Probing the Meta-Stability of Oxide Core/Shell Nanoparticle Systems at Atomic Resolution

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ABSTRACT: Hybrid nanoparticles allow exploiting the interplay of confinement, proximity between different materials and interfacial effects. However, to harness their properties an in-depth understanding of their (meta)stability and interfacial characteristics is crucial. This is especially the case of nanosystems based on functional oxides working under reducing conditions, which may severely impact their properties.
In this work, the in-situ electron-induced selective reduction of Mn$_3$O$_4$ to MnO is studied in magnetic Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticles by means of high-resolution scanning transmission electron microscopy combined with electron energy-loss spectroscopy. Such in-situ transformation allows mimicking the actual processes in operando environments. A multi-stage image analysis using geometric phase analysis combined with particle image velocity enables direct monitoring of the relationship between structure, chemical composition and strain relaxation during the Mn$_3$O$_4$ reduction. In the case of Fe$_3$O$_4$/Mn$_3$O$_4$ core/shell the transformation occurs smoothly without the formation of defects. However, for the inverse Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell configuration the electron beam-induced transformation occurs in different stages that include redox reactions and void formation followed by strain field relaxation via formation of defects. This study highlights the relevance of understanding the local dynamics responsible for changes in the particle composition in order to control stability and, ultimately, macroscopic functionality.

**Keywords:** Magnetic oxides, core-shell nanoparticles, scanning transmission electron microscopy, in-situ transformations
Highlights

• In situ, atomic resolution, electron microscopy study of the electron beam induced Mn3O4 → MnO reduction reaction in Fe3O4/Mn3O4 and Mn3O4/Fe3O4 core/shell nanoparticles.

• In depth analysis of the evolution of the microstructure with complex analysis techniques, including particle image velocimetry analysis and geometric phase analysis.

• Very different evolution of the reaction depending on the morphology, i.e., Mn3O4 in the core or in the shell.

• Multistage complex processes taking place in the reduction process in the Mn3O4/Fe3O4 with generation of voids, stress relaxation, ionic transport and formation of defects.
1. Introduction

Hybrid nanoparticles [1] are very attractive as model systems for fundamental studies [2-6] and also for a very wide range of applications. The combination of reduced dimensionality effects with the interplay between the properties of the constituents can give rise to multiple, and sometimes new, functionalities in systems of interest for catalytical [7], optical [8,9] magnetic [10,11] or biomedical applications [12-15]. Such multifunctional nanoparticles can be obtained in different configurations, being the dimer and core/shell geometries the most usual archetypes [1]. Interestingly, their inherent metastability can also be exploited to design novel architectures such as those in galvanic replacements [16-20]. Dimer systems exploit proximity effects as source of extra functionality, but the contact interface area is limited thus precluding a large contact surface to promote interaction between the components. On the other hand, the encapsulation of core material into a core/shell structure promotes a large contact surface area, since the whole surface of the core becomes the interface with the shell. Additionally, it allows controlling the structure degree of freedom in the form of lattice constraints that permit modifying the system structure and properties. Interface related lattice constraints can result in thermodynamically metastable systems which may phase-segregate or homogenize due to ion diffusion upon heating, during electrochemical cyclic, or in long term ambient conditions [21], relevant to processes such as redox limited performance of energy materials. In this context, the study of processes that mimic the destabilization of hybrid core/shell nanoparticles under controlled conditions can facilitate the understanding of their behavior in real applications (e.g., in the study of cathode materials for batteries or electrochemical capacitors) and open new avenues to create improved structures [22-24].
This is particularly the case with nanostructured systems based on complex oxides, which are very robust materials displaying a wide plethora of macroscopic functionalities which are extremely sensitive to the oxygen stoichiometry. Minuscule changes in the local O content may result in large macroscopic responses, both due to variations in local electronic properties and also in lattice strains. In this context, investigations of the in-situ reduction of oxide nanoparticles provide a unique tool to understand the transformations that take place when heated [25] or upon redox reactions [16,17,26], likely scenarios in many of their applications [27-30]. Unfortunately, more often than not these processes take place in a very inhomogeneous fashion within nanometric length scales. Thus, monitoring nanometric reaction fronts in real space with atomic resolution becomes a key task which may pave the way to harnessing novel heterostructured nanosystems [31-34]. Macroscopically-averaged diffraction techniques cannot always provide a clear picture of the intrinsically discontinuous nature of such systems, hindering our understanding of the mechanisms underlying the behavior of such small active regions. Real space probes capable of providing a comprehensive view of the structure, chemistry and electronic properties in real space constitute the keystone towards harnessing hybrid nanosystems. Owing to the success of the correction of optical aberrations, advanced electron microscopy and spectroscopy techniques have undergone a major revolution within the last decade and can provide atomically resolved views of complex materials at the sub-Ångström scale with single-atom sensitivity [35-37]. Armed with such tools, we are in a unique position to address the study of heterogeneous nanostructured materials in real-time.

In this work, we present an atomic resolution live (in-situ) study of electron beam-induced transformations in core/shell Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ nanoparticles by
means of high-resolution scanning transmission electron microscopy combined with electron energy-loss spectroscopy (STEM-EELS). Interestingly, the manganese oxide-iron oxide nanoparticles system is appealing not only for its unique magnetic properties (e.g., antiferromagnetic interface coupling or magnetic proximity effects) [10,38-40] but also for its applications in diverse fields like catalysis and environmental remediation [41,42], batteries and supercapacitors [43,44], microwave components [45] or biomedicine [46,47]. Our results, acquired within time scale of minutes, show that the nanoparticles under electron irradiation can develop a variety of defects, including formation of voids and dislocations, as the Mn$_3$O$_4$ component is forced to reduce to MnO. Such transformations depend on factors such as the morphology and they occur in different stages that include both redox reactions and generation of voids followed by stress relaxation, ionic transport and formation of defects.

2. Materials and Methods

2.1 Synthesis of the Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ nanoparticles

Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticles were synthesized using a seeded-growth approach [10], where pre-synthesized Fe-oxide and Mn-oxide nanoparticles were used as seeds (cores) for the growth of Mn-oxide and Fe-oxide shells, respectively. Fe$_3$O$_4$/Mn$_3$O$_4$ nanoparticles were synthesized using a slurry of dispersed iron oxide cores in dibenzyl ether heated up to 220 °C. 10 mL of a dibenzyl ether solution containing manganese (II) acetylacetonate, 1,2-hexadecanediol, and a mixture of oleylamine and oleic acid as surfactants was injected rapidly to the slurry. The temperature was maintained at 220 °C for 1 hour after the injection under magnetic stirring. Alternatively, Mn$_3$O$_4$/Fe$_3$O$_4$ nanoparticles were synthesized by dispersing manganese oxide cores in a dibenzyl ether solution of iron (III) acetylacetonate, 1,2-
hexanedicandiol, oleic acid and oleylamine. The reaction flask was heated to 250 °C for 20 minutes. The reaction vessel was removed from the heat and allowed to cool down to room temperature under argon. Both types of nanoparticles were washed by several cycles of coagulation with ethanol, centrifuging at 2000 g, disposal of supernatant, and redispersion in n-hexane.

2.2 Scanning Transmission Electron Microscopy (STEM)

STEM samples were prepared by sonicating the nanoparticles in methanol to make a suspended solution. One drop of the solution was deposited on a holey carbon TEM grid, and then the grid was air-dried. The experiments were performed on a Nion UltraSTEM200, equipped with a cold field-emission electron source and a fifth -order aberration corrector, operating at accelerating voltages of 60 kV and 200 kV. High angle annular dark field (HAADF) images were used to measure the lattice parameters of the different compounds. For this aim, a script was used to assign atomic column positions to the brightest pixel locations. Once all the coordinates of all atomic columns were obtained, distances between nearest neighbors were measured between them.

2.3 Electron energy-loss spectra (EELS)

Electron energy-loss spectra (EELS) were collected using a Gatan Enfinium spectrometer, with an energy dispersion of 0.25 eV per channel; the convergence semi-angle for the incident probe was 30 mrad, with an EELS collection semi-angle of 36 mrad. Electron beam currents were kept in the range of a few tens of pA, in a relatively high dose range estimated to be near $5 \times 10^7$ e$^-$/Å$^2$. EEL spectrum images were acquired for the different elements of interest. Mn, Fe and O maps were produced using the Mn $L_{2,3}$, Fe $L_{2,3}$, and O K edges, respectively. Spectrum images were acquired on nanoparticles that were off-axis in order to prevent channeling related artifacts during quantification. In order to carry out a meaningful quantification the background was
subtracted using a power-law fit and the intensity under every absorption edge was integrated using a 25 eV wide window. We used the built-in quantification process available in Digital Micrograph (Gatan Inc) using the available cross sections. Principal component analysis (PCA) [48] was used to improve the signal-to-noise level in the spectrum images. Multiple linear least square fits were also used to analyze the spatial distribution of the different components (see Fig. S1 in the Electronic Supplementary Material (ESM) for an example). Note that at least three different on-axis, randomly selected particles were analyzed in each sample by means of atomic resolution STEM-EELS (previously, low magnification TEM characterization was carried out to ensure sample homogeneity).

2.4 STEM-HAADF simulations

STEM-HAADF simulations were obtained by a multi-slice method using the JEMS software. The experimental parameters used for the simulation are $C_s = 0.009$ mm (spherical aberration) and $C_5 = 10$ mm (fifth-order spherical aberration), and all other axial aberration coefficients were neglected.

2.5 Geometric phase analysis (GPA)

Geometric phase analysis (GPA) was used to measure the distortion of lattice fringes with respect to a reference region in an HRTEM image of the particles [49]. A careful analysis was carried out to rule out the influence of scanning artifacts on the analysis. The lattice parameters were determined from the STEM images by averaging the distance between consecutive atoms in a given atom row (see example in Fig. S2 in the ESM). Geometric phase analysis was performed using 220 MnO/ 440 Fe$_3$O$_4$ and 002 MnO/ 004 Fe$_3$O$_4$ reflections using a cosine mask with a spatial resolution of 0.9 nm. The reference was chosen in the Fe$_3$O$_4$ unstrained region (top left region of the particle).

2.6 Particle image velocimetry (PIV)
The analyses of the STEM-HAADF images by particle image velocimetry (PIV) were carried out using the PIV plugin [50] for Image J [51]. Prior to the analysis, one hundred STEM-HAADF images (recorded every 2 seconds) were aligned to correct the drift, and a set of twenty images, obtained by summing up every five subsequent STEM-HAADF images, were used for the PIV analysis.

3. Results and Discussion

The studied core/shell Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ particles are highly crystalline and due to the topotactical growth of the spinel structures they exhibit defect free core/shell interfaces, as depicted in Fig. 1a and 1b. The morphology of the two systems is slightly different, where Fe$_3$O$_4$/Mn$_3$O$_4$ nanoparticles present an incomplete shell while Mn$_3$O$_4$/Fe$_3$O$_4$ nanoparticles exhibit a continuous, complete shell. The EELS maps shown in Fig. 1 reveal that the Fe$_3$O$_4$/Mn$_3$O$_4$ nanoparticles indeed consist of an iron oxide core with an overgrown manganese-rich shell (Fig. 1c-d and g-h). The stoichiometry of the Fe$_3$O$_4$/Mn$_3$O$_4$ core/shell nanoparticles was assessed by EELS off-axis imaging [52] both at 60 kV (Fig. 1e-f) and 200 kV (Fig. 1i-j). The false-color map displaying the relative oxygen content [53] of the Fe$_3$O$_4$/Mn$_3$O$_4$ core/shell (Fig. 1e) obtained at 60 kV reveals a fairly homogeneous stoichiometry across the nanoparticle. The laterally averaged line scan profile (Fig. 1f, where O stands for oxygen and M for the metal species) indicates relative oxygen to metal content of approximately 57 at%, in good agreement with the expected stoichiometry of the Fe$_3$O$_4$ and Mn$_3$O$_4$ phases. On the other hand, the analysis of equivalent data acquired at 200 kV (bottom panels) exhibits a severe inhomogeneity of the relative O content revealed through a significant O reduction near the surfaces (Fig. 1i). The laterally averaged profile (Fig. 1j) shows that the oxygen content decreases from ca. 57-59 at% near the central portion of the
particle down to 50 at% at the surface of the Fe$_3$O$_4$/Mn$_3$O$_4$ core/shell, suggesting that the stoichiometry of the shell is, in fact, closer to that of MnO. We conclude that at 200 kV an electron beam-induced reduction of Mn$_3$O$_4$ to MnO takes place, in agreement with earlier reports [54,55]. Interestingly, this beam-induced phenomenon also takes place in the case of the Mn$_3$O$_4$/Fe$_3$O$_4$ configuration (note the presence of MnO at the core of the nanoparticle in Fig. S1 in the ESM), when a Mn$_3$O$_4$ core is completely surrounded by Fe$_3$O$_4$ and no free Mn oxide surface is available, indicating that this is not a surface-related effect. Typically, the reduction of the spinel phase into the rock salt phase is observed at relatively high temperature [56] and pressure but it can also be triggered by an induced reduction process [57]. Note that no electron-induced reduction of Fe (III) to Fe (II) is observed in neither the Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nor the Fe$_3$O$_4$/Mn$_3$O$_4$ core/shell configurations. This finding can be understood both in terms of the difference in Gibbs free enthalpy between the Mn$_3$O$_4$ → MnO and Fe$_3$O$_4$ → FeO transformation [54] and the instability of Fe$_{1-x}$O [58] (see ESM).

The local processes underlying the reduction transformation can be monitored via atomic resolution imaging. High-resolution STEM-high angle annular dark field (STEM-HAADF) images recorded along the spinel [1-10] zone axis allow tracking in-situ electron beam-induced selective reduction of Mn$_3$O$_4$ in both Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell systems. As summarized in Fig. 2a-d (detail in e and f), a contrast change is observed in a temporal series of consecutive images, which is associated with the formation of MnO upon electron exposure. Both core/shell systems display a high crystallinity and more coherent and defect free interfaces before the transformation of the spinel-type Mn$_3$O$_4$ to a rock salt-type MnO (see Fig. 2a and e; detail in e-f). Comparing Fig. 2a and 2c allows identifying the location of the manganese oxide (right outer most edge of the particle) and iron oxide regions.
Similarly, it is possible to observe the location of the manganese oxide in the center of the particle shown in Fig. 2c-d. All of the involved phases are based on an FCC oxygen sublattice that is preserved upon reduction, permitting a high lattice coherency.

An in-depth analysis of the images of the Fe₃O₄/Mn₃O₄ nanoparticle, shown in Fig. 2a, evidences that the lattice parameter of the cubic spinel (Fd̅3m) Fe₃O₄ core lies around \( a(\text{Fe}_3\text{O}_4)_{\text{core}} = 8.4 \text{ Å} \), i.e. similar to the bulk [58] value, which gradually increases towards the nanoparticle surface. The shell shows a change of lattice parameter, in which \( \sqrt{2}a(\text{Mn}_3\text{O}_4)_{\text{shell}} = 8.5 \text{ Å} \) and \( c(\text{Mn}_3\text{O}_4)_{\text{shell}} = 9.5 \text{ Å} \), denoting an expansion of \( \Delta a(\text{Mn}_3\text{O}_4)_{\text{shell}} \approx +4\% \) (compared to the bulk [58] value, for the tetragonal spinel (I4̅1/amd) Mn₃O₄, \( a = 5.76 \text{ Å} \) and \( c = 9.44 \text{ Å} \) due to the interface with the Fe₃O₄ core (Figs. S2 and S3 in the ESM). In addition, upon the reduction of the manganese oxide shell, an overall decrease around \( \sim 2\% \) of the nanoparticle size is observed, which should be mostly related to oxygen removal. The lattice parameters obtained for the Mn oxide shell after reduction (MnO; rock salt, Fm̅3m) are \( a(\text{MnO})_{\text{shell}} = 4.6 \text{ Å} \), about 4% larger than the bulk.

In the case of the Mn₃O₄/Fe₃O₄ core/shell nanoparticle, before the transformation, the analysis of the image shown in Fig. 2c indicates that the lattice parameters of Mn₃O₄ and Fe₃O₄ lie around \( \sqrt{2}a(\text{Mn}_3\text{O}_4)_{\text{core}} = 8.5 \text{ Å} \) and \( c(\text{Mn}_3\text{O}_4)_{\text{core}} = 9.0 \text{ Å} \) and \( a(\text{Fe}_3\text{O}_4)_{\text{shell}} = 8.5 \text{ Å} \), i.e., \( \Delta a(\text{Mn}_3\text{O}_4)_{\text{core}} \approx +4\% \) (expansion) and \( \Delta c(\text{Mn}_3\text{O}_4)_{\text{core}} \approx -5\% \) (compression) and \( \Delta a(\text{Fe}_3\text{O}_4)_{\text{shell}} \approx +1\% \), with respect to the bulk. In this case, the smaller relative changes in the lattice parameters of the shell reflect its large volume fraction, which dominates over the core. In general, such comparisons to the bulk values show that both the Mn₃O₄ and Fe₃O₄ contract and/or expand to keep the system energy low by generating a coherent interface [2,60,61] in which the configuration, i.e., Mn₃O₄ in the core or in the shell, and the volume fractions of each phase are very important to
understand which phase undergoes the largest deformation. In fact, for Fe$_3$O$_4$/Mn$_3$O$_4$, with an incomplete shell, only the shell suffers strains to accommodate the coherent interface (see Fig. S4 in the ESM), whereas for Mn$_3$O$_4$/Fe$_3$O$_4$, with a fully developed core/shell structure, both counterparts need to strain their structures to stay structurally coherent. This indicates that the amount of elastic energy contained in the particles is still lower than that required to break the interface by forming an incoherent, defect rich state. Notably, these changes in their lattice parameters (i.e., strains) can ultimately lead to a large modification of their functional properties, e.g. their magnetic ground state, since the compressed (or expanded) lattice parameters effectively reduces (or enlarges) the interatomic distances. In the case of magnetic materials, a change of interatomic distances and/or angles directly influences the exchange coupling constant [62]. For example, for bulk MnO a reduction of the Mn-O bond length increases its Néel temperature [63,64], while strains can strongly influence the Verwey transition temperature of Fe$_3$O$_4$ [65,66].

The spinel and rock salt structures (e.g., Mn$_3$O$_4$ and MnO) transform topotactically [63] into one another via redistribution of cations from/to the interstitial tetrahedral positions to/from the octahedral positions while maintaining the oxide fcc-sublattice along with release/uptake of oxygen. The transformation observed in both core/shell systems (i.e. Mn$_3$O$_4$ → 3MnO + ½O$_2$) involves a volume loss associated with oxygen loss and a lattice expansion/contraction (i.e., $\Delta a$(Mn$_3$O$_4$) → MnO = +8 % and $\Delta c$(Mn$_3$O$_4$)→MnO = –6 % for bulk materials). The comparison of the electron beam-induced reduction of the Mn$_3$O$_4$ in the two systems (Fig. 2b and 2d) suggests that the underlying mechanism of reduction in the Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticles might be more complex than in the case of Fe$_3$O$_4$/Mn$_3$O$_4$. When the Mn$_3$O$_4$ composes the shell, its reduction towards MnO modifies the lattice parameters as the reduction of Mn(III) to Mn(II) removes the
Jahn-Teller distortion and increases the cation radius [67-70]. The associated large changes of lattice parameters (i.e., $\Delta a({\text{Mn}_3{\text{O}}_4} \rightarrow \text{MnO} \approx +12\%$ and $\Delta c({\text{Mn}_3{\text{O}}_4} \rightarrow \text{MnO} \approx -3\%$) are easily accommodated by the absence of confinement which allows lattice rotation that maintains a defect-free coherent interface (see Fig. 2a-b and Fig. S4 in the ESM). Moreover, the lattice expansion/contraction evolves as expected during the transformation (compare the expansion/contraction sign of the lattice parameters with bulk materials). Conversely, in the case of the $\text{Mn}_3{\text{O}}_4/\text{Fe}_3{\text{O}}_4$ core/shell system, the $\text{Mn}_3{\text{O}}_4$ reduction comprises more intricate effects such as the concomitant formation of a void as the $\text{Mn}_3{\text{O}}_4$ core transforms into MnO (white circle in Fig. 2d) and the lattice expansion along the $c$-axis. Fig. 2e and 2f show a zoomed-in region of the interface at the core-shell before and after reduction, respectively. Since some Mn/Fe interdiffusion can take place during the transformation, it was allowed within the simulations to include a possible chemical intermixing, although no major effect was found. A comparison of the real image with the simulations indicate that the interface is relatively sharp on the order of one unit cell. Considering that the reduction of $\text{Mn}_3{\text{O}}_4$ to MnO in the $\text{Mn}_3{\text{O}}_4/\text{Fe}_3{\text{O}}_4$ case is more complex than for the $\text{Fe}_3{\text{O}}_4/\text{Mn}_3{\text{O}}_4$ case, we will study in more depth the evolution of the $\text{Mn}_3{\text{O}}_4/\text{Fe}_3{\text{O}}_4$ nanoparticles.

In order to monitor the atomic column movement associated with the reaction front in the $\text{Mn}_3{\text{O}}_4/\text{Fe}_3{\text{O}}_4$ nanoparticles, we used particle image velocimetry (PIV, see Fig. S5 in the ESM) [71,72] in a series of subsequent high-resolution STEM-HAADF images at early transformation times. Remarkably, the PIV analysis (see Fig. 3, S5 and Video V1 in the ESM) reveals an inhomogeneous pattern of column displacements (for our analysis we are assuming that the small thickness of the particles does not constitute an issue). The larger shifts are mainly localized at the top-left region of the particle shell towards the nanoparticle core, with a maximum displacement value of 1.5 Å (see Fig.
S5 in the ESM). The relatively small displacements observed at the right bottom corner are probably due to the pinning of the particle by other neighboring nanoparticles. The column displacement towards the core occurs as a strain relaxation mechanism related to oxygen loss during reduction of Mn₃O₄ to MnO. According to the PIV analysis, some cation columns move around the network during the beam-induced transformation. However, confinement effects within the MnO core in the Mn₃O₄/Fe₃O₄ core/shell nanoparticle prevent stress relief mechanisms via lattice rotation from taking place, a mechanism which is available for the Fe₃O₄/Mn₃O₄ configuration (see Fig. S4 in the ESM). As a consequence, misfit dislocations appear, as evidenced in Fig. 4 (see below). Surprisingly, in the Mn₃O₄/Fe₃O₄ core/shell nanoparticle the reduction of the Mn₃O₄ spinel structure brings about a global expansion of the lattice parameters along the c-axis of the core, (i.e., Δc Mn₃O₄ → MnO ≈ +4 %, see Fig. S2 in the ESM). The unexpected lattice expansion of the core along the c-axis also compensates for the oxygen loss to maintain the coherent interface between the core and the shell of the nanoparticle. The reduction of Mn(III) to Mn(II) also results in an expected expansion of the a-axis within the core of about Δa (Mn₃O₄ → MnO) ≈ +5 %. The expansion of the a-axis in the core results in the expansion of the a-axis of the shell as well.

Geometric phase analysis (GPA) [49] can be used for further analysis of the strains and defects resulting from the loss of oxygen and redistribution of cationic columns. Fig. 4 shows a quantification of the strain along the c-axis and along the [110] direction. The GPA analysis, using Fourier filtering, highlights the presence of defects in the Mn₃O₄/Fe₃O₄ core/shell nanoparticle upon electron exposure. The magnified STEM-HAADF images of the Mn₃O₄/Fe₃O₄ core/shell interface reveal structural changes upon electron exposure (Fig. 4a-d). The coherent spinel domain transforms into a metastable MnO-like phase (Fig. 4b), followed by a cation redistribution across the core/shell
We believe that upon prolonged exposure time the formation of an intermediate \( \text{Mn}_x\text{Fe}_{3-x}\text{O}_4 \) phase is possible (Fig. 4d and also the detailed interface in Fig. 2f) although a definitive conclusion in this respect cannot be drawn from our data. The formation of this intermediate phase would be driven by an uneven cation distribution over the whole core/shell nanoparticle combined with a vacancy gradient between the inner and outer boundaries [73]. In fact, the deformation map shown in Fig. 4e shows that the core of the nanoparticle at early stages of exposure is positively strained (3-12\%) along the \( c \)-axis with respect to the \( \text{Fe}_3\text{O}_4 \) shell (the down-right region of the nanoparticle shell is used as reference area for GPA), pointing towards the heterogeneous distribution of Mn and Fe cations [69]. Note that an absence of observable dislocations (from this particular zone axis) at the core/shell interface is observed in Fig. 4e-h (note that dislocations would be noticeable as discontinuities, i.e., sudden color change from yellow to blue, on the deformation map). This fact indicates that the nanoparticle core/shell boundary acts as a strained buffer layer of intermediate metastable cation arrangement in between the normal and inverse spinel structure, as previously reported on similar \( \gamma \)-\( \text{Fe}_2\text{O}_3/\text{Mn}_3\text{O}_4 \) core/shell nanoparticles [74,75]. The deformation maps shown in Fig. 4f-h reveal a positively strained core along with the presence of interfacial defects. The dislocations act as a strain relief at the core/shell nanoparticle interface allowing the nanoparticle to maintain a stable interface since the confinement of the core does not allow lattice rotation. These defects are misfit dislocations that are characterized by a shift of \( \frac{1}{4} \) along [001] with respect to the \( \text{Fe}_3\text{O}_4 \) lattice, or alternatively by a shift of \( \frac{1}{2} \) [001] with respect to the \( \text{MnO} \) lattice, as evidenced by white circles in panel j. The misfit dislocation pairs, in turn, propagate around the core/shell interface but do not annihilate (see Fig. 4k,l). The deformation maps along [110] indicate a relatively homogeneous strain throughout the nanoparticle
with some small local interfacial strain (see Fig. 4m-p). This finding indicates that the 
Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticle is anisotropically strained, so possible strain 
relaxation mechanisms involving cation redistribution along the $c$-axis towards the 
interface can be expected upon annealing (external or under the electron beam). Also, 
the core of the nanoparticle remains in a relatively highly strained state (3-12%) as the 
volume reduction associated with oxygen loss contributes to increasing the strain at the 
nanoparticle interface. Defects such as misfit dislocations are observed (Fig. 4q-t). The 
formation of MnO implies a large displacement of cations from tetrahedral into 
octahedral positions. This fact, together with the presence of newly generated Mn(II) 
ions have been reported in similar cases before, is again likely to promote the formation 
of interfacial Mn$_3$Fe$_{3-x}$O$_4$ [3,74,75].

4 Conclusion

In summary, we have demonstrated that a selective electron-induced structural 
transformation of Mn$_3$O$_4$ into MnO in Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell 
nanoparticle systems can take place through different mechanisms depending on the 
core/shell configuration. The confinement of the Mn$_3$O$_4$ in Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell 
nanoparticle prevents stress relief by lattice rotation and results in the formation of 
misfit dislocation pairs. Prolonged exposure of the particles, may ultimately lead to 
annealing of the core/shell nanoparticles leading to an equilibrium single phase. We 
have shown how in-situ STEM-HAADF monitoring of the atomic column positions and 
image analysis combining GPA and PIV are useful tools to study chemical 
transformations and diffusion processes in solid materials as they allow direct 
observation of the relationship between the structure and strain relaxation during the 
Mn$_3$O$_4$ reduction. Such electron beam-induced transformations can be tuned by varying
the acceleration voltage, which can be exploited to further modulate the energy transfer in a controlled and reproducible manner. The selective reduction of confined heterostructured materials based on functional oxides (e.g., core/shell nanoparticles) systems offers innovative possibilities for locally tuning or patterning their electronic, magnetic and other functional properties (e.g., Mn$_3$O$_4$ is ferrimagnetic while MnO is antiferromagnetic) in order to optimize functionality in fields as diverse as catalysis, magnetism, biomedicine, or sensors.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at:

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Fig. 1. High Resolution HAADF images of both types of core|shell nanoparticles. (a) Fe₃O₄/Mn₃O₄ and (b) Mn₃O₄/Fe₃O₄. Elemental mapping of two different Fe₃O₄/Mn₃O₄ core/shell nanoparticles at (c-f) 60 kV and (g-i) 200 kV. EELS maps obtained from the Fe L₂,₃ edge at (c) 60 kV and (g) 200 kV. Mn L₂,₃ edge at (d) 60 kV and (h) 200 kV. Maps of the relative oxygen content at (e) 60 kV and (i) 200 kV. Data along the direction outlined with a red arrow in (e) and (i), respectively, are shown in panels (f) and (j).
Fig. 2. Electron-induced transformation from Mn$_3$O$_4$ to MnO in Fe$_3$O$_4$/Mn$_3$O$_4$ and Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticles. (a-b) High-resolution STEM-HAADF images recorded along the [1-10] Fe$_3$O$_4$ axis of a Fe$_3$O$_4$/Mn$_3$O$_4$ core/shell nanoparticle (a) at an early stage and (b) after 10 minutes of electron beam exposure. (c-d) High-resolution STEM-HAADF images recorded along the [1-10] Fe$_3$O$_4$ axis of a Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticle (c) at an early stage and (d) after 30 minutes of beam exposure. The white circled area in (d) indicates the position of a void. (e-f) Magnified STEM-HAADF images of the highlighted (orange box) regions in (c) and (d), respectively, with the unit cells overlaid (oxide anions not shown). (Insets) Multislice simulation of a 12-nm thick Mn$_{3-x}$Fe$_x$O$_4$ spinel and a MnO rock salt-type structures along [1-10].
**Fig. 3.** Atomic column displacement analysis during phase transformation of the Mn$_3$O$_4$ core into MnO. (a) High-resolution STEM-HAADF image of the Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticle at the initial electron exposure. (b) Magnified STEM-HAADF image of the boxed area in image (a). (c) Displacement map obtained by PIV analysis of the STEM-HAADF images in (b) and (d). The false color scale indicates the atomic column displacement in Å. (d) Magnified STEM-HAADF image of the boxed area in image (e). (e) High-resolution STEM-HAADF image of the Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell nanoparticle after 3.5 minutes of electron exposure.
**Fig. 4.** Lattice deformation upon electron exposure in the Mn$_3$O$_4$/Fe$_3$O$_4$ core/shell system. (a-d) High-resolution STEM-HAADF images of the nanoparticle interface (area marked with a square in the deformation maps) at (a) initial exposure time, (b) after 30, (c) 45, and (d) 48 minutes respectively. (e-h) Deformation maps obtained by GPA using 002 MnO/004 Fe$_3$O$_4$ reflections. (i-l) Fourier-filtered images of (a-d) using 002 MnO/004 Fe$_3$O$_4$ reflections, respectively. (m-p) Deformation maps obtained by GPA using 220 MnO/440 Fe$_3$O$_4$ reflections. (q-t) Fourier-filtered images of images in (a-d) using 220 MnO/440 Fe$_3$O$_4$ reflections. The false color scale depicts the strain in %. White circles highlight misfit dislocation in the Fourier-filtered images. Scale bars represent 1 nm.
Probing the Meta-Stability of Oxide Core/Shell Nanoparticle Systems at Atomic Resolution

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In-situ electron-induced selective reduction of magnetic oxides is studied in core/shell nanoparticles using high-resolution electron microscopy and spectroscopy. This study highlights the relevance of understanding the local dynamics responsible for nanoscale changes to control stability and, ultimately, macroscopic functionality.

ToC figure:
Supplementary Information

Probing the Meta-Stability of Oxide Core/Shell Nanoparticle Systems at Atomic Resolution

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Figure S1: Multiple linear least squares (MLLS) coefficient maps of a Mn$_3$O$_4$/Fe$_3$O$_4$ nanoparticle, obtained at 200kV, for the different reference spectra: (a) Fe$_3$O$_4$, (b) Mn$_3$O$_4$ and (c) MnO, respectively. (d) Corresponding RGB colored map where the previous maps are overlaid with Fe$_3$O$_4$ in red, MnO in green, and Mn$_3$O$_4$ in blue. Image lateral size is 12nm x 12 nm.
Figure S2. Lattice spacing in Mn$_3$O$_4$|Fe$_3$O$_4$ core|shell nanoparticle at different electron exposure times. (a, b) STEM-HAADF high resolution images of a Mn$_3$O$_4$|Fe$_3$O$_4$ core|shell nanoparticle recorded along <110> at initial exposure stage and after 30 minutes, respectively. (c, d) Line profiles of the marked regions in (a) along (c) [110] and (d) [001]. (e, f) Line profiles of the marked regions in (b) along (e) [110] and (f) [001]. Scale bar: 2nm.
Figure S3: Theoretical values and model for Mn$_3$O$_4$, MnO and Mn$_x$Fe$_{3-x}$O$_4$. Spinel and rock-salt structures (top panel), with their respective simulated electron diffraction patterns (bottom panel).
Figure S4. GPA analysis for a Mn$_3$O$_4$/Fe$_3$O$_4$ nanoparticle. (a) High-resolution STEM-HAADF image of the nanoparticle interface at initial exposure time. (b) Fourier filtered image of (a) using the 044$_{Fe3O4}$|022$_{MnO}$ reflection. (c) Deformation map obtained by GPA using the 044$_{Fe3O4}$|022$_{MnO}$ reflection. (d) Model of the coherent domain corresponding to a spinel structure of Mn$_x$Fe$_3$O$_4$. (e) Fourier filtered images of (a) using the 004$_{Fe3O4}$|002$_{MnO}$ reflection. (f) Deformation map obtained by GPA using the 004$_{Fe3O4}$|002$_{MnO}$ reflection. (g) High-resolution STEM-HAADF image of the nanoparticle interface after 15 minutes exposure time. (h) Fourier filtered image of (g) using the 044$_{Fe3O4}$|022$_{MnO}$ reflections. (i) Deformation map obtained by GPA using the 044$_{Fe3O4}$|022$_{MnO}$ reflection. (j) Model of Fe$_3$O$_4$MnO domains with a 5° lattice rotation. (k) Fourier filtered images of (g) using the 004$_{Fe3O4}$|002$_{MnO}$ reflection. (l) Deformation map obtained by GPA from (k) using the 004$_{Fe3O4}$|002$_{MnO}$. The false color scale depicts the strain in %. The dashed lines inset depict the interface between Fe$_3$O$_4$ and Mn$_3$O$_4$. Scale bars: 2 nm.
Figure S5. Series of subsequent high-resolution STEM-HAADF images extracted from the PIV study. The arrows in the top left panel show the crystallographic directions of the Mn₃O₄ structure. The scale bar represents 2 nm.

Also, Video 1 (V1) including the PIV images is attached as a separate file as additional supporting information.
Thermodynamics of the Fe\(_3\)O\(_4\) and Mn\(_3\)O\(_4\) phases

The electron-induced reduction of Fe(III) to Fe(II) is not observed in neither of the core|shell systems. This finding can be understood both in terms of the difference in Gibbs free enthalpy between the Mn\(_3\)O\(_4\)→MnO and Fe\(_3\)O\(_4\)→FeO transformation and the instability of Fe\(_{1-x}\)O.

Table S1. Bibliography Enthalpy data\(^1\):

| Oxide   | \(\Delta H_f^o\) (KJ/mol) |
|---------|-----------------|
| Fe\(_3\)O\(_4\) | -1118.4         |
| FeO     | -272.0          |
| MnO     | -385.2          |
| Mn\(_3\)O\(_4\) | -1387.8         |

Reactions involved in Fe\(_3\)O\(_4\)|Mn\(_3\)O\(_4\) and Mn\(_3\)O\(_4\)|Fe\(_3\)O\(_4\) core|shell nanoparticles:

- Fe\(_3\)O\(_4\)→3FeO + \(\frac{1}{2}\) O\(_2\)
  \[\Delta H_r = 3 \ (-272.0) - (-1118.4) = 302.4 \text{ KJ/mol} \text{ Fe}_3\text{O}_4 \rightarrow 3.1 \text{ eV} \quad (1)\]

- Mn\(_3\)O\(_4\)→3MnO + \(\frac{1}{2}\) O\(_2\)
  \[\Delta H_r = 3 \ (-385.2) - (-1387.8) = 232.2 \text{ KJ/mol} \text{ Mn}_3\text{O}_4 \rightarrow 2.4 \text{ eV} \quad (2)\]

[1] Data from Chase, M. W., NIST-JANAF Thermochemical Tables, Fourth Edition J. Phys. Chem. Ref. Data Monograph 9, 1998.