Spectroscopic study of the electric field induced valence change of Fe-defect centers in SrTiO$_3$

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The electrochemical changes induced by an electric field in Fe-doped SrTiO$_3$ have been investigated by X-ray absorption spectroscopy (XANES and EXAFS), electron paramagnetic resonance (EPR) and Raman spectroscopy. A detailed study of the Fe dopant in the regions around the anode and cathode reveals new insights into the local structure and valence state of Fe in SrTiO$_3$ single crystals. The \textit{ab initio} full multiple-scattering XANES calculations give an evidence of the oxygen vacancy presence in the first coordination shell of iron. Differences in the length and disorder of the Fe–O bonds as extracted from EXAFS are correlated to the unequivocal identification of the defect type by complementary spectroscopical techniques to identify the valence state of the Fe-dopant and the presence of the Fe – VO complexes. Through this combinatorial approach, novel structural information on Fe – VO complexes is provided by X-ray absorption spectroscopy, and the relation of Fe–O bond length, doping level and oxidation state in SrTi$_{1-x}$Fe$_x$O$_3$ is briefly discussed.

1 Introduction

In recent years, considerable effort has been spent on the investigation of the valence state of 3d transition metal dopants in the perovskite SrTiO$_3$ by the use of X-ray absorption spectroscopy. In Fe-doped SrTiO$_3$, the mixed valence (Fe$^{3+}$/Fe$^{4+}$) character of the dopant creates a charge imbalance with respect to the lattice, which is compensated by the creation of oxygen vacancies\(^1\) to conserve charge neutrality. Redox-type reactions of the crystal affect the balance between Fe$^{3+}$ and Fe$^{4+}$, and therefore the oxygen vacancy concentration in the material. It has been shown that such redox reactions, induced by thermal treatment under oxidizing or reducing atmosphere, influence the X-ray absorption near-edge structure (XANES) of transition metal dopants, while the XANES of the Ti K-edge remains unchanged.\(^2\) The strong ionicity of the constituent species makes SrTiO$_3$ susceptible to chemical changes induced by an electric field,\(^3,4\) resulting in a concentration polarization of oxygen vacancies that is visible as a color-change taking place at the anode and cathode (electrocoloration).\(^5\) This effect gained new interest in the last decade since the related insulator-to-metal transition (electroformation) in SrTiO$_3$ and other transition metal oxides can be exploited for the fabrication of non-volatile resistive random access memory (RRAM) devices.\(^6,9\) Microfocus X-ray absorption spectroscopy\(^8\) at the K-edge of the Cr-dopant proved the redistribution of oxygen vacancies during the electroforming process in SrTiO$_3$. However, despite of one singular report,\(^10\) the valence change in the Ti sublattice is generally insufficient to be detected at the Ti K-edge since oxygen vacancies are preferentially formed at the acceptor sites.\(^5\)

The chemical equilibrium between charged defects poses an inherent challenge for XAFS investigations since it is almost impossible to obtain “pure” samples having only one type of defect. It is therefore necessary to deconvolute the measured signal using either appropriate standards or theoretical approximations. In addition, the absorption characteristics are strongly influenced by the polarized bonding environment in transition metal oxides. The connection between the formal valence state and the position and structure of the absorption edge is complicated by the fact that a preferred ground state configuration may be stabilized by a ligand-hole,\(^11\) and by the influence of distortions in the coordination polyhedra, coordination number and site symmetry, all of which may be present in the probed volume.

In the present work, we investigate the valence state and the local environment of Fe in a Fe-doped SrTiO$_3$ single crystal after electrocoloration by a dc electric field using a combinatorial approach of three spectroscopic techniques. The concentration polarization associated with the electrocoloration creates an oxygen-rich region around the anode and a strongly
oxygen-deficient region around the cathode. Using the fact that Raman spectroscopy and electron paramagnetic resonance (EPR) are sensitive only to a specific defect configuration of Fe in SrTiO$_3$, it is the scope of this work to probe the respective defects using locally sensitive experimental techniques such as X-ray absorption spectroscopy. It should be noted that the concentration polarization induced by the dc electric field is only indirectly connected to the insulator-to-metal transition taking place locally confined between the electrodes.

2 Experiment

A 0.2 wt% Fe-doped SrTiO$_3$ single crystal was electrocolored using two Au-electrodes evaporated onto the (001)-oriented, polished crystal. A constant current was increased stepwise from 1 nA to 0.1 mA over the duration of 2 weeks, until finally a breakdown in resistance occurred and a current of 10 mA flowed. The crystal was kept at a temperature of 300 °C at a vacuum pressure of $10^{-8}$ mbar. The coloration of the crystal is depicted in Fig. 1, along with an indication of the anodic and cathodic regions that were investigated in this work. After XAFS and Raman measurements, the crystal was cut into slices for the EPR spectrometer.

XAFS experiments were performed at beamlines BM29 (ESRF, Grenoble) and A1 (HASYLAB, Hamburg). X-Ray absorption near edge spectroscopy (XANES) and extended X-ray absorption fine structure (EXAFS) were measured in fluorescence step-by-step (single crystal) and transmission mode (powder references) using monochromatic X-ray radiation, using a Si(111) double-crystal monochromator with an energy resolution of $\Delta E/E < 10^{-4}$ at 7000 eV photon energy. The size of the beam at the sample was typically $1 \times 8$ mm$^2$. XANES of the Fe K-edge (Ti K-edge) spectra were typically recorded within an interval of 100 eV (50 eV) around the absorption edge, with a constant energy step of 0.5 eV (0.2 eV). The Fe K- and Ti K-edge EXAFS spectra were recorded using an optimized energy mesh with a step of 0.2 eV around the absorption edge and a constant $\Delta k$ step far above the edge.

Micro-Raman spectra were collected at 20 °C using a confocal microscope with spectrometer Nanofinder-S (SOLAR TII, Ltd.). The measurements were performed through a Nikon Plan Fluor 40 × (NA = 0.75) optical objective. The Raman spectra were excited by a He–Cd laser (441.6 nm, 50 mW cw power) and dispersed by 600 grooves mm$^{-1}$ diffraction grating mounted on the 520 mm focal length monochromator. A Peltier-cooled back-thinned CCD camera (ProScan HS-101H, 1024 × 58 pixels) was used as a detector. The elastic laser light component was eliminated by an edge filter (Omega, 441.6ÅELP-GP). Similar results (not shown here) were also obtained under excitation by a 532 nm DPSS solid state laser. The EPR spectra have been recorded with a X-band ($\nu = 9.323$ GHz) spectrometer with a modulation of 100 kHz. In order to study the angular dependence, thin slices of the Fe-doped single crystal were cut along the [010]-axis and aligned with [010] parallel to the rotation axis of the spectrometer. The dimensions of each piece were about $1 \times 10 \times 1$ mm$^3$. The samples were mounted on the $xyz$ orthogonal laboratory frame chosen for the EPR anisotropy pattern and related to the crystallographic axes as follows: $x \parallel [001]$, $z \parallel [010]$, $y \parallel [100]$. The presented measurements were done at room temperature, while additional measurements at liquid nitrogen temperature confirm the presented results.

3 Data analysis

The XANES/EXAFS data treatment and fitting was done using the EDA software package. The XANES region of the absorption edge was extracted by background subtraction and normalization to 100 eV above the edge. The EXAFS signal was extracted up to $k = 10$ Å$^{-1}$ after standard background subtraction, and the single-scattering contributions of the first shell were isolated by Back Fourier Transformation (BFT) in the interval 0.8–2.1 Å. The single-shell fitting of the EXAFS spectra was done using the single-scattering curved-wave formalism in $k$-space after BFT. In this paper, the EXAFS data were analyzed using two different approaches: the backscattering phases and amplitudes were either calculated by the FEFF8 code or obtained experimentally from the polycrystalline reference compound LaFeO$_3$.

As the threshold energy of the photoelectron $E_0$ is defined in the FEFF8 code relative to the Fermi level and depends on the muffin-tin radii, the spectra have to be corrected to avoid $E_0$ difference errors in the fitting process. Consequently, the phase differences between theoretical and experimental spectra were set to zero at low $k$-values, according to Bunker and Stern’s criterion, and $E_0$ was allowed to vary for fine adjustment during the fitting procedure.

The calculations by the FEFF8 code were performed for a cluster having a radius of 8 Å, which mimics the environment of the Fe ions in an SrTiO$_3$ crystal (lattice parameter $a_0 = 3.905$ Å) and centered on a Fe atom that has been substituted for Ti. A muffin-tin cluster potential was calculated in a self-consistent way, and the complex Hedin–Lundqvist exchange–correlation potential was used to account for inelastic effects (see the details in ref. 17). Experimental backscattering phases and amplitudes were extracted from the experimental EXAFS data measured on a polycrystalline LaFeO$_3$ powder at room temperature.
These functions were obtained assuming the iron coordination number $N = 6$ and medium Fe–O interatomic distance $R = 2.007 \, \text{Å}$, from the crystallographic data, and the mean-square relative displacement (MSRD) Fe–O $\sigma^2 = 0.0033 \, \text{Å}^2$. The MSRD value was obtained from the best fit of the Fe K-edge EXAFS signal for LaFeO$_3$ using the single-shell Fe–O model with the theoretical phase and amplitude, calculated by the FEFF8 code for the LaFeO$_3$ structure. While LaFeO$_3$ is orthorhombic (space group $Pnma$), the distortion of the Fe–O$_6$ octahedra is smaller than the measurement uncertainty ($\sim 0.005 \, \text{Å}$), and therefore the first shell was treated as a perfect octahedron ($R_{Fe-O} = 1.994 \pm 0.005 \, \text{Å}$). The good quality of the fit and the small MSRD obtained for the Fe–O bond ($\sigma^2 = 0.0033 \, \text{Å}^2$) in LaFeO$_3$ confirms the validity of this approach.

Note that the use of experimental phases and amplitudes for EXAFS allows, to a certain extent, the compensation of systematic errors as they include the contribution of the mean free path, the multi-electron amplitude reduction factor $S^0$, glitches and resolution, thus reducing the number of adjustable parameters and consequently increasing the reliability of the fitted results. To ensure the phases and amplitudes transferability, and to allow estimation of systematic errors, all the data were analyzed in a similar way, using also the theoretical phases and amplitudes, after procedures and parameters.

The fitting procedure was done using a parameter set of the disorder parameter MSRD $\sigma^2$ (describing the disorder in the bond length), the distance $R$, the amplitude reduction factor $S^0$, glitches and resolution, thus reducing the number of adjustable parameters and consequently increasing the reliability of the fitted results. To ensure the phases and amplitudes transferability, and to allow estimation of systematic errors, all the data were analyzed in a similar way, using also the theoretical phases and amplitudes, after procedures and parameters.

To provide a framework for the interpretation of the single crystal spectra, the EXAFS of powder references SrTi$_{0.9}$Fe$_{0.1}$O$_3$ and SrTi$_{0.9}$Fe$_{0.1}$O$_{2.95}$ were fitted as well. Note that the data on SrTi$_{0.9}$Fe$_{0.1}$O$_3$ and SrTi$_{0.9}$Fe$_{0.1}$O$_{2.95}$ calculated with the theoretical phase and amplitude were previously published. In the first step, the experimental spectrum of powder reference compound LaFeO$_3$ was analyzed using the theoretically calculated scattering phase and amplitude. In the second step, all experimental spectra from powder references and the single crystal were treated with the experimental phase and amplitude extracted from the LaFeO$_3$ spectra, as well as with theoretical phase and amplitude.

Self-consistent, real-space multi-shell-scattering XANES calculations of the Fe K-edge in Fe-doped SrTiO$_3$ have been performed by the FDMNES code$^{19,20}$ in the one-electron electric dipole ($\Delta l = 1$, $1s \rightarrow n\sigma$) and quadrupole ($\Delta l = 2$, $1s \rightarrow nd$) approximations. We considered four structural models based on the clusters, centered at the absorbing Fe atom and constructed from the cubic SrTiO$_3$ crystal structure (lattice parameter $a_0 = 3.905 \, \text{Å}$). The radius of the clusters was equal to 6 Å, being sufficiently large to reproduce the main features in the experimental Fe K-edge XANES signal. In the first model (SrTiO$_3$:Fe), the substitution of one Ti atom by the Fe atom was performed. In the second model (SrTiO$_3$:Fe with JT), the Jahn–Teller (JT) distortion$^{17}$ of the first coordination shell of the absorbing iron atom was taken into account: the [FeO$_6$] octahedron was distorted by placing four in-plane oxygen atoms at 1.89 Å from iron and two oxygen atoms at 2.10 Å. In the third model (SrTiO$_3$:Fe + VO$^-$), no JT distortion was present, but an oxygen vacancy (VO$^-$) was introduced into the first coordination shell of the Fe atom, i.e. one of the six oxygen atoms was eliminated, so that the iron atom becomes five-fold coordinated. Finally, in the fourth model (SrTiO$_3$:Fe + VO$^{2\text{+}}$), no JT distortion was present, but an oxygen vacancy (VO$^-$) was introduced into the second coordination shell of the Fe atom. The self-consistent cluster potential was calculated within the non-muffin-tin approximation using the finite difference method (FDM), in which the full cluster potential is set on an equally spaced 3D-grid. The real Hedin–Lundqvist exchange-correlation potential was employed in the calculations, and the final excited state of the photoelectron included a screened core-hole. Note that the calculated XANES signals were intentionally not broadened to underline the fine structure details.

### 4 Results and discussion

The generally accepted model of electrocoloration in Fe-doped SrTiO$_3$ is based on the field-driven migration of oxygen vacancies and the corresponding chemical changes at the Fe centers that give rise to the color change. In the low concentration (dilute) regime where the Fe centers can be considered as isolated, a set of equations can be used to describe the defect equilibria in the ionic crystal. Of special interest for this work are the conservation conditions:

\[
[F]_{\text{tot}} = [Fe^{3+}] + [Fe^{4+}] \tag{1}
\]

\[
2[VO] + [\mathcal{E}] = [Fe^{3+}] + [\mathcal{E}] \tag{2}
\]

In particular, the weak band-gap excitation (no free charge carriers) allows the simplification of eqn (2):

\[
2[VO] \approx [Fe^{3+}] \tag{3}
\]

This charge neutrality condition imposes some constraints on the defect structure that can be present in the material. Most notably, even under completely reduced conditions, eqn (3) demands that the oxygen vacancy concentration will not be more than half of the total Fe concentration, assuming that every Fe center has been reduced to Fe$^{3+}$ and that other mechanisms of charge compensation can be neglected. For strongly reduced crystals, the charge neutrality condition has to be expanded to accommodate the compensation of oxygen vacancies by free electrons, causing the appearance of Ti$^{3+}$ ions through the occupation of conduction band states. To verify the validity of eqn (3) for the present work, it is necessary to monitor the chemical changes induced by the electrocoloration at both the Fe- and Ti-site. Since there is no observable change in the Ti K-edge XANES (Fig. 2), and the average Ti–O bond distance as extracted from EXAFS data is identical in the anodic and cathodic regions (within the measurement uncertainty, data not shown), it follows that the oxidation/reduction process takes place preferentially at the Fe-site and that the changes at the Ti-site are too low to be detectable.

Therefore, the following sections are concerned with the structural and chemical changes at the Fe-site induced by the electrocoloration process, and eqn (3) will be regarded as valid.

#### 4.1 EPR studies

Fig. 3 shows the EPR spectra of Fe$^{3+}$ of the cathodic (upper, red line) and anodic regions (lower, black line) of the doped...
SrTiO$_3$ single crystal, with the static magnetic field parallel to the crystallographic [010] direction. The EPR signal in the anodic region consists of a strong, isotropic central line, with 4 symmetrically split satellite lines showing an angular variation identical to that found in ref. 22, with the isotropic g-factor of 2.00 corresponding to cubic Fe$^{3+}$ centers and with a cubic splitting parameter of $a \approx 0.02$ cm$^{-1}$.

The EPR signal of the cathodic region consists of two structurally non-equivalent centers, a cubic center equivalent to that in the anodic region and an axial center with a strongly anisotropic g-factor, as is well described in the literature. Kirkpatrick et al. found these lines with g$_{||}$ = 5.993 and g$_{\perp}$ = 2.0054 to correspond to cubic Fe$^{3+}$ centers and with a cubic splitting parameter of $a \approx 0.02$ cm$^{-1}$.

The angle of rotation corresponds to the magnetic field lines and the crystallographic [100] axis.

We can conclude that after the electrocoloration process, the axial Fe$^{3+}$ center in the anodic part has disappeared almost completely, while the signal of the cubic Fe$^{3+}$ center is decreased by about a factor of 7 ($\pm$1) with respect to the cathodic part. It should be mentioned that the Fe$^{4+}$ center created in the anodic region is a “non-Kramers” ion due to the large contribution of the d$^5$ configuration to the ground state, and therefore does not appear in the spectra. There are no other EPR-active impurity centers with significant contributions to the EPR spectra.

### 4.2 Raman spectroscopy

Since SrTiO$_3$ at room temperature has a cubic structure (space group Pm$ar{3}$m) with a center of inversion, the undoped crystal exhibits no first-order Raman scattering, and only second-order effects are observed in the Raman spectra. An extensive examination of the Raman spectra of oxidized and reduced SrTi$_{1-x}$Fe$_x$O$_3$, solid solution powders has been given in ref. 17, and the existence of a first-order Raman peak at 690 cm$^{-1}$ has been attributed to doping with Fe$^{4+}$ ions. In addition, three “forbidden” phonon modes have been weakly visible: LO3, TO4 and LO4 at 475 cm$^{-1}$, 545 cm$^{-1}$ and 795 cm$^{-1}$, respectively. On the other hand, Fe$^{3+}$ does not induce any change in the Raman spectrum with regard to that of an undoped crystal. Therefore, the appearance of the strong peak at 690 cm$^{-1}$ can be seen as a “fingerprint” indicating the presence of Fe$^{4+}$ in the crystal.

In the present work, Raman spectroscopy was performed close to the anode and cathode of the Fe-doped SrTiO$_3$ single crystal, using a confocal Raman microscope with a spatial resolution of 1 µm. The recorded spectrum (Fig. 5) of the anodic (solid black line) region shows the distinct features that characterize the oxidized SrTi$_{1-x}$Fe$_x$O$_3$ powders as described above. In the spectrum of the cathodic region (dashed red line), the additional peak as well as the weak phonon modes are notably absent, indicating that the amount of Fe$^{4+}$ in the cathodic region is too low to be detected by Raman spectroscopy. From the Raman data, we can conclude that the cathodic region does not contain significant amounts of Fe$^{4+}$, while the anodic region contains predominantly Fe$^{4+}$.
4.3 XANES studies

A comparison of the experimental Fe K-edge XANES for the cathodic and anodic regions in the Fe-doped SrTiO₃ single crystal (Fig. 6(a)) indicates that while the pre-edge peak located at 7115 eV is similar in both cases, the fine structure located at higher energies is different. In particular, the shoulder at 7123 eV is more pronounced in the cathodic region, and the main absorption edge shifts by about \( \sim 0.5 \) eV to higher energies in the anodic region. Reference XANES data (from ref. 17) of completely oxidized and reduced \( \text{SrTi}_1-x\text{Fe}_x\text{O}_3 \) solid solution powders show an almost identical shift of the main absorption edge (Fig. 6(b)). Comparing the XANES data for the single crystal to the reference data, it is obvious that the changes taking place during electrocoloration affect the valence state of the Fe atoms in a reduction/oxidation process, leaving Fe centers predominantly oxidized in the anodic region and reduced in the cathodic region.

Deeper understanding of the Fe K-edge XANES signals has been obtained using theoretical simulations, performed for the four models as described in Section 3. The results of the XANES calculations are shown in Fig. 7 and are compared with the experimental data for the cathodic and anodic regions in Fig. 8. There are four features (A, B, C, D) in the XANES signals that can be distinguished below the main absorption edge \( E \). At the Fe K-edge, the \( 1s^2(\text{Fe}) \) core-electron is excited, following dipole selection rules, to empty final-states with \( p \)-character, located above the Fermi level and being relaxed due to the presence of the positively charged core-hole. Such transitions give rise to peaks starting from B and above (Fig. 7). The transition of the \( 1s \) electron to the conduction band of cubic Fe-doped \( \text{SrTiO}_3 \) (peak A in Fig. 7), composed mainly of the \( 3d \)-states of the Ti(Fe) ions, is forbidden in the dipole approximation. However, it becomes allowed through symmetry lowering due to the Jahn–Teller distortion, the presence of an oxygen vacancy in the first coordination shell or within the quadrupole approximation. Note that generally, the small cross section of the \( 1s \rightarrow 3d \) quadrupole transition is compensated by a high density of \( 3d \) states.

Next we will discuss the peculiarities of the Fe K-edge XANES signals for the four theoretical models (Fig. 7). By looking at the first and second models, corresponding to the regular and JT-distorted iron environments, respectively, one can see that the presence of the JT distortion leads mainly to a decrease of the main peak \( E \) amplitude and a splitting of peak C. Thus the JT distortion cannot explain an increase of the absorption in the energy range from 10 to 15 eV and, also, the origin of the shoulder around 30 eV in the experimental data (Fig. 8).

Next two models take into account the possible presence of the oxygen vacancy (\( V_{\text{O}} \)) in the first or second coordination shell of iron. Contrary to the case of the JT distortion, an
introduction of the single oxygen vacancy into the first coordination shell of iron (the third model) makes the theoretical Fe K-edge XANES signal closer to the experimental one (Fig. 8). The most prominent effect is observed in the range of 4–12 eV above the Fermi level. Here two new peaks B and D appear, and peak C becomes split. The position of peak D corresponds well to the energy region, where the experimental XANES signals for the cathodic and anodic regions differ. Therefore, the occurrence of peak D can be used as a “fingerprint” to trace the presence of an oxygen vacancy in the first coordination shell of iron.

Note that placing the single oxygen vacancy into the second coordination shell of iron (the fourth model) results in the Fe K-edge XANES signal being very similar to that without vacancy (Fig. 7). Such result can be expected since there are 24 oxygen atoms in the second shell, so that the absence of one atom produces negligible influence on the total XANES signal.

To conclude, we attribute a variation of the absorption intensity observed between the two experimental Fe K-edge XANES signals from the anodic and cathodic regions of the Fe-doped SrTiO3 single crystal to the appearance of an oxygen vacancy in the first coordination shell of iron.

The experimental EXAFS $\chi(k)$ spectra for the anodic and cathodic regions are shown in Fig. 9. Their Fourier transforms (FTs), shown in Fig. 10, indicate a change of the first peak position (better seen in the imaginary part of FTs) at ~1.4 Å due to the first coordination shell of iron. At the same time, the position of the second and third coordination shells (peaks in the region 2.5–4 Å, Fig. 10) remains unchanged. This suggests that the electrocoloration process induces a difference in the average Fe–O bond distance in the anodic and cathodic regions of the single crystal.

4.4 EXAFS studies

The experimental EXAFS $\chi(k)$ spectra for the anodic and cathodic regions are shown in Fig. 9. Their Fourier transforms (FTs), shown in Fig. 10, indicate a change of the first peak position (better seen in the imaginary part of FTs) at ~1.4 Å due to the first coordination shell of iron. At the same time,
Fe$^{3+}$ ($d^6$ configuration) in SrTiO$_3$.

From the EXAFS and XANES analyses as well as Raman, it follows that Fe is predominantly tetravalent in the anodic region, albeit with a small admixture of Fe$^{3+}$ (cubic center, $d^5$ configuration) as shown by EPR (see Section 4.1).

In contrast, the Fe–O bond length in the cathodic part (1.972 Å) is 4 pm longer than in the anodic region, while the MSRD decreases significantly. The EPR and Raman results show the existence of cubic Fe$^{3+}$ centers in this region, albeit with an admixture of Fe$^{3+} – V_O$ complexes, and no Fe$^{4+}$. The Fe$^{3+}$–O distance in reduced SrTi$_{0.9}$Fe$_{0.1}$O$_{2.95}$ (1.955 Å) is intermediate between that in SrTi$_{0.5}$Fe$_{0.5}$O$_{2.5}$ (1.960 Å) and the Ti–O distance in SrTiO$_3$ (1.951 Å), while the Fe–O distance in the cathodic region of the single crystal is significantly longer. In addition, the slightly larger MSRD value observed in SrTi$_{0.9}$Fe$_{0.1}$O$_{2.95}$ compared with the cathodic part suggests a larger distortion in the first coordination shell around Fe$^{3+}$ in the reduced solid solution.

5 Discussion

The experimental determination of the Fe$^{3+} – V_O$ concentration has been shown to be possible via double-integration EPR techniques, and also attempted via linear combination of XANES spectra. In principle, the relative concentrations of Fe centers can be estimated from linear combination of the Fe-O bond distances using suitable standards. Care has to be taken to determine which type of Fe center is present in the analyzed sample. Following the EPR and Raman results, we can assume that the only defect centers present in the anodic region of the single crystal are 6-fold coordinated Fe$^{3+}$ and Fe$^{4+}$ without any Fe$^{5+} – V_O$. Suitable standards are LaFeO$_3$ and SrFeO$_3$ with Fe–O distances of 1.994 Å and 1.925 Å, respectively. For these compounds, it has been shown that A-site substitution of Sr for La decreases the size of the unit cell, despite the smaller Shannon radius of the La$^{3+}$ ion, and the Fe–O bond distance has been shown to decrease continuously with increasing amount of Fe$^{4+}$ going from LaFeO$_3$ to SrFeO$_3$ without systematic deviations from the Vegard’s law. Assuming a linear relation of the Fe–O distance to concentration, the Fe$^{3+}$ concentration in the anodic part can be estimated to be approximately 10%. Integrating the EPR signals, we can further estimate the ratio (cathode to anode) of Fe$^{4+}$ to be about 7 (±1), which puts the concentration of the cubic Fe$^{3+} – V_O$-center at roughly 70% (±10%) in the cathodic region. Since Raman spectroscopy demonstrates the absence of Fe$^{4+}$ in this region, consequently the amount of Fe$^{3+} – V_O$ can be estimated as 30%. Unfortunately, the above estimation cannot be used to gain information on the Fe–O bond length in the Fe$^{3+} – V_O$ complexes because of the tetragonal distortion of the axial site.

Merkle and Maier demonstrate that the degree of association of Fe and oxygen vacancies shows a tendency to decrease with increasing doping level and temperature. For a similar doping concentration as used in this study, the degree of association at room temperature is given as ~25%, while values approaching 50% are only seen for lower concentrations and temperatures below 273 K. Since these values are in agreement within the error bar specified above, a Fe$^{3+} – V_O$ concentration of 30% is a realistic estimation.

From the EXAFS and XANES analysis as well as from EPR data, it is reasonable to conclude that Fe exists mainly as a cubic Fe$^{3+}$ center with Fe–O distance $R = 1.972$ Å in the cathodic region of the single crystal. The concentrations of the cubic Fe$^{3+} – V_O$ and the axial Fe$^{3+} – V_O$ center are in agreement with the defect chemical constraint $[V_O]/[Fe^{3+}] < 0.5$ for the dilute regime (eqn (3), Section 4), with the amount of the Fe$^{3+} – V_O$ present around the cathode estimated to be ~30%. As SrTi$_{0.9}$Fe$_{0.1}$O$_{2.95}$ is far outside the dilute regime, the constraints described by eqn (3) are no longer valid and Fe$^{3+} – V_O$ with an Fe–O distance of $R = 1.955$ Å becomes the predominant Fe center.

6 Summary and conclusions

In this work the electrochemically induced changes in the local structure and valence state of iron atoms have been studied in electrocolored Fe-doped SrTiO$_3$ single crystals by EPR, Raman scattering and X-ray absorption (XANES and EXAFS).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Best-fit structural parameters ($N$ is the coordination number, $R$ is the interatomic distance, and $\sigma^2$ is the mean-square relative displacement (MSRD)) for the first coordination shell of iron obtained from the analysis of the Fe K-edge EXAFS spectra using experimental backscattering phase and amplitude functions extracted from LaFeO$_3$</th>
<th>SrTi$<em>{0.9}$Fe$</em>{0.1}$O$_{3}$</th>
<th>SrTi$<em>{0.9}$Fe$</em>{0.1}$O$_{2.95}$</th>
<th>Fe:SrTiO$_3$ (anode)</th>
<th>Fe:SrTiO$_3$ (cathode)</th>
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<tr>
<td>$N$</td>
<td>$5.4 \pm 0.1$</td>
<td>$5.0 \pm 0.1$</td>
<td>$5.7 \pm 0.1$</td>
<td>$5.4 \pm 0.1$</td>
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<tr>
<td>$R$(Fe–O)/Å</td>
<td>$1.931 \pm 0.005$</td>
<td>$1.955 \pm 0.005$</td>
<td>$1.932 \pm 0.005$</td>
<td>$1.972 \pm 0.005$</td>
<td></td>
</tr>
<tr>
<td>$\sigma^2$(Fe–O)/Å$^2$</td>
<td>$0.0094 \pm 0.0005$</td>
<td>$0.0072 \pm 0.0005$</td>
<td>$0.0098 \pm 0.0005$</td>
<td>$0.0056 \pm 0.0005$</td>
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spectroscopy. With EPR spectroscopy, we demonstrate the presence of axial Fe$^{3+}$ – V$_O$ centers in the cathodic region, with the characteristic angular dependence of the axial center, although the majority of Fe centers in the cathodic region are cubic Fe$^{3+}$ centers. The anodic region of the electrocolored crystal is highly oxidized since the signal associated with the Fe$^{3+}$ – V$_O$ complexes vanishes almost completely. Since EPR is not sensitive to Fe$^{4+}$ ions (d$^0$ configuration), Raman spectroscopy complements the EPR technique as the Raman line at 690 cm$^{-1}$ is a ”fingerprint” of Fe$^{4+}$ in Fe-doped SrTiO$_3$. The strong appearance of this line in the anodic part is due to the predominance of Fe$^{4+}$ in this region, whereas its disappearance in the cathodic part demonstrates the absence of Fe$^{2+}$.

As shown by ab initio full-multiple-scattering XANES calculations, a variation of the experimental Fe K-edge XANES observed between the anodic and cathodic regions can be attributed to the change in the amount of oxygen vacancies in the first coordination shell of iron atoms, in agreement with the EPR results.

EXAFS analysis yields a significant increase of the average Fe–O distance by $\sim 0.03$–0.04 Å in the cathodic region, compared to both the anodic region and the SrTi$_{0.97}$Fe$_{0.03}$O$_3$ powder reference. By combination of EPR and EXAFS, we show that this increase is caused mainly by the presence of cubic Fe$^{3+}$ centers in the cathodic region, whereas the smaller Fe–O distance in the powder can be attributed to a much higher concentration of oxygen vacancies compared to the single crystal. The shorter Fe–O distance in the anodic part is characteristic of Fe$^{4+}$ centers, in good agreement with the SrTi$_{0.97}$Fe$_{0.03}$O$_3$ powder reference and observations by Raman spectroscopy.

In the dilute regime, the Fe-centers can be considered as isolated and non-interacting, and the large Fe$^{2+}$–O bond length is evidence of a local expansion of the crystal lattice around cubic Fe-centers (Fe$^{3+}$–O$_6$). This suggests that the formation of Fe$^{3+}$ – V$_O$ complexes is favored by the minimization of strain energy in the crystal in addition to the association enthalpy. Above the dilute limit, the reduced distance between Fe-centers provides some strain relaxation which results in smaller Fe–O distances even in the reduced material. These considerations enable the determination of the Fe valence state in the dilute regime based on the X-ray absorption characteristics.

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**References**