

Structural insights into magnetic clusters grown inside virus capsids

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Abstract

Magnetic nanoparticles have multiple applications in materials science. In particular, virus capsids have been suggested as promising templates for building up nanometric-sized magnetic clusters by taking advantage of their inner cavity as a nanoreactor. In this study we investigate the magnetization of individual cobalt-filled cowpea mosaic virus empty virus-like particles using atomic force microscopy. We also combine the analysis of the effects of dehydration on the structure of virus particles with a comparison of their magnetic signal to that provided by commercially available magnetic nanoparticles of similar size. These two approaches allow the evaluation of the structure of the metallic cluster grown inside the virus capsid. We conclude that, rather than forming solid clusters, cobalt inside viruses forms a discontinuous structure which does not completely fill the viral cavity and reaches about 10% of its volume.

Introduction

Virus capsids, and other protein cages, are increasingly being used as multifunctional nanocontainers for applications in materials synthesis, drug encapsulation and delivery, and catalysis¹⁻⁷. Virus-like particles (VLPs) resemble viruses but are non-infectious as they lack the viral genetic material. Their monodispersity, morphological and structural uniformity, biocompatibility, safety, amenability to chemical and genetic modifications, and their stability over a wide range of conditions makes them ideal nanocontainers. Cowpea mosaic virus (CPMV) empty VLPs (eVLPs) can be produced in large quantities in plants using an expression system known as pEAQ vector^{8,9}. Previously, we have reported the loading of the CPMV eVLPs with cobalt and iron oxide¹⁰. In order to move closer toward the development of CPMV eVLPs for such applications as magnetic hyperthermia treatments,^{11,12} it is essential to investigate the magnetic moment of individual particles because they are required to be easily manipulated by magnetic fields. Their magnetic properties are related, in turn, with the amount of metal accumulated in the eVLP cavity.

The cavity of icosahedral virus capsid architectures is an ideal size-constrained environment where the interior surface can direct the nucleation of nanomaterials. In particular, the positively charged interior surface of cowpea chlorotic mottle virus (CCMV), which has an interior cavity diameter of approximately 24 nm, has been used for directing mineralization reactions to form spatially constrained nanoparticles such as of β -TiO₂.⁵ In addition, the interior surface of the CCMV capsid can be altered to change its charge from positive to negative, thereby allowing the mineral nucleation of transition metal oxides such as γ -FeO(OH)². This internal mineralization takes advantage of the interaction between cationic ions and the inner cavity of the virus particles, which acts as a nucleation surface for metal formation. Besides, the virus capsid does not require any further modification. In other words, a gradual growth of the nanoparticle occurs simultaneously to the entry of the precursors within the inner cavity of the virus particles. This approach results in completely filled virus capsids.²

Recently, the related virus CPMV has been developed extensively as a template for external mineralization, to produce narrowly disperse metallic nanoparticles, and CPMV eVLP for metallo-nanoparticle growth within the cavity.^{4,10} CPMV and CPMV eVLPs can be readily isolated from plants in high yield. Metallic nanoparticles can be grown inside CPMV eVLPs capsids by incubation with a solution containing a metal salt and, once the precursor is inside the virus, adding a reducing agent that induces the deposition of metal within the capsid¹⁰. In this case, the virus is used as a nanoreactor and the deposition reaction is not started until the virus is completely loaded with the corresponding precursor. Specifically, this procedure has used cobalt chloride as the metal precursor and sodium borohydride as a reducing agent. Electron microscopy (EM) imaging provides a comparison between eVLPs and loaded viruses (Co-VLPs) by the different contrast inside the particles; unmodified particles were invisible under the

transmission electron microscopy (TEM) ¹⁰. The TEM data suggested that the virus cages are completely filled with metallic cobalt (Figure 1).

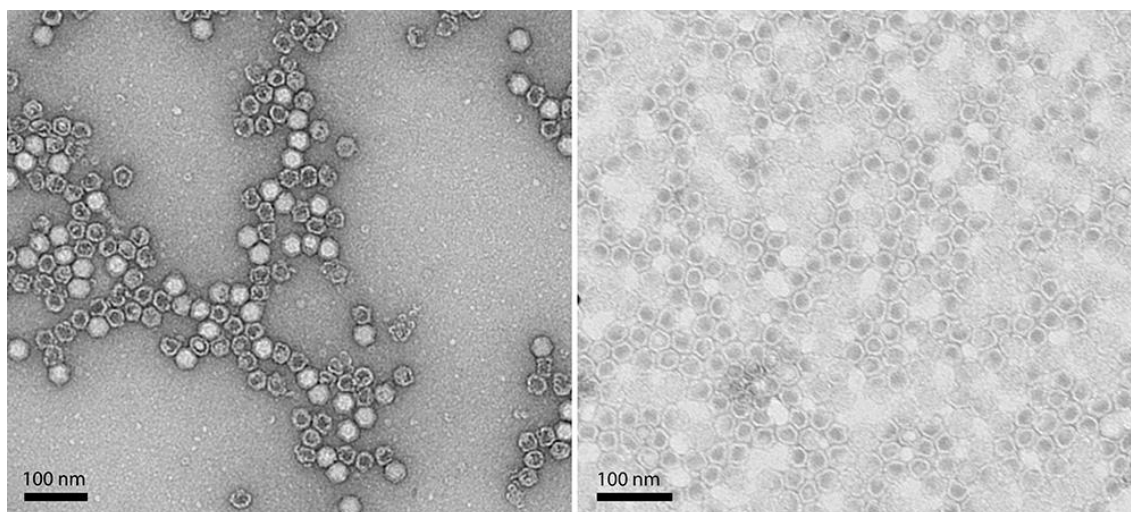


Figure 1. TEM images of CPMV particles. Left, 2% uranyl acetate stained particles of CPMV eVLPs and right, unstained Co-VLPs showing the cobalt as a core and a white ring around it representing the virus coat protein.

These observations inform about the presence of metal inside the virus but not about its structure. However, current chemical understanding of such systems indicates that one incubation cycle does not necessarily result in a high enough concentration of the precursors to completely fill the virus capsid.

Single molecule techniques provide detailed information not only on the structure, but also about the key physical properties of nanoparticles. Specifically, atomic force microscopy (AFM), in contrast to EM, gives information about the 3D topographies of virus particles in air and liquid environments ¹³. In addition, AFM provides information about a variety of physical properties, which may reveal crucial information about the internal structure of viruses ¹⁴. The ultimate proof of functional magnetic particles would be detection of the magnetic signal from individual entities. Although, magnetic force microscopy (MFM) has been suggested as a convenient tool for measuring the magnetism of single particles ¹⁵⁻¹⁹, a correct interpretation of the MFM images of low-coercivity and low moment magnetic nanoparticles (MNPs) remains elusive ²⁰

This paper focuses on the structural and magnetic characterization of individual cobalt-loaded CPMV eVLPs (Co-VLPs) and provides new insights on the metallic nanostructure formed within the eVLP.

Materials and Methods

Empty and loaded virus particles.

Empty CPMV virus particles were propagated and isolated from plants, purified and loaded with cobalt by published procedures ^{9,10}. The eVLPs were treated with chymotrypsin to enzymatically

cleave off the C-terminal 24-amino acid peptide prior to loading to ensure homogeneity of the particles and so as to maximize internal loading²¹.

Procedures for the adsorption of particles on surfaces

The virus particles were stored in a solution of phosphate-buffered saline (PBS) 100 mM, pH 7.0. To prepare AFM samples, 30 μ L were placed in highly ordered pyrolytic graphite (HOPG), to provide a smooth surface for the AFM studies and left for 30 minutes for the particles to settle on the surface. Thereafter, the samples were washed with Milli-Q water and dried with argon for AFM and MFM characterization. For AFM measurements in liquid solutions, samples were washed in the same buffer (10 mM sodium phosphate buffer pH 7.2) in which the viruses were diluted in order to avoid the sample drying out and to remove the unattached particles.

To obtain a homogeneous dispersion of cobalt nanoparticles (CoNPs), 1 mg of carbon-coated cobalt nanoparticles (Sigma Aldrich) were suspended in 1 mL of Milli-Q water containing sodium dodecyl sulfate (SDS, Sigma Aldrich) 0.01%. These suspensions were sonicated for 2 hours (frequency: 24 kHz, amplitude: 210 microns) on a UP400S Hielscher ultrasonic tip, Model H3-Tip3 (3-mm diameter). From the resulting colloidal suspension 40 μ L were deposited by drop-casting on a mica surface functionalized with 3-aminopropyltriethoxysilane (APTS, Sigma Aldrich). After one hour, the samples were dried under argon, washed with Milli-Q water and dried again under argon.

Magnetic Force Microscopy (MFM)

The magnetic characterization was performed with a commercial MFM from Nanotec Electronica S.L. (Madrid, Spain). A magnetic probe attached to the end of a cantilever was used as the force sensor. Commercial Nanosensors PPP-MFMR and Budget Sensors MFM ($k=1.5$ N/m and $f = 75$ KHz) were used. Before each experiment, the probes were magnetized along their pyramid axis to ensure a maximum interaction between the tip and the sample. To measure a magnetic signal with MFM dynamic modes were employed (Amplitude Modulation in the case of ambient conditions, Drive Amplitude Modulation at High Vacuum)²¹. The interaction between the magnetic moments of the tip and the sample, F_z , induces a change in the resonance frequency of the cantilever with the distance z . This frequency shift ($\Delta\omega$) can be correlated with the force gradient as is shown in equation (1):

$$\Delta\omega = \frac{\omega_0}{2k} \frac{\partial F_z}{\partial z} \quad (1)$$

Where ω_0 is the resonance frequency and k is the force constant of the cantilever.

The isolation of electrostatic forces, which are simultaneously present to magnetic forces, is achieved with Kelvin Probe Force Microscopy as explained elsewhere²² When very low magnetic signals are measured the presence of other long range interactions, like electrostatic, can hide the magnetic information. We utilized HOPG (a conductor) for Co-CPMV to induce the charge migration and isolate the magnetic signal from the electrostatic one. In the case of cobalt

nanoparticles the magnetic signal was large enough so to combine MFM + KPFM HOPG was not needed.

Thermogravimetric analysis (TGA)

Experiments were conducted using a Perkin Elmer Diamond TG/DTA instrument between room temperature and 900 °C at a heating rate of 100 °C min⁻¹ under an air flow of 200 mL min⁻¹. Zero grade air was supplied from BOC gases. The sample was loaded to a clean pan supported by a precision balance and the mass of the sample was monitored during the experiment. The purified Co-VLPs were either freeze-dried (-50 °C under vacuum and pressure) prior to TGA analysis or the Co-VLPs were loaded to the pan and the temperature raised to 100 °C and held at that temperature for 15 minutes to ensure the removal of all the moisture before continuing the experiment.

Results

To determine the capability of MFM to detect the magnetism of the cobalt within a single Co-VLP, a control experiment was performed on a similar-sized commercial carbon-coated cobalt nanoparticle (CoNP) which possesses high magnetization values at room temperature (160 emu/g). CoNPs were immobilized on a mica surface. The CoNPs size distribution was determined experimentally to be 19 ± 4 nm (average ± standard deviation, 50 particles), in agreement with the manufacturer's data. Figure 2 (a & b) presents simultaneous topography and magnetic images of CoNPs, respectively, obtained at a retrace distance of 20 nm in air conditions.

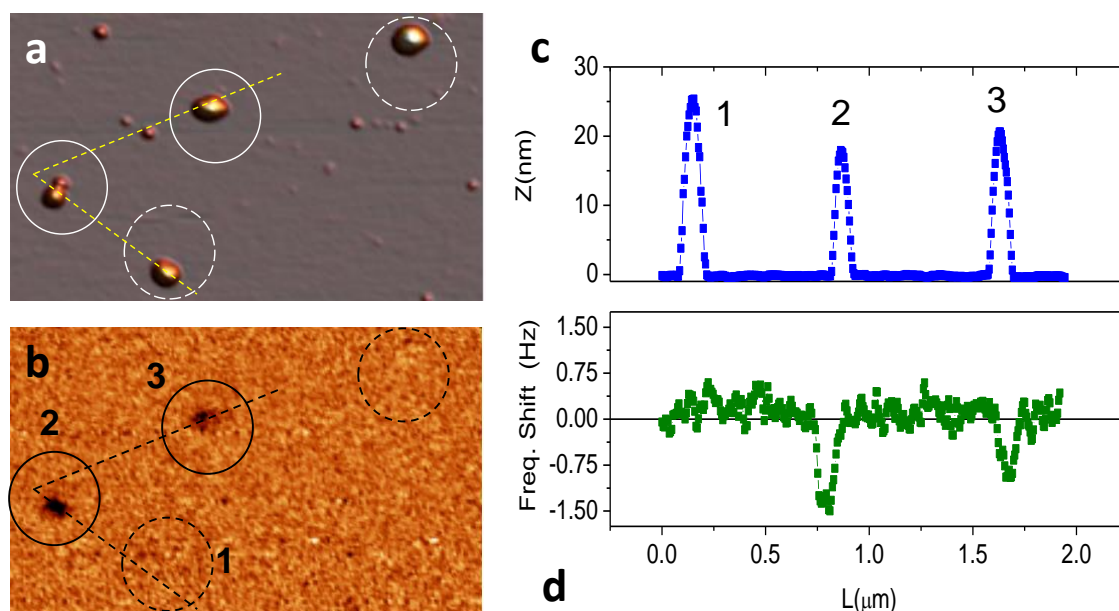


Figure 2. Magnetic characterization of commercial CoNPs particles. a) The topography of CoNPs, b) the frequency shift caused by the tip-particle magnetic interaction, c) and d) the profiles of topography and magnetic interaction, respectively, along the corresponding lines 1-2-3. Images dimension 1.9 μm × 1.4 μm.

In the case of protein shells, both empty and Co-filled virus particles (Co-VLPs) were explored by immobilizing them on HOPG and SiO₂ surfaces. Protein shells are severely affected by dehydration. In particular, virus structures are prone to collapse wall-to-wall because of the capillary forces of water menisci during the last stages of desiccation processes¹³. This effect is modulated by the presence of the genetic material within wild-type particles, which prevents the wall-to-wall collapse. In order to unveil the effects of desiccation, AFM topographies for eVLPs and Co-VLPs adsorbed on HOPG were obtained in both water and air as shown in Figure 3 (a & b), respectively. Figure 3c compares typical topographical profiles of viruses obtained in water (left) and air (right). The AFM data in water revealed that the height of eVLPs and Co-VLPs are similar between them and 4 nm below their nominal expected diameter²³, as expected. Viral particles typically deform upon absorption on surfaces because of their interaction with the surface¹⁴.

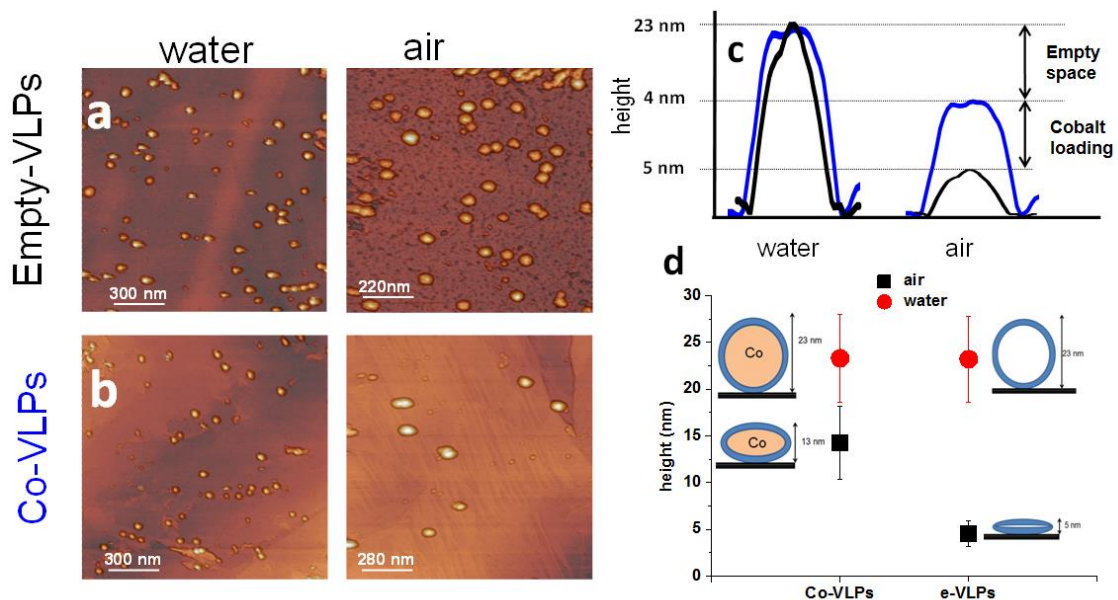


Figure 3. AFM images of desiccation effects on CPMV particles. a) eVLPs on the surface in water (left) and air (right). b) Co-VLPs on the surface, in water (left) and air (right). c) Topographical profiles of single eVLPs and Co-VLPs, in black and blue, respectively. d) Average height of eVLPs and Co-VLPs in air (black) and water (red). The number of eVLPs in water and air are 46 and 54, respectively. The number of Co-VLPs in water and air are 34 and 40, respectively. Error bars refer to the standard deviation. Insets depict cartoons of the role of the filling on the height of the particle.

In addition, although desiccated Co-VLPs have a lower height than eVLPs in liquid, they have a greater height than dehydrated eVLPs. Indeed, a detailed statistical analysis (Figure 3d) indicates that eVLPs collapse wall-to-wall to 5 nm in height when desiccated, while Co-VLPs retain a height of about 14 nm. In the case of using SiO₂, and due to weak particle-surface interactions, both eVLPs and Co-VLPs show larger heights (less collapse), although still Co-VLPs are bigger than eVLPs (Supplemental Information -SI-). Thus the adsorption force of particles on HOPG is higher

than when SiO₂ is used. As a consequence, eVLPs do not collapse wall-to-wall in SiO₂ and Co-VLPs' height is less sensitive to the amount of Co mass than when adsorbed in HOPG. Hence, HOPG data was utilized for analysis.

In general, magnetic nanoparticles present a low magnetic moment at room temperature and heterogeneous electrostatic interactions with the tip²⁰, impairing a direct interpretation of these experiments. MFM and Kelvin probe force microscopy (KPFM) experiments were performed in air and inside a high vacuum (HV) chamber, magnetic signal only being observed when measuring in HV. The principal advantage of measuring at HV is the enhancement of the sensitivity due to the increment of the quality factor of the resonance²⁴ (we have increased the signal to noise ratio by a factor of ~6). The topography and magnetic data obtained in HV for Co-VLPs are presented in Figure 4 (a & b). Figure 4c shows the corresponding profiles that allow an adequate identification of each particle with its magnetization.

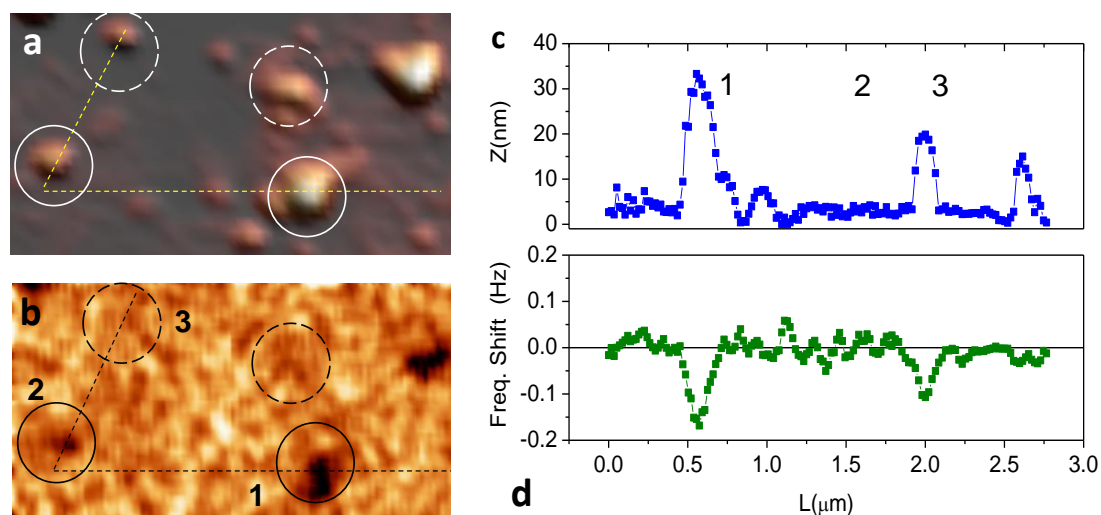


Figure 4. Magnetic characterization of Co-VLPs. a) The topography of Co-VLPs, b) the frequency shift caused by the tip-particle magnetic interaction, c), and d) the profiles of topography and magnetic signal, respectively, along the corresponding lines 1-2-3. Images dimension 1.9 μm × 1.4 μm.

We also utilized a bulk technique to estimate the nature of Co-VLPs to analyze the amount of cobalt inside viral particles. Thermogravimetric analysis (TGA) (Figure 5) determined for Co-VLPs that there was up to 55% mass loss from the starting material. If this mass loss is entirely attributed to the oxidative degradation of the protein shell, the remaining mass (45%) may be ascribed to the remaining metallic cobalt.

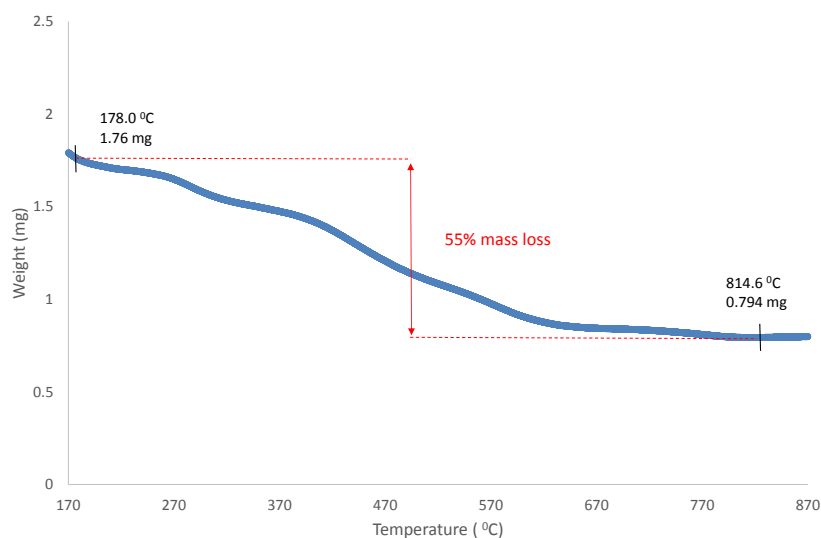


Figure 5. TGA curve of Co-VLPs lyophilized. The initial weight loss of samples observed around 100 °C is due to loss of moisture. By increasing the temperature, the weight residue declined at around 300 °C, probably due to the breakdown of side chain groups of amino acid residues of the virus capsid as well as the cleavage of peptide bonds.

Discussion

When CPMV eVLPs were incubated with cobalt ions, the cobalt cations interact with the negatively charged amino acids within the virus capsid. The subsequent addition of the reducing agent (NaBH₄) generated solid metallic cobalt within the virus capsid. TEM suggested that the cavity of CPMV was completely filled with cobalt (Figure 1)¹⁰. Our work aimed to interrogate the quality of the VLPs from a practical point of view, i.e., how large is the magnetic moment of a single Co-VLP? The magnitude of this magnetic dipole would determine the capacity of these particles to be manipulated by magnetic fields. The magnetic dipole is related to the amount of Co inside the particle. Therefore, the primary objective was to evaluate the magnetic dipole of individual particles to establish a relationship with the amount of Co. The measurement of the magnetization of Co-VLPs at the single-particle level is attained with MFM²⁵⁻²⁷.

It is well known that AFM can be used to detect different short, medium and long range interactions. However, the unknown contribution of every kind of force to the total force measured hinders the amount of quantitative information that can be obtained from the measurements²⁸. It should be noted that all the tip-sample interactions induce changes in the total force, i.e., they modify the cantilever state. In MFM it is a usual procedure to separate the topography and the magnetic signal by scanning at a certain height such that the long range tip-sample interactions dominate (typically 25-50 nm of retrace distance). An additional problem appears if several different long range interactions are present between the tip and sample. In such a case the best option is a combination of KPFM^{22, 29} and MFM to compensate the electrostatic contribution to the frequency shift signal^{28, 30}.

Most MFM measurements are based on the overall magnetic signal generated from the particles regardless to the possible variation between individual particles³¹ Besides, single particle

experiments can determine the uniqueness of each entity and the uniformity or the lack of it between particles within the same sample. MFM characterization of individual Co-VLPs requires the selection of differential contrast experiments where the positive results should not be influenced by other variables such as topography, mechanical properties or electrostatic properties. However, magnetic measurements of the commercial CoNPs revealed such inhomogeneous magnetization. This can be observed in the corresponding profiles of two particles with similar size as shown in Figure 2 (c & d). Commercial CoNPs are passivized (coated) with a carbon layer that would avoid cobalt oxidation; any damage to this coating layer would result in cobalt oxidation, which will give CoNPs with variable magnetization uncorrelated with their size.

Since Co-VLPs and commercial CoNPs are of similar size, we expected to find similar magnetization values in the case of solid filling of Co-VLPs. However, the magnetic signal of Co-VLPs was very low when measured in ambient air. Therefore, high vacuum conditions were needed to increase the sensitivity of the magnetic measurements. Figure 4c reveals that the magnetization of Co-VLPs is not correlated with their size, thus showing that the amount of Co inside the viruses varies between particles. Furthermore, the Co-VLP signal is ~8 times lower than that of the commercial CoNP, indicating that the amount of the magnetic Co inside virus particles is less than expected. Thus, we compared the Co-VLPs' signal with commercial Co-NPs of similar size, where it is known that Co spans the whole volume of the particle. We expected to find the Co-VLPs frequency shift comparable to the commercial particles. Since Co-VLPs show a 8-fold less frequency shift (~0.2 Hz) than Co-NPs (~1.5 Hz), it suggests that Co is not filling the available volume of the CPMV capsid. By attending to these data, we could roughly affirm that Co-VLPs have about 8 times less Co than commercial Co-NPs. Nevertheless, we pursued further AFM topography analyses in order to estimate the amount of Co inside the CPMV cavity. CPMV eVLPs and Co-VLPs show heights close to 23 nm in water (Figure 2), which coincides with the virus nominal size²³. However, in air conditions, AFM topographies revealed a dramatic decrease of eVLP height to ~5 nm on desiccation. In contrast, Co-VLPs height decreased to 14 nm in height after desiccation. In a similar case, it is known that DNA prevents partially the wall-to-wall collapse of bacteriophage ϕ 29 virus cages under desiccation (11). To what extent this collapse is prevented in Co-VLPs may signify the degree of filling of the virus cavity. The difference between heights observed for collapsed Co-VLP and eVLP (~9 nm) establishes a top limit for the size of the cobalt cluster inside the viruses.

There are several possible explanations for the lack of material inside the Co-VLPs. First, cobalt could escape from the particles by the release of Co^{2+} cations upon oxidation, which would be apparent in the visible spectrum by the intense pink coloration of hexaaquacobalt(II) complex. However, this phenomenon was not observed. Second, the AFM topographies are not compatible with the idea of solid nanoparticles filling the whole inner cavity of the virus. Our experiments inform about two facts: first, Co does not span the whole virus cavity, and second, its structure is not compact enough to support a dehydration process. There are two hypothetical structural motifs compatible with the experimental observations: 1) Cobalt might be structured in a thin layer carpeting the inner virus wall; 2) It could present a porous structure, corresponding to a non-continuous metallic cluster like a sponge. These hypotheses are still consistent with the TEM images, which do not distinguish between solid nanoparticles

occupying all the inner space of the virus and either a hollow Co shell or a sponge-like Co cluster (Figure 6). These hypotheses imply the presence of a lower amount of magnetic material, consistent with the low magnetic signal revealed by our MFM experiments.

A rough approach to make a quantitative estimate of the cobalt inside one virus shell is to consider the virus as spherical shell with an inner cavity of ~ 12 nm in radius. By using the density of cobalt, the mass of metal filling a particle is 6.4×10^{-17} g (SI). Likewise, the mass of the proteinaceous shell reaches 5.6×10^{-18} g, thus indicating that for a 100% cobalt filled particle, 8% of the material is the protein shell (protein%) and 92 % of the sample accounts for the loaded cobalt (cobalt%). Therefore the ratio $\frac{\text{protein}\%}{\text{cobalt}\%} = 0.086$ for a solid filling. However, the TGA analysis (Figure 5) reveals that for the Co-VLPs this ratio increases to $\frac{55}{45} = 1.22$, suggesting that the virus particles are not fully loaded with cobalt. We can perform a similar estimation with the AFM topographic data of Figure 3 (SI) to find a ratio of $\frac{55}{45} = 1.22$. Interestingly, both bulk and single-molecule topography techniques coalesce in estimating the amount of cobalt inside the Co-VLPs: the cobalt fills about 10% of the viral cavity volume (SI), supporting the absence of a solid filling.

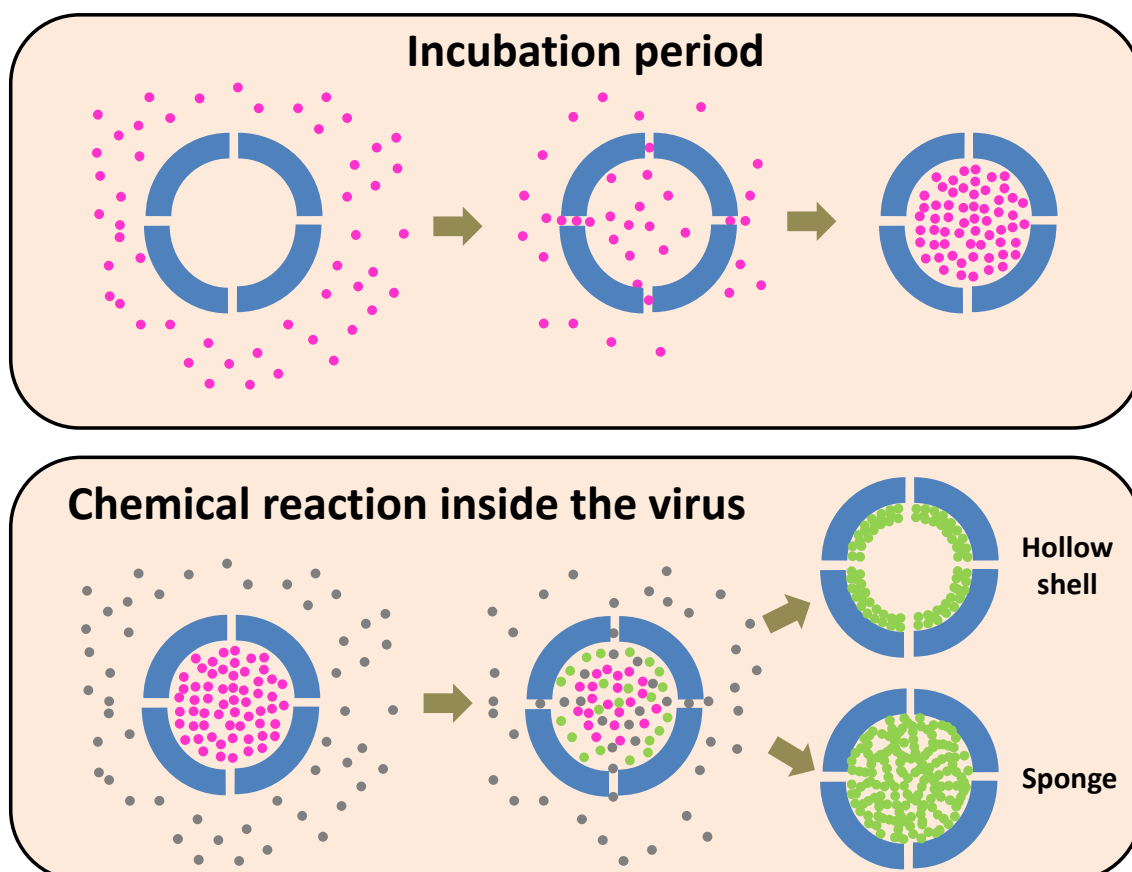


Figure 6. Schematic representation of the steps involved in the formation of cobalt within the virus capsid and of the chemical reaction for filling the capsid. The small pink spheres represent Co cations diffusing through the 12 pores on the virus capsid to the interior, and the incubation time allows interactions with the negatively charged amino acids present inside the capsid. The subsequent addition of the reducing agent NaBH₄ (black spheres) generates cobalt either as a hollow shell or a discontinuous cobalt cluster.

Under the experimental conditions used, our findings may challenge the concept that the cavity of a virus can be completely filled by a nanoparticle grown *in situ* by a single chemical reaction inside the virus capsid. However, modification of the loading process might, in the future, lead to fuller loading by means of successive incubation-reduction cycles.

Conclusions

Herein, we challenge the way to think about the use of empty viruses as nanoreactors to obtain size constrained magnetic nanoparticles. It is known that the spontaneous nucleation of metal oxides, simultaneous with the entry of the precursors inside a virus cavity, results in the gradual growth of nanoparticles². However, not much attention has been paid to the architecture and specific performance of the formed structure at single particle level. In our work we have utilized a twofold approach to characterize individual cobalt structures grown within CPMV eVLPs, including both AFM topographies upon desiccation and magnetic force microscopy of individual particles. Both approaches suggest that the cobalt grown inside the virus shell can be described either as a hollow shell coating the inner wall of the virus or a porous sponge-like cobalt structure, revealing an incomplete filling of the virus cavity (10%), which we relate to the single incubation step of eVLPs in a solution containing a metal salt. Such non-compact metal distribution can be due to the fact that the amount of metal ions incorporated inside the virus during a single incubation period is never enough to fill all the empty space after chemical reduction. Consequently, an increase in the amount of magnetic material inside the capsid would require the implementation of multi-step protocols including successive incubation-metal-reduction cycles.

Acknowledgements

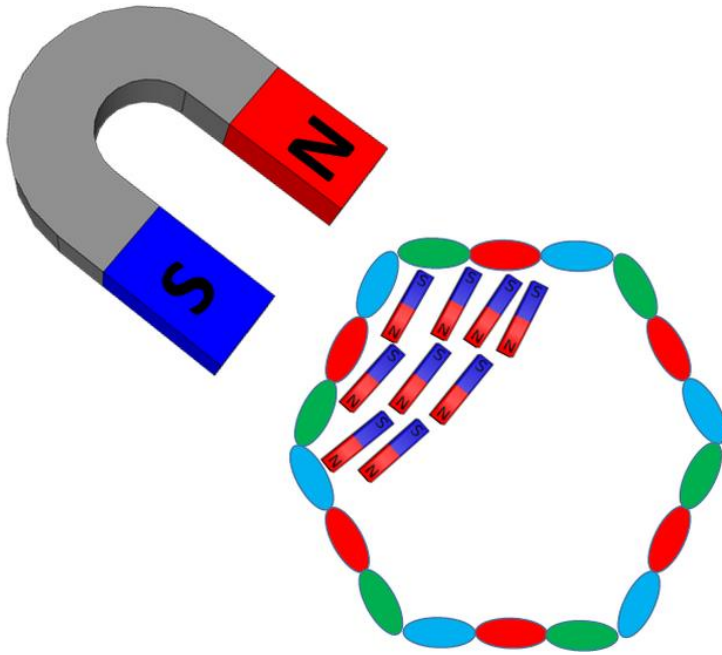
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