Encapsulation of Inorganic Nanomaterials inside Virus-Based Nanoparticles for Bioimaging

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Abstract

Virus-based nanoparticles (VNPs) can serve as containers for inorganic nanomaterials with excellent physical and chemical properties. Incorporation of nanomaterials inside the inner cavity of VNPs has opened up lots of possibilities for imaging applications in the field of biology and medicine. Encapsulation of inorganic nanoparticles (NPs) in VNPs can achieve the labeling of VNPs with nanoprobes and maintain the original outer surface features of VNPs at the same time. In return, VNPs enhance the stability and biocompatibility of the inorganic cargoes. This review briefly summarizes the current typical strategies to encapsulate inorganic nanomaterials in VNPs, i.e. mineralization and self-assembly, as well as the applications of these hybrid nanostructures in the field of bioimaging, including in vitro and in vivo fluorescence imaging, magnetic resonance imaging, and theranostics. Nanophotonic studies based on the VNP platform are also discussed. We anticipate that this field will continue to flourish, with new exciting opportunities stemming from advancements in the rational design of VNPs, the development of excellent inorganic nanomaterials, the integration of multiple functionalities, and the regulation of nano-bio interfacial interactions.

Key words: virus-based nanoparticle, encapsulation, mineralization, self-assembly, bioimaging, theranostics.

Introduction

Viruses are infectious agents in the biological sense, while in the chemical sense, they are highly ordered assemblies of biomacromolecules, with well-defined structures and monodisperse sizes in the range from tens to hundreds of nanometers. Generally, a virus has a proteinaceous capsid with the genomic nucleic acid inside, and enveloped viruses possess an additional membrane layer wrapping the capsid. Viruses are sometimes looked upon as natural nanomaterials due to their characteristic sizes and structures. Accordingly, the term virus-based nanoparticles (VNPs) have come into being. They are formulations based on viruses or virus-like particles, which are often assembled from multiple copies of viral capsid proteins (CPs) with a hollow inner cavity [1-3].

Encapsulation is a common strategy used by viruses to protect and deliver their genetic materials. In fact, it is also used by lots of other biostructures for specific biological reactions and processes under confinement, e.g. carboxysome [4], Eut microcompartment [5], and ferritin [6, 7]. In recent years, encapsulation of artificial cargoes-like inorganic nanomaterials, synthetic RNA, small molecule drugs, metal ions, and organic fluorophores in VNPs, has emerged as a unique strategy to integrate various functionalities for diverse purposes [1, 8, 9]. This review seeks to summarize the encapsulation of
inorganic nanomaterials with emphasis on the encapsulating strategies and bioimaging applications. Encapsulation of inorganic materials in VNP{s} is a typical and beautiful example of the interplay between biology and nanotechnology. The inorganic cargoes render VNP{s} new physical or chemical properties [2, 3]. In return, VNP{s} increase the biocompatibility and facilitate addressable functionalization of the inorganic nanomaterials. In combination, the hybrid nanoparticle-containing VNP{s} (NP@VNP{s}) provide a versatile nanoplatform for the fabrication of theranostic nanostructures or nanodevices.

**Strategies for encapsulation of inorganic nanomaterials in VNP{s}**

A variety of VNP{s} have been explored to encapsulate inorganic nanomaterials. There are two ways to load inorganic nanomaterials inside VNP{s}, i.e. mineralization in the inner cavity of VNP{s} and encapsulation of pre-formed nanomaterials through controllable self-assembly (Table 1).

**Mineralization**

The inner cavity of VNP{s} can serve as a nanoreactor for mineralization of inorganic nanomaterials with controllable sizes. In order to mineralize specifically inside VNP{s}, precursor metal ions should be attracted and trapped in the inner cavity rather than on the outer surface of VNP{s}. This can be achieved through electrostatic interaction, using ion affinity tags, or equipping seeds for mineralization.

In 1998, Douglas and co-workers first described the mineralization of paratungstate and decavanadate inside CCMV VNP{s} by means of electrostatic interaction. The pores on the VNP shell exhibit a pH-dependent gating mechanism. They swelled at pH>6.5, allowing the negatively charged polynometallic salts to be attracted by the positively charged interior interface, and closed at pH<6.5, resulting in the entrapment and oligomerization of polynometallic salts (Fig. 1A) [10]. Similarly, β-TiO₂ nanoparticles (NP{s}) were also successfully synthesized inside CCMV VNP{s} [11]. The charge of the interior surface of VNP{s} can be modified through genetic engineering to extend the spectrum of precursors. The nine basic residues at the N-terminus of the CCMV subunit were replaced with glutamic acid to switch the interior surface of VNP{s} from cationic to anionic, which favored strong interaction with positively charged ferric precursors and led to size-constrained iron oxide formation inside CCMV VNP{s} [12]. For CPMV, the inner surface is rich in negatively charged amino acids, which has been used to mineralize iron oxide and Co NP{s} [13,14]. Electrostatic interaction has also been exploited to mineralize iron oxide and Co NP{s} inside VNP{s}. Neutral or alkaline condition was used to reduce the positive charge of the outer surface and increase the negative charge on the inner surface of TMV VNP{s}. At pH 7.0, cationic silver precursors were preferentially sequestered within the central channel, which led to the growth of aligned discrete Ag NP{s} within the internal channel of TMV VNP{s} [15]. Another rod-shaped VNP, ToMV, has also been used for fabrication of 3 nm magnetic NP{s} (Co/Pt alloy) aligned within its channel by introducing lysine facing the inner surface to increase positive charges [16]. The flexibility of VNP{s} to be rationally engineered to switch interior surface charges based on their structural information enriches the possibilities of synthesizing different species of NP{s} inside VNP{s}.

Genetically introducing metal affinity tags is another approach to tune inorganic NP deposition inside VNP{s}. P22 VNP is assembled from 420 identical copies of CP with approximately 300 copies of scaffolding protein (SP) binding on the inner surface. SP can be genetically modified with different affinity peptides to provide nucleation sites for a variety of inorganic NP{s}. For examples, Fe₂O₃, TiO₂, and CdS NP{s} were successively synthesized inside P22 VNP{s} by this approach [17-20]. In the case of TMV VNP{s}, mutation of threonine 103 to cysteine dramatically enhanced the assembly capability and stability of TMV rod-like structure due to the formation of disulfide bonds between adjacent subunits [21]. Then growth of AuNP chains (AuNC{s}) and Au nanorods (AuNR{s}) was realized in the internal channel of the mutant TMV VNP{s}, mediated by the strong coordination interaction between Au³⁺ and sulfhydryl of the introduced cysteine (Fig. 1B) [22]. The single mutation of cysteine 103 successively played two important roles, namely, stabilizing TMV rod-like template and mediating the deposition of Au.

Equipping inorganic seeds on the inner surface is an alternative route for NP mineralization inside VNP{s}. Taking the advantages of natural affinity to Pd or Pt, a series of nanowires (NWs), such as Ni, Co, Cu, Co/Fe, Co/Pt, and FePt NWs of 3 nm in diameter and a few hundred nanometers in length have been synthesized inside the central channel of TMV VNP{s} after the activation of Pd or Pt followed by electroless deposition [23-27]. In addition to direct nucleation of seeds on the inner surface, preformed NP{s} can also be equipped inside the inner cavity of VNP{s} as seeds for subsequent growth of NP{s}. Most recently, we have established such a route with SV40 VNP{s} to mineralize AuNP{s} and Au@Ag core-shell NP{s}. A preformed 5 nm AuNP was encapsulated inside SV40
VNP as the seed, which was followed by further growth of an outer metal layer. This method circumvents the limitation that only gentle reaction conditions can be used for seed synthesis inside viral protein cages to avoid protein impairment. The encapsulation of the preformed 5 nm seed was realized by virtue of the self-assembly process of SV40 VNP, which will be discussed in the next section [28].

It is worth mentioning that in addition to VNPs, other protein cages, for example, the ferritin family, have also been versatile templates for constrained synthesis of nanomaterials. There are several excellent reviews to be referred to in this regard [29,30].

**Table 1. A brief summary of NP@VNPs and their bioimaging applications.**

| Viruses                          | Encapsulation strategies | Encapsulated inorganic nanomaterials                                      | Applications in bioimaging               |
|---------------------------------|--------------------------|---------------------------------------------------------------------------|-----------------------------------------|
| CCMV (cowpea chlorotic mottle virus) | Mineralization           | Paratungstate and decavanadate [10], β-TiO2 [11], iron oxide [12] NPs     | Nanophotonics [62]                      |
| BMV (brome mosaic virus)        | Self-assembly            | AuNP [63-65]                                                              | Fluorescence imaging in cells [49]      |
| CPMV (cowpea mosaic virus)      | Mineralization           | CoNP [14]                                                                 | MRI [41,57]                             |
| RCNMV (red clover necrotic mosaic virus) | Self-assembly          | Au, CoFeOx, and CdSe NPs [37,38]                                          | MRI [41,57]                             |
| MS2 (bacteriophage)             | Self-assembly            | AuNP [62], CdSe@ZnS QD [67]                                               | Fluorescence imaging in cells [49]      |
| P22 (bacteriophage)             | Mineralization           | Fe3O4 [17], TiO2 [18], GdS [19, 20] NPs                                   | MRI [41,57]                             |
| SV40 (simian virus 40)          | Mineralization           | CdSe@ZnS QD [35, 49, 59, 61, 69, 70], AuNP [69, 71], Ag5 S QD [50, 60], MNP [66] | MRI [41,57]                             |
| HBV (hepatitis B core virus-like particles) | Self-assembly          | Fe3O4 NP [41], FeO/FeO, Ni [72], γ-Fe3O4@AuNP [57]                         | MRI [41,57]                             |
| Alpha-virus (nucleo-capsid)     | Self-assembly            | AuNP [73]                                                                 | Fluorescence imaging in cells [39]      |
| PTLV (pseudotyped HIV-1-based lentivirus) | Self-assembly       | CdSe@ZnS QD [39]                                                          | Fluorescence imaging in cells [40]      |
| HIV-1 (human immunodeficiency virus type 1) | Self-assembly     | CdSe@ZnS QD [40]                                                          | Fluorescence imaging in cells [40]      |
| TMV (tobacco mosaic virus)      | Mineralization           | Ni [24, 25], Co [24], Cu [27], Co/Fe [26], Co/Pt and FePt NWs             | MRI [41,57]                             |
| ToMV (tomato mosaic virus)      | Mineralization           | Co/Pt NP [16]                                                             | MRI [41,57]                             |

**Figure 1.** Encapsulation of nanomaterials inside VNPs by mineralization. A) Upper: Schematic illustration of the synthetic approach for mineralization within the CCMV VNPs. Step I involves the removal of viral RNA and purification of the empty VNPs by sucrose gradient centrifugation. Step II involves the selective mineralization of an inorganic paratungstate species within the confines of the VNPs, at pH 5.5 and 6°C. Lower: TEM images of paratungstate-mineralized VNPs. a) An unstained sample showing discrete electron dense cores; b) A negatively stained sample of (a) showing the mineral core surrounded by the intact VNP protein cage. Scale bar: 100 nm. Reproduced with permission [10]. B) Site-specific biomineralization of gold nanostructures in TMV rod-like VNPs. The cysteines arrayed in the interior of mutated nanotubes are specifically bound with the gold precursors, which initializes the controlled growth of AuNCs (a) or AuNRs (b). Scale bars: 10 nm. Reproduced with permission [22].
**Self-assembly**

VNPs can undergo the disassembly-assembly cycle controlled by buffer conditions, which has been widely utilized to encapsulate guest cargoes. Inorganic nanomaterials of different surface properties, components, and sizes have been encapsulated into VNPs of different viruses (Table 1).

For encapsulation in VNPs, the surface coatings of inorganic NPs, which play an important role in determining the interactions between CPs and inorganic NPs, have been the primary consideration. What surface properties are required for an inorganic NP to be encapsulated by VNPs? The central idea is to mimic the packaging of viral genome. Therefore, negatively charged coatings are often used to make NPs to mimic viral nucleic acids to initiate the assembly of VNPs. However, different species of negatively charged surface coatings may present different performance in initiating the assembly of VNPs. In an early report by Chen et al., AuNPs coated by carboxylate-terminated thiolalkylated tetraethylene glycol (TEG) reached a high encapsulation efficiency of 95% by BMV VNPs. However, in the case of citrate-coated AuNPs, serious aggregation occurred during the reassembly procedure (Fig. 2A) [31]. Similarly, quantum dots (QDs) coated by H5-poly(ethylene glycol) (PEG)-COOH showed the highest ability in promoting BMV VNP assembly in comparison to QDs functionalized with phospholipid micelle, streptavidin-biotin-DNA, or dihydrolipoic acid (DHLA) [32]. Those surface coatings, which can keep the stability of inorganic NPs as well as provide efficient nucleation and/or propagation sites for the assembly of VNPs, are excellent in promoting the encapsulation process [31]. Furthermore, the effect of surface charge density of NPs on the encapsulation by BMV VNPs was studied with AuNPs functionalized by a mixture of HS-TEG-COOH and HS-TEG-OH. It was found that a critical surface charge density was required for BMV VNP assembly [33]. According to these experimental observations, the authors presented a model to describe the encapsulation of spherical particles functionalized with weakly acidic surface groups, which demonstrated that surface charge regulation in combination with size polydispersity of the NPs has a large effect on the encapsulation efficiency [34]. In contrast, a quite different case is presented by SV40 VNP. It showed that the encapsulation of NPs does not rely on the surface charge of NPs, as negatively charged mercaptopropanoic acid (MPA)-coated QDs and DNA-coated QDs, neutrally charged methoxy-terminated PEG-coated QDs, and positively charged amine-terminated PEG-coated QDs could be encapsulated by SV40 VNP with comparable efficiencies [35]. The lack of dependence of NP encapsulation by SV40 VNP on electrostatic interaction may be related to the unique structural organization of natural SV40 virion. SV40 genome DNA forms a minichromosome with host histones that neutralize the negative charges of DNA, and the interaction between the capsid and the minichromosome is mediated by two minor CPs [36]. Such a structure feature suggests that genome DNA encapsulation of SV40 depends much less on electrostatic attraction than those of viruses like CCMV and BMV.

Packaging signals of specific nucleotide sequences have evolved in many viruses to guide selective encapsulation of viral genomes. Such specific interactions between nucleic acids and CPs have been exploited to direct NP encapsulation by VNPs. As a proof of concept, oligonucleotide mimics of the origin of assembly sequence (OAS) in RCNMV were attached to NPs to recruit RCNMV CPs and to direct the assembly of VNPs. Using this strategy, Au, CoFe2O4, and CdSe NPs ranging from 3 nm to 15 nm in diameter were successfully encapsulated inside RCNMV VNPs (Fig. 2B). Attempt to encapsulate NPs larger than 17 nm did not result in well-formed RCNMV VNPs, which is consistent with the presence of a 17 nm cavity in native virion [37,38]. This strategy has also been used to encapsulate NPs inside enveloped virus in living cells. QDs conjugated with modified genomic RNAs (gRNAs), were packaged into enveloped virus together with the gRNAs (Fig. 2C) [39]. Besides oligonucleotide sequences, other inner components of virus can also be used to mediate the encapsulation of NPs. Recently, QDs modified with the accessory protein Vpr have been encapsulated into HIV-1 virions during virus assembly [40]. In addition to learning from natural packaging mechanisms, other specific interactions have also been utilized to facilitate the encapsulation of NPs by VNPs. For example, nickel-nitritolactric acid (Ni-NTA) functionalized Fe3O4 NPs efficiently templated the assembly of HBV VNP with carboxyl-terminally appended polyhistidine tags [41].

The size of inorganic NPs is another factor affecting the encapsulation by VNPs. AuNPs with different diameters guided the assembly of BMV VNPs into T=1, T=2 or T=3 icosahedral structures, respectively, resembling to the three classes of BMV viral particles found in host cells (Fig. 2D) [42]. Phospholipid micelle-coated magnetic NPs which was larger than the inner cavity of native BMV capsid was also found to be able to template the assembly of BMV CPs into VNPs larger than the native T=3 BMV assembly. 

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For HBV VNPs, the VNP diameter followed the increase of cargo size linearly until it reached the value expected for the $T=4$ capsid, and then remained unchanged upon further increase of the cargo diameter up to 18% [44]. The flexibility of VNPs to encapsulate various sizes of NPs to form polymorphic assemblies, makes them more versatile in biomedical applications, because shape and size are two important factors that affect the in vivo behaviors of VNPs [45].
Besides the surface properties and size effect of cargo NPs, environmental parameters also play a role in the encapsulation by VNP. For instance, the buffer conditions, which not only control the disassembly-assembly cycle, but also influence the assembly pathway. It was reported that depending on the pH value and ionic strength, the assembly of BMV VNP around anionic AuNPs could be cooperative or non-cooperative adsorption. At high pH and low ionic strength, non-cooperative Langmuir adsorption of the BMV CPs around the anionic core occurred, whereas at low pH and low ionic strength, initial cooperative CP-AuNP association followed by CP rearrangement took place to form a regular capsid-like structure [46]. In addition, the buffer conditions may influence encapsulation by affecting cargo stability. Some protein cages, such as MS2 VNP and ferritin, need an ultra-low pH treatment to disassemble. When NPs are mixed with the disassembled CP oligomers for encapsulation, they would be prone to precipitation and fail to be encapsulated. This problem can be solved by finding out either proper surface coatings of NPs or new strategies to control the disassembly-assembly cycle of VNP.

NP@VNP hybrid nanostructures for bioimaging

Encapsulation of inorganic nanomaterials in VNP has inspired people to come up with new ideas about biolabeling and bioimaging. In this section, we review the recent advances of bioimaging applications of NP@VNPs, including live cell imaging, \textit{in vivo} imaging, magnetic resonance imaging (MRI), and nanophotonics. The most recently designed hybrid multifunctional NP@VNPs as a novel nanotheranostic tool is introduced. Future opportunities and challenges of NP@VNPs in bioimaging field are also discussed.

Fluorescence imaging

Fluorescence imaging is now an evolving field that has already achieved significant advances, due to the accumulative availability of fluorescent proteins, dyes and probes, as well as the development of optical imaging technologies [47]. As a typical representative fluorescent nanoprobes, QDs emit in the visible and near infrared wavelength region with high brightness and excellent photostability [48]. However, it has been controversial that labeling QDs on virus or VNP outer surfaces would affect virus-host cell interactions and might cause artifacts. Therefore, encapsulation of QDs in VNP cavities provides an alternative way that has attracted increasing interests.

The first demonstration of imaging viral behaviors in living cells with encapsulated QDs was reported with the self-assembling VNPs of SV40 in 2009. By real-time and long-term tracking of the cellular uptake and intracellular transportation, it was found that SV40 QD@VNP was internalized by cells through caveolae-mediated endocytosis, then travelled along the microtubules and accumulated in the endoplasmic reticulum, like the wild-type SV40 (Fig. 3A) [49], suggesting that the early infection steps of SV40 are basically determined by the major CP VP1.

Encapsulation of QDs inside VNP eliminates in principle the disadvantages caused by QD modification on outer surface of viruses and avoids controversies over the modification strategies. QD encapsulation therefore enables high-fidelity imaging, which should benefit the investigation of infection events of other viruses. However, the most challenging is to label and track live viruses, not merely VNP, using such a strategy.

Subsequent efforts succeeded in encapsulating QDs inside the core of vesicular stomatitis virus glycoprotein (VSV-G) pseudo-typed HIV-1-based lentivirus (PTLV) during the assembly process of the virus in living cells. Revealed by the real-time single particle tracking of QD@PTLVs, there was significant colocalization between the virus and early endosomes. In the following steps, the trajectory and velocity of single virus particle on microtubules were recorded, and a microtubule-dependent QD@PTLVs cellular entry route was clearly drawn [39]. Most recently, productive entry of HIV-1 into human primary macrophages has been fluorescently tracked at the single particle level by taking advantages of encapsulated QDs, which revealed that the infection process requires clathrin-mediated endocytosis, endosomal escape of the viral core, and actin dynamics (Fig. 3B) [40]. Both studies are successful examples of live virus labeling and imaging by the QD encapsulation strategy. The high brightness and photostability of QDs endow real-time virus tracking to the single particle level.

When \textit{in vivo} deep tissue imaging is needed, challenges of high absorbance, serious scattering and strong autofluorescence of the tissue to the visible fluorescence should be overcome to reach deep tissue imaging with desired temporal and spatial resolution. The second near infrared (NIR-II) optical window (1000-1700 nm) of tissues has been utilized as a solution to the problem, because tissues have minimal optical extinction and autofluorescence in the NIR-II range. AgS QDs have proved to be an excellent NIR-II probe with high brightness, stability, and good biocompatibility. By encapsulating the AgS QDs,
dynamics and biodistribution of SV40 VNPs in live mice were clearly monitored in real time. It was found that Ag$_2$S@VNPs quickly accumulated in liver, spleen, and bone marrow within 5 min post injection. Surface PEGylation of VNPs showed comprehensive impact on the in vivo behaviors of SV40 VNPs, including greatly elongated blood circulation time and significantly reduced accumulation in the reticuloendothelial system (Fig. 3C). Encapsulation of Ag$_2$S QDs enabled real-time monitoring of surface chemistry-dependent in vivo behaviors of VNPs with unprecedented spatiotemporal resolution [50]. The hybrid Ag$_2$S@VNPs might be a useful nanoplatform for future theranostics.

**Iron oxide NP@VNPs for magnetic resonance imaging**

MRI is now an advanced technique very useful in biomedical research and disease diagnosis. It is a non-invasive imaging tool that produces three-dimensional anatomical images without damaging radiation. Most MRI contrast agents enhance imaging contrast by changing the relaxation rates ($1/T_1$ or $1/T_2$) of tissue water protons, therefore improving the sensitivity and specificity for detection [51]. Viral capsids provide the advantages of suitable sizes and shapes that can increase the chances to enhance the relaxation rates as well as an ideal modification platform for contrast agents labeling [52]. In 2005, Douglas et al. developed VNPs as MRI contrast agents by chemically attaching Gd$^{3+}$ to CCMV VNPs [53]. Subsequently, several other VNPs have been investigated as potential MRI contrast agents labeled with Gd$^{3+}$. Detailed labeling strategies and applications of Gd$^{3+}$-VNPs contrast agents have been summarized in recent review papers [3,9,54,55]. Here we concentrate on diverse MRI applications of encapsulated iron oxide NPs in VNPs.

![Figure 3.](http://www.ntno.org)

**Figure 3.** Real-time fluorescence bioimaging through encapsulation of QDs in VNPs. A) Live fluorescence microscopy of Vero cells transiently expressing caveolin-1-CFP and incubated with SV40 QD@VNPs, showing that SV40 QD@VNPs (red) colocalized with the caveolae marker, caveolin-1 (green). Scale bar: 10 μm. Reproduced with permission [49]. B) Visualization of HIV-1 dynamic productive entry via fusion of the viral envelope with Rab5A-positive endosome. The cellular boundary of the macrophage is highlighted by a dashed line. Dual-labeled QD@HIV-Dio (green and red) was shown to colocalize with ECFP-Rab5A-marked endosome (blue) in a macrophage. Scale bar: 10 μm. Reproduced with permission [40]. C) In virtue of encapsulation of the Ag$_2$S QDs, NIR-II fluorescence imaging clearly distinguished the real-time distribution of the naked and PEGylated VNPs of SV40 in living mice at 12 h post injection. Reproduced with permission [50].
While Gd³⁺ is the most popular agent for positive MRI contrast, superparamagnetic iron oxide can effectively reduce the transverse relaxation time (T₂), producing negative contrast effect [54]. Application of VNPs-encapsulated iron oxide for MRI was firstly reported in 2007. Iron oxide NPs of tunable sizes were packaged inside icosahedral BMV VNPs via in vitro self-assembly [43]. The behaviors of iron oxide-encapsulating BMV VNPs in the host plant were examined. In this work, the cubic iron oxide exhibited outstanding characteristics as a negative contrast agent in two aspects. Firstly, the 18.6 nm cubic iron oxide NPs with enhanced charge density was chosen as inner core that could increase the stability of BMV VNPs and encapsulation efficiency. Secondly and more importantly, the cubic iron oxide NPs have much higher r₂/r₁ (transverse/longitudinal relaxivity) ratio than that of commercial contrast agents with similar sizes such as Feridex® and Supravist®, and other Gd³⁺-labeled VNPs. By taking transport assay inside N. benthamiana leaves, it was demonstrated that iron oxide-encapsulating VNPs penetrated cell wall into cytoplasm and cell junctions of intercellular space, and thus took cell-to-cell long-range transport, while free iron oxide NPs were blocked at the exterior of plant cells (Fig. 4A) [56]. This VNP-based contrast agent may be meaningful for investigation of plant growth and development. Also, it offers a new opportunity for high-performance contrast probes design based on VNP platforms. Certainly, the size of the iron oxide core encapsulated in the VNPs should be considered to maintain natural structure of viral capsids [43]. For example, a well-designed core-shell NP, γ-Fe₂O₃@AuNP11 nm in size was optimal for VNP encapsulation and presented significantly higher r₂/r₁ ratio at 115.8 [57].

Figure 4. Iron oxide NPs encapsulated in VNPs as MRI contrast reagents. A) T₂-weighted MR images of HBV VNPs encapsulating a) 3.4 nm, b) 6.1 nm, and c) 11.7 nm of Fe₃O₄ NPs. Increasing the Fe concentration significantly changes the signal intensity in the MR images. B) Plot of 1/T₂ versus Fe concentration of the Fe₃O₄ NPs of various sizes encapsulated in HBV VNPs with the slope indicating the specific relaxivity (r₂). Reproduced with permission [41].
verified [59]. This work integrates three functionalities including fluorescent imaging, molecular targeting, and drug delivery, and represents an important step in developing multifunctional NP@VNPs for theranostics. More endeavors are further needed to advance such kinds of designer nanostructures to clinical practices.

**NP@VNPs for nanophotonics**

As VNPs are highly symmetrical and monodisperse in a suitable size range, they provide an ideal design platform to explore the photonic interactions between metal NPs and/or fluorophores.

The first observation on the coupling of surface plasmon resonance (SPR) in AuNP@VNPs was taken as an obvious color change caused by encapsulation of multiple AuNPs in a single VNP [60]. Later, interactions between AuNPs and QDs were described from a quantitative view based on a hybrid AuNP-QD system templated by SV40 VNPs, in which AuNPs were assembled onto the outer surface of SV40 VNPs encapsulating a QD in the cage. By tuning the number of AuNPs on QD@VNPs from one to twelve, the optical interaction between QDs and AuNPs could be quantitatively evaluated. It was observed that the fluorescence of QDs decreased dramatically with increasing AuNP:QD ratio. No fluorescence enhancement was observed because of the long distance of 8 nm between AuNPs and QDs (Fig. 6A) [61]. Later, Francis et al. designed a well-defined hybrid system for analyzing the interactions caused by nanoscale distance between NPs and fluorescent organic dyes. They firstly encapsulated AuNPs within the VNPs of bacteriophage MS2, and then introduced aniline groups by inserting p-aminophenylalanine amino acid to connect exterior dye-labeled DNA strands. By adjusting the length of DNA to control the distance between dyes and AuNPs, AuNP-enhanced fluorescence was observed using a total internal reflection fluorescence (TIRF) microscope (Fig. 6B) [62]. Herein, by encapsulating the AuNPs, MS2 VNPs served not only as a scaffold for controllable docking of fluorescent dyes, but also as a physical layer to avoid direct sticking of the dyes onto AuNPs, leading to fluorescence quenching.

**Conclusions and Outlooks**

It has been nearly twenty years since the first study on loading inorganic nanomaterials in VNPs was reported. Great potentials have been recognized with these protein-inorganic hybrid nanostructures. This review has briefly summarized the development of inorganic NP encapsulation inside VNPs and bioimaging applications thereof. Various methods have been established for NP encapsulation, which can be categorized into mineralization and self-assembly. The fabricated hybrid NP@VNP nanostructures have shown unique advantages in bioimaging applications in terms of sensitivity, resolution and fidelity.

Although NP@VNPs provide a promising platform for theranostic purposes, their biomedical application is still in its infant stage. Many of the bioimaging studies reported so far have been conducted in vitro, with only a few performed in living animal models. Currently, a major challenge is to target NP@VNPs to a specific region of a living body. Investigations in interactions between VNPs and biomolecules in body fluid, in vivo stability and fates of NP@VNPs, as well as the rational design of VNPs will help to find out solutions. At the same time, more endeavors are needed to integrate multiple functionalities into NP@VNPs to meet clinical requirements. Owing to the addressability, versatility, and designability of the VNP nanoplatform, these challenges will be hopefully overcome.

![Figure 5](http://www.ntno.org)
Figure 6. NP-NP and NP-dye photonic interactions investigated using the NP@VNPs platform. A) In SV40 VNP-templated 3D discrete hybrid AuNP (yellow)/QD (red) nanoarchitectures (Left), very weak SPR coupling of AuNP clusters (Middle) and AuNP number-dependent shortening of life-time of QD fluorescence (Right) were observed. Curves i-viii correspond to the number of AuNPs per VNP being 1, 3, 5, 6, 8, 10 and 12, respectively; ix, instrument response. Reproduced with permission [61]. B) Dye-labeled DNA was attached to the exterior of MS2 VNPs that encapsulate a 10 nm AuNP. The dye-AuNP distance was tuned using DNA strands with different lengths. The fluorescence life-time (τ) of the dye was shortened as the dye and AuNP got closer (Left). Distance-dependent fluorescence enhancement of the dye by AuNPs was quantitatively determined using TIRF microscopy (Right). Enhancements of 2.2-fold and 1.2-fold were observed for 3 bp and 12 bp separations, respectively, while a 24 bp separation showed no effect. Reproduced with permission [62].

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Competing Interests

The authors have declared that no competing interest exists.

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