symmetry of the antibonding orbitals of the uranyl in a Cs₂UO₂Cl₄ single crystals, which contains oxygen ligands only in the uranyl species (Figure 54). Comparison of the parallel and perpendicular polarized spectra with the molecular orbital diagram of the uranyl cation calculated by DFT with the ADF code, allows assignment of the three main peaks of the oxygen K-edge XAS spectrum to 3σ_u(5f⁶), 5σ_g(5f⁶), 4σ_g(6d⁰), and 2σ_g(6d⁰). A further study of these antibonding valence states of U⁶⁺ was performed on Ba₂ZnUO₆ where all oxygen in the U coordination sphere are interstitial oxygens and the hyperstoichiometric nature of the surface. The thesis of Pizzi on the influence of the partial pressure of oxygen on the oxidation state of uranium in UO₂ used the oxygen K-edge to investigate the contribution of oxidized phases. Pizzi highlighted the difference between oxygen K-edge data obtained in TEY vs TFY modes: the TEY data from Wu et al. contain the contribution of an oxidized layer, which is absent in the data measured in TFY.
ing π* from the σ* orbitals, and pDOS calculations allow assigning the transitions to the 5f and 6d. The intensity of oxygen K-edge XANES features can be used as a fingerprint for the hybridization strength. Fillaux et al. showed that the nature of the equatorial ligands strongly modifies the relative energies of the antibonding molecular orbitals.

In the field of geochemistry, Ward et al. published a compilation of oxygen K-edge data of common uranyl-bearing minerals, including carbonates, oxyhydroxides, phosphates, and silicates. All spectra present a first sharp peak assigned to the 3p(σ) of the uranyl and other features at higher energy depending on the compound. The authors showed that the uranyl compounds can be distinguished based on their oxygen K-edge spectra. This opens perspectives for mineral identification using STXM.

8. OUTLOOK

In this review, we have discussed the general analysis of the oxygen K-edge X-ray absorption spectra of atoms, molecules, and solids. An important experimental fact is that the oxygen K-edge spectrum is broadened by the oxygen 1s core–hole lifetime broadening of 300 meV. Since 1987, the resolution of most synchrotron beamlines is superior to this resolution, and this is also the case for most EELS measurements. This implies that the published oxygen K-edges essentially did not change in resolution over the last 30 years.

We have reviewed the theoretical analysis and simulations of oxygen K-edges. In principle, the procedure to calculate oxygen K-edges is known, and in the first approximation, the oxygen K-edge corresponds to the empty oxygen p-projected states. More detailed calculations calculate the electron–hole excitations with TD-DFT or BSE. Some open issues remain in the interpretation of oxygen K-edges, in particular, (a) the best procedure to describe the core–hole effect, (b) the accurate calculation of the exact excitation energies, and (c) the (calculational) best way to describe dynamics.

Experimental progress is made in the combination of oxygen K-edges with (i) time-resolution with femtosecond resolution, (ii) spatial resolution both with X-rays and electrons, (iii) operando conditions, and (iv) resonant experiments such as RIXS, which we briefly discuss with regard to their possible future developments.

Time

Concerning the future of time-resolved oxygen K-edge measurements, there is much progress in the development of HHG sources and XFEL sources with femtosecond time-resolution. We expect that in the near future it will become possible to measure the pump–probe excited-state oxygen K-edges with good statistics and well-determined femtosecond time-resolution. Such measurements of the femtosecond dynamics of oxides will be important for the fundamental studies on strongly correlated oxides as well as for the study of the fundamental steps in chemical reactions.

Space

Electron microscopes have atomic resolution, and one can expect further detailed study of the oxygen K-edge spectra with atomic resolution of thin film oxides. X-ray microscopes will likely improve to better than 5 nm resolution, allowing the measurement of oxygen K-edges of systems that cannot be measured with electron microscopes. Combining X-ray microscopy with transmission, fluorescence yield, and electron yield, one can specifically study (a) the surface of active systems and (b) the distribution and chemical nature of dilute species.

Operando

With the further development of nanotechnology, one can expect that the design of dedicated reactor cells for active systems will improve, allowing the study of the oxygen K-edge of active systems. This includes the important applications in the fields of batteries and (electro)(photo)catalysis. An issue that will remain to complicate operando studies is the constant struggle with X-ray induced effects. A way to avoid X-ray induced effects is to constantly replace samples. For example, liquid jet design is developing quickly, allowing the measurement of sensitive systems.

Resonance

We expect that many new experiments will become possible using resonant experiments, including RIXS, resonant elastic scattering, and resonant diffraction. The experimental resolution and the detection efficiency of RIXS beamlines are constantly improving. This will allow a new level of RIXS experiments, in combination with time and/or spatial resolution. The time-resolved RIXS and spatially resolved RIXS experiments are expected to bring much new information on the detailed (dynamics of) the electronic structure.

In conclusion, there is a very interesting future for oxygen K-edge X-ray absorption spectra, not so much in the progress of their specific spectral shapes (as limited by the 1s lifetime), but rather in the combination of oxygen K-edges with time-resolved, spatially resolved, operando, and resonant experiments.

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Figure 54. Polarized oxygen K-edge spectrum of the uranyl UO2+2 species in Cs2UO2Cl4 single crystal. The σ and π characters of the probed states are revealed with the incident light polarization respectively parallel and perpendicular to the uranyl bond.
Notes
The authors declare no competing financial interest.

Biographies

Federica Frati received her bachelor in chemistry in 2014 from Perugia University, where she did undergraduate research under the direction of Professor F. Tarantelli and Leonardo Belpassi. She obtained her master degree with Professor Mauro Stener and Remco W.A. Havenith at Trieste University in 2016. She is currently a PhD candidate of the Department of Inorganic Chemistry and Catalysis at Utrecht University. Her primary research efforts are directed toward oxygen K-edge modeling in molecular and solid systems.

Myrtille Hunault is a beamline scientist at the SOLEIL synchrotron, France. She graduated from both the ESPCI ParisTech and the University Pierre and Marie-Curie in Paris, France, in 2011. She received her PhD in material chemistry in 2014 for the spectroscopic study of the colors of ancient glasses. After a 1-year postdoctoral fellowship to study the colors of the 15th century rose of the Sainte-Chapelle of Paris, she then went for a second postdoctoral fellowship in the Chemistry Department of Utrecht University to work on the fundamentals of X-ray spectroscopy of metal oxides. Since 2017, she has been working at the SOLEIL synchrotron, France. Her research focuses on the speciation and coloring role of transition metals and actinide ions in oxides, using optical and X-ray spectroscopies and theoretical calculations.

Frank de Groot is Professor of X-ray spectroscopy at the Chemistry department of Utrecht University. He received his MSc degree in Chemistry from Nijmegen University in 1986 and a PhD in Chemistry from Nijmegen University in 1991. He then went on to the LURE synchrotron in Orsay, France, and in 1995 to a KNAW academic research position at Groningen University. Since 1999, he has been working in the Chemistry Department of Utrecht University. His research is focused on X-ray spectroscopy for the study of the electronic and magnetic structure of condensed matter, in particular, for transition metal oxides and heterogeneous catalysts under working conditions.

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