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# Nitroxyl Modified Tobacco Mosaic Virus as a Metal-Free High-Relaxivity MRI and EPR Active Superoxide Sensor

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Supporting Information

**ABSTRACT:** Superoxide overproduction is known to occur in multiple disease states requiring critical care; yet, noninvasive detection of superoxide in deep tissue remains a challenge. Herein, we report a metal-free magnetic resonance imaging (MRI) and electron paramagnetic resonance (EPR) active contrast agent prepared by "click conjugating" paramagnetic organic radical contrast agents (ORCAs) to the surface of tobacco mosaic virus (TMV). While ORCAs are known to be reduced in vivo to an MRI/EPR silent state, their oxidation is facilitated specifically by reactive oxygen species—in particular,



superoxide—and are largely unaffected by peroxides and molecular oxygen. Unfortunately, single molecule ORCAs typically offer weak MRI contrast. In contrast, our data confirm that the macromolecular ORCA-TMV conjugates show marked enhancement for  $T_1$  contrast at low field (<3.0 T) and  $T_2$  contrast at high field (9.4 T). Additionally, we demonstrated that the unique topology of TMV allows for a "quenchless fluorescent" bimodal probe for concurrent fluorescence and MRI/EPR imaging, which was made possible by exploiting the unique inner and outer surface of the TMV nanoparticle. Finally, we show TMV-ORCAs do not respond to normal cellular respiration, minimizing the likelihood for background, yet still respond to enzymatically produced superoxide in complicated biological fluids like serum.

KEYWORDS: tobacco mosaic virus, reactive oxygen species, magnetic resonance imaging, electron paramagnetic resonance, bioconjugation, organic radical contrast agents

#### INTRODUCTION

The upregulation of reactive oxygen species (ROS), in particular, superoxide, is associated with certain cancers, neurodegenerative disorders<sup>1,3,4</sup> such as Parkinson's disease,<sup>5</sup> diabetes, 6,7 mitochondrial diseases such as Friedreich ataxia, 8 Leber's hereditary optic neuropathy (LHON), mitochondrial encephalomyopathy, lactic acidosis, stroke-like episodes (MELAS), <sup>10</sup> myoclonic epilepsy with ragged red fibers (MERRF), <sup>11</sup> and Leigh syndrome (LS). <sup>3,4,12,13</sup> The detection of superoxides, both extra and intracellularly, has consequently become an area of research interest in efforts to study and target diseased tissues. 14,15 Superoxide sensing in biological tissues has focused mainly on fluorescent imaging, as many dyes are easily quenched in the presence of ROS, 16,17 and superoxide in particular.<sup>18</sup> Moreover, the superoxides produced by people with diabetes can cause serious health complications such as heart injuries, and sensing of superoxide overproduction in these situations is also done using fluorescence imaging. 19,20 An

unavoidable issue with fluorescence imaging in general, however, is limited tissue penetration, making it problematic for tissue imaging but ideal in guided surgery applications. Consequently, the need to detect cellular and biological events in deep tissue noninvasively has driven the development of "smart" probes for magnetic resonance imaging  $(MRI)^{21-23}$  or electron paramagnetic resonance (EPR) modalities. 14,24 On the MRI front, paramagnetic chemical exchange saturation transfer (paraCEST) and other stimuli-responsive contrast agents have begun to generate considerable interest for their ability to detect changes in extracellular pH or production of specific

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biological markers like lactate.<sup>25–27</sup> While redox active paraCEST agents based on lanthanides have emerged as potential sensors of singlet oxygen<sup>28</sup> and peroxide,<sup>29</sup> none have clearly emerged as candidates for superoxide.<sup>25</sup> Paramagnetic nitroxide organic radical contrast agents (ORCAs), on the other hand, have been making inroads toward superoxide detection in vivo and in vitro.<sup>14,30–32</sup> While sterically unencumbered nitroxide radicals like TEMPO (Figure 1b)

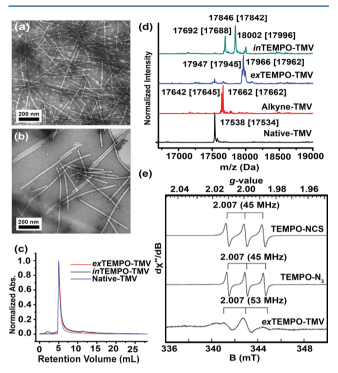


Figure 1. Characterization of TMV after bioconjugation reactions. The TEM images of (a) exTEMPO-TMV and (b) inTEMPO-TMV shows the expected rod-like morphology. (c) A single peak in the SEC chromatogram (at 260 nm) of modified TMV confirms that the solution dispersity was unchanged. (d) Bioconjugation of the TEMPO radical to the TMV was characterized by ESI-MS. Native TMV has a peak at 17 538 m/z. The loss of this peak and emergence of a new peak at 17 662 m/z confirms that TMV was completely modified to alkyne-TMV. ESI-MS of exTEMPO-TMV shows complete conjugation of TEMPO-N $_3$  to the alkyne-TMV. The peak at 17 966 m/zrepresents the attachment of the TEMPO radical to the TMV coat protein, whereas the peak at 17 942 m/z is the attachment of the TEMPO radical to the alkyne group with an unidentified elimination first noted by Francis (see ref 44) that has taken place during diazonium reaction. The interior surface of TMV consists of several glutamate residues, and three sites were modified via EDC coupling with TEMPO—one (17692 m/z), two (17846 m/z), or three (18 002 m/z) TEMPO molecules were installed. (e) X-band EPR spectra of exTEMPO-TMV. The exTEMPO-TMV sample was prepared in a capillary tube to minimize interaction between high dielectric aqueous solvent and the electric field of the incident microwave radiation.

are easily reduced to diamagnetic hydroxylamines in biological media, making their use as long-lived MRI contrast agents problematic, this property has been utilized to map redox activity in tumors ex vivo and in vivo.<sup>33</sup> On the other hand, reoxidation is kinetically favored by superoxide, while less reactive ROS, and in particular hydrogen peroxide, do not significantly oxidize hydroxylamines back.<sup>1</sup> Consequently, TEMPO can serve as a "turn-on" MRI sensor for superoxide.

The benefit of "turn-on" over "turn-off" sensors is that they can have nearly infinite contrast against background.<sup>34</sup>

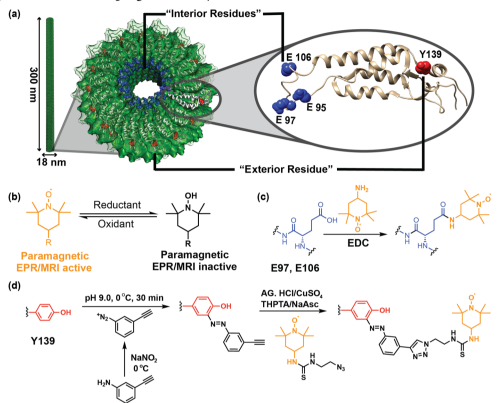
Nevertheless, ORCAs have historically suffered from low proton relaxation rates compared to metal-based contrast agents. These poor relaxivities have been cleverly addressed using polymeric systems, <sup>33,35-39</sup> as these multivalent scaffolds allow for high local concentrations of paramagnetic ORCAs. While such multimeric polymeric scaffolds allow for attachment of both MRI and fluorescence agents on the same platform, ORCAs are well-known and potent quenchers of adjacent fluorophores.<sup>33</sup> This quenching process means that when an ORCA is MRI/EPR active, adjacent fluorophores are typically quenched. While this has been elegantly exploited, <sup>32,33,40,41</sup> a true ratiometric sensor capable of assessing the probe concentration by fluorescence and the relative ROS levels by EPR/MRI signal would require a constant fluorescence emission regardless of the oxidation state of the ORCA.

Viral nanoparticles (VNPs) provide distinct structural features that make them well-suited to resolving some of these issues. For instance, they contain spatially well-defined functionalities, are rigid and monodisperse, and contain discrete topological surfaces that provide a distinct "outer" and "inner" surface. These distinctions are well-suited for EPR active spin systems prone to spin pairing and relaxation when in close contact with other radical species. Tobacco mosaic virus (TMV) is a 300  $\times$  18 nm rod-shaped RNA virus that contains 2130 individual coat proteins.  $^{43-45}$ 

Each of these coat proteins contains an easily functionalizable tyrosine residue (Y139) on the exterior surface, which is spatially separated from the next closest accessible tyrosine 46 by 2.3-2.6 nm. 47,48 Furthermore, TMV contains a 4 nm narrow pore lined with glutamic acid residues (E97 and E106), likewise easily functionalized, and electronically inaccessible to the exterior surface. These features provide two distinct advantages over existing polymeric nanostructures containing nitroxide radicals: (i) radical systems on the surface are spaced far enough apart so that spin-spin coupling is minimized even if 100% of the available functional groups are utilized, and (ii) fluorophores placed on the interior of the capsid will not be quenched by electron transfer processes from radical TEMPO moieties placed on the exterior surface. Previous studies have shown that VNPs and virus-like particles (VLPs) decrease the  $T_1$  and  $T_2$  relaxation times of attached metal-based MRI contrast agents. 41,48,52-63 The increase in  $r_1$  (1/ $T_{1p}$ , p = paramagnetic contribution; mM<sup>-1</sup>·s<sup>-1</sup>) relaxivity in these systems has been attributed to an increase in molecular diffusion and translation correlation times brought about by attachment to the surface of relatively massive, rigid, and slowly diffusing protein ensembles.

Here we present a method to create a sensitive EPR/MRI probe for superoxide detection in vitro with enhanced  $r_1$  and  $r_2$  relaxivities that rival clinically used molecular lanthanide-based MRI probes. The observed  $r_2/r_1$  ratio allows TEMPO-conjugated TMV probes to operate as both a  $T_1$  and  $T_2$  contrast agent, making these probes valuable for both clinical (<3 T) and preclinical (>3 T) MRI scanners. We have also designed a bimodal TEMPO-TMV probe for ratiometric fluorescence experiments that is functionalized with a fluorescent dye on the interior and with an ORCA for MRI/EPR imaging on the exterior. The unique topology of the VNP scaffold allows for the insertion of an internal fluorescent probe as a concentration marker that is insensitive to the oxidation

Scheme 1. (a) Representations of a Single TMV Nanoparticle<sup>a</sup> and a Single TMV Coat Protein Highlighting Solvent Exposed Amino Acid Residues Available for Bioconjugation Reactions, (b) Oxidation States of Nitroxide Radicals,<sup>b</sup> (c) Bioconjugation to the Interior Surface of TMV via EDC Coupling Reaction To Make *in*TEMPO-TMV, and (d) TMV Bioconjugation on the Exterior via Stepwise Diazonium Coupling Followed by a CuAAC Reaction To Make *ex*TEMPO-TMV



<sup>a</sup>Created in Chimera using PDB ID of 2tmv. <sup>b</sup>Paramagnetic radical species can be detected by both EPR and MRI, whereas the reduced diamagnetic species is neither MRI nor EPR active.

state of the ORCA probe. Finally, we present the evaluation of these probes in both cells and serum.

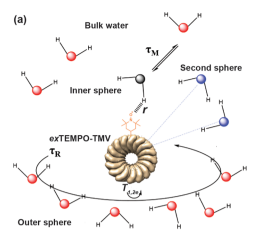
#### RESULTS AND DISCUSSION

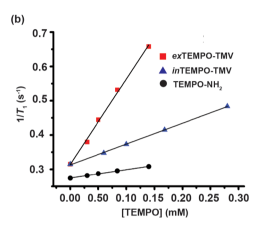
Attaching ORCA to the Surfaces of TMV. As shown in Scheme 1a, TMV contains two distinct surfaces: (i) an outer surface with few charged functional groups and uninhibited access to bulk water and (ii) a 4 nm narrow inner channel with a net negative surface charge and less access to diffusing bulk water. We thus first set to determine if there would be any difference in measured relaxivities if ORCAs were placed on the inner vs the outer surface of TMV. The interior surface of TMV contains solvent exposed glutamic acid residues, which we functionalized via EDC coupling with TEMPO-NH2 (Scheme 1c). High-performance liquid chromatography/electron spray ionization mass spectroscopy (HPLC/ESI-MS) confirms that three sites were modified following the EDC coupling reaction to yield inTEMPO-TMV. As shown in Figure 1d, peaks at 17 692 Da, 17 846 Da, and 18 002 Da represent the attachment of one, two, and three molecules of TEMPO, respectively, to the coat protein monomers of TMV. While only glutamate residues E97 and E106 have been identified as reactive sites, 49 the third peak at 18 002 Da likely corresponds to E95. Though three conjugations to this inner channel have been observed<sup>5</sup> before, the residue of the third modification site has never been identified. Separately, the exterior surface of TMV was functionalized initially using a diazonuim coupling reaction<sup>49</sup> (alkyne-TMV) with in situ prepared 3-ethynylphenyldiazonium

salt. This quantitative reaction was followed by a copper-catalyzed azide—alkyne cycloaddition  $^{64}$  (CuAAC) with TEMPO-N<sub>3</sub> to yield *ex*TEMPO-TMV (Scheme 1d).

The TEMPO radical is known to be reduced in the presence of sodium ascorbate, so we modified a procedure initially described by Finn<sup>65</sup> by forming Cu<sup>I</sup>(THPTA) separately by mixing a 1:5:1 molar ratio of CuII sulfate, the ligand THPTA, and sodium ascorbate in water. Starting with Cu<sup>II</sup> sulfate as opposed to directly using a Cu<sup>I</sup> source to form the Cu<sup>I</sup>(THPTA) complex provided consistently high yields, likely because of the more favorable aqueous solubility of Cu<sup>II</sup> sulfate. This solution was transferred quickly in open air to an aqueous solution of TMV under N<sub>2</sub>. Degassing of solutions were not necessary, and the successful attachment of the new functional groups was again confirmed by HPLC/ESI-MS. As shown in Figure 1d, after the quantitative diazonium coupling reaction, all of the TMV coat protein monomers were modified to the azo adduct (17 666 Da) in line with previous literature reports.<sup>66</sup> The CuAAC reaction on the alkyne-TMV with TEMPO-N<sub>3</sub> proceeded quantitatively, evidenced by the loss of the alkyne peak at 17 666 Da. The morphology and dispersity of TMV were checked after each bioconjugation reaction and were found unchanged by transmission electron microscopy (TEM) and size exclusion chromatography (SEC), respectively (Figures 1a-c and S11).

Finally, EPR spectroscopy was used to characterize the presence of paramagnetic species on the TMV surface. As shown in the Figure 1e, small molecule TEMPO radicals

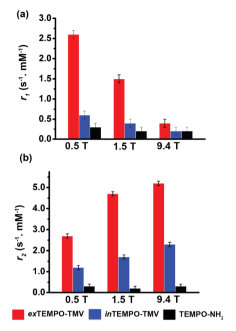




**Figure 2.** (a) Schematic representation of the *ex*TEMPO-TMV with one coordinated water molecule (inner-sphere water, its oxygen is colored black) in solution (bulk water, oxygens are red). Second-sphere water molecules (water oxygens are blue). The parameters that govern the relaxivity are also represented: NO–H distance (r), the mean lifetime  $(\tau_{\rm m})$  of the water molecule(s) in the inner sphere, the rotational correlation time  $(\tau_{\rm R})$ , and the electronic spin relaxation times  $(T_{\rm 1e}$  and  $T_{\rm 2e})$ . (b) Plot of  $1/T_1$  (s<sup>-1</sup>) versus TEMPO concentration (mM) for *ex*TEMPO-TMV, *in*TEMPO-TMV, and TEMPO-NH<sub>2</sub> at 23 MHz in 0.1 M pH 7.4 KP buffer and 310 K.

showed a characteristic  $^{14}$ N-triplet (I=1;  $A\sim45$  MHz) centered at a g-value of 2.007, which is typical of the nitrogencentered radical. The sharp line width and isotropic g/A-values are consistent with rotational averaging. By contrast, the exTEMPO-TMV (Figure 1e) and inTEMPO-TMV (Figure S12) results in significantly broader and more anisotropic EPR spectra. The distributed g/A-values and attenuated signal intensity are attributed to decreased rotational and translational mobility of the radical. This significantly increases the observed line width of the mobilized radical. High density of radicals can also result in dipolar spin—spin exchange, which also contributes to the increased line width.

**Proton Relaxation Properties of ORCA Loaded TMV Rods.** To characterize the relaxation behavior of the TEMPO-TMV particles, we performed proton relaxometric measurements at different fields. It is worth noting that nitroxide radicals normally bind noncovalently to at least one water molecule, <sup>67</sup> which is similar to that found with metal-based complexes with one or more water molecules coordinated to the inner sphere of the metal centers. <sup>34,68</sup> The overall proton relaxation behavior is determined by the sum of the inner-



**Figure 3.** Determined (a)  $r_1$  and (b)  $r_2$  relaxivities for the agents exTEMPO-TMV, inTEMPO-TMV, and TEMPO-NH<sub>2</sub> at different fields in 0.1 M pH 7.4 KP buffer and 310 K.

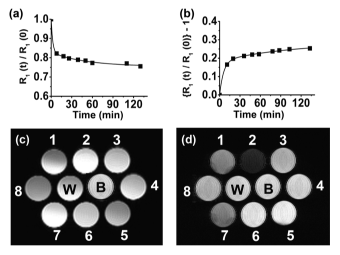


Figure 4. (a) Evolution of the relative water proton paramagnetic relaxation rate of a 0.14 mM aqueous solution of exTEMPO-TMV in the presence of 100-fold molar excess of ascorbate. Plot normalized to the initial value  $R_1^{(t)}/R_1^{(0)}$  as a function of the time (min) and measured at 23 MHz, 0.1 M pH 7.4 KP buffer, 310 K. (b) Evolution of the relative water proton paramagnetic relaxation rate of a 0.14 mM aqueous solution of exTEMPO-TMV (reduced) in the presence of 100-fold molar excess of KO<sub>2</sub>. Plot normalized to  $(R_1^{(t)}/R_1^{(0)}-1)$  as a function of the time (min) and measured at 23 MHz, 0.1 M pH 7.4 KP buffer, 310 K. The solid lines through the data points represent the pseudo-first-order reaction fits. (c)  $T_1$ -weighted and (d)  $T_2$ -weighted fast spin-echo images (TE = 200 ms) of (1) exTEMPO-TMV oxidized with KO<sub>2</sub>, (2) exTEMPO-TMV (0.14 mM/TEMPO), (3) exTEMPO-TMV reduced with ascorbate, (4) ascorbate in PBS, (5) TEMPO-NH<sub>2</sub> (1.4 mM), (6) TEMPO-NH<sub>2</sub> (0.14 mM), (7) inTEMPO-TMV (0.14 mM/TEMPO), (8) KO<sub>2</sub> in PBS, (W) water at pH 7.0, (B) PBS buffer at pH 7.0 taken at 9.4 T and 293 K.

sphere and outer-sphere contributions.<sup>34</sup> While the outer-sphere contributions cannot be neglected, the inner-sphere contributions are typically dominant.<sup>69</sup> For nitroxide radicals, under the extreme narrowing condition,<sup>70,71</sup> the primary inner-

Scheme 2. (a,b) inFITC-exTEMPO-TMV Bimodal Imaging Probe and (c) Diagram of TMV Showing the Relative Placement of the Fluorescent Dye on the Interior Residues and the ORCAs on the Exterior<sup>a</sup>

<sup>a</sup>The exterior surface was first modified with TEMPO radicals via diazonium coupling followed by a CuAAC reaction, and then, the interior surface of TMV was modified by EDC coupling with propargylamine followed by a CuAAC reaction.

sphere relaxation mechanism is determined by dipolar and Curie dipolar contributions. <sup>72,73</sup> However, at higher magnetic fields, line broadening predominantly originates from Curie spin relaxation processes. The dipolar and Curie mechanisms from the inner- and outer-sphere contributions can be modeled as described in previous detailed reports. <sup>73–75</sup> Large macromolecules, like the TMV presented here, are characterized by slow rotation dynamics in solution <sup>33,35</sup> commonly represented by the rotational correlation time ( $\tau_R$ ). Other factors that have a high impact on the relaxation processes for  $T_1$  and  $T_2$  are the electronic relaxation rates ( $1/T_{eij}$ , i=1,2) and solvent exchange lifetimes ( $\tau_m$ ) as defined by the correlation times ( $\tau_{cij}$ , i=1,2)

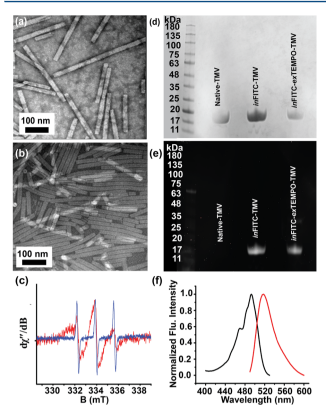
$$\tau_{ci} = \frac{1}{T_{ei}} + \frac{1}{\tau_{m}} + \frac{1}{\tau_{R}}, i = 1, 2$$
(1)

We can make several simplifying assumptions. In systems where  $T_{\rm e}$  is short, as is the case of TEMPO radicals  $(T_{\rm ei} \geq 20~{\rm ns})$ ,  $^{39}$  the molecular rotation and water exchange dominate  $\tau_{\rm ci}$ . The water exchange rate  $(k_{\rm ex}=1/\tau_{\rm m})$  for nitroxide radicals has been estimated to be quite slow  $(k_{\rm ex}\sim 2\times 10^{-4}~{\rm s})$ ,  $^{39}$  and it is expected from theory that the low-field total transverse relaxivity  $(T_2)$  will be small and similar to the longitudinal relaxivity, since it is modulated by  $T_{\rm e}$ . At higher magnetic fields, the  $r_2$  relaxivity is mostly affected by the square of the field (not  $r_1$ ) and increases with field strength. As shown in Table S2, exTEMPO-TMV, the most efficient system (Figure 2b), shows a significant increase in  $r_2$  at 9.4 T to 5.2 s<sup>-1</sup> mM<sup>-1</sup> per TEMPO compared to lower fields where  $r_1$  and  $r_2$  were similar.

Our data show that the TEMPO-TMV conjugates are efficient  $T_1$  and  $T_2$  contrast agents at low field and present

remarkable properties as  $T_2$  agents at high field, 35 with the greatest enhancement seen for exTEMPO-TMV (Table S2). Indeed, exTEMPO-TMV shows an enhancement of  $r_1$  and  $r_2$  at low field nearly 1 order of magnitude higher than that of the corresponding small molecule TEMPO-NH<sub>2</sub>, and at high-field, the  $r_2$  shows a 13-fold increase compared to that of TEMPO- $NH_2$ . A comparison of  $r_1$  and  $r_2$  values for exTEMPO-TMV and inTEMPO-TMV (Figure 3a,b) show that the exterior functionalization yields higher relaxivity values at all fields and both  $r_1$  and  $r_2$  relaxivities compare very favorably to existing polymeric ORCAs and small molecule contrast agents (Table S3). These data suggest that the TEMPO spins located on the inner cavity of the TMV macromolecule may have less access to bulk water preventing optimal exchange between the innersphere water molecules and those in the close vicinity of the TEMPO spin.

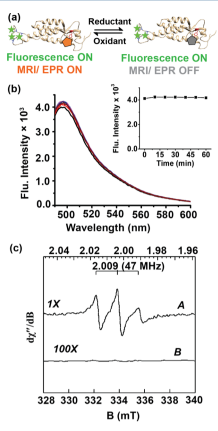
Nitroxide-based ORCAs typically undergo rapid reduction to diamagnetic hydroxylamines in the presence of reducing agents such as ascorbate. Relaxometric studies of exTEMPO-TMV, shown in Figure 4a, confirm that, in the presence of an equimolar amount of ascorbate, about 20% of the paramagnetic TEMPO spins on the exTEMPO-TMV are reduced within a few minutes. This was also confirmed by EPR spectroscopy (Figure S13a). As proof of principle that TEMPO-TMV compounds could be used as potent indicators of strong ROS, we performed the relaxometric experiments over the same time span, but in the presence of an excess of KO<sub>2</sub> at 0.4 T. In this case, the  $T_1$  of water protons recovered to nearly the original values (Figure 4b). This same reoxidation experiment was confirmed using EPR spectroscopy (Figure S13b).We inves-



**Figure 5.** Synthesis and characterization of *in*FITC-*ex*TEMPO-TMV; TEM images: (a) *in*FITC-TMV, (b) *in*FITC-*ex*TEMPO-TMV. (c) EPR spectrum of *in*FITC-*ex*TEMPO-TMV (red) and TEMPO-NH<sub>2</sub> (blue). SDS-PAGE to confirm that FITC has been successfully attached to TMV; SDS-PAGE was visualized by (d) Coomassie brilliant blue and (e) UV. (f) Excitation (black line) and emission (red) spectra of *in*FITC-*ex*TEMPO-TMV.

tigated the rate of reduction of the *ex*TEMPO-TMV under pseudo-first-order conditions using a 100-fold excess of ascorbate in pH 7.4 phosphate-buffered saline (PBS). Fits were obtained for relaxometric data collected over a 2 h period. Pseudo-first-order rate constants,  $k^\prime$ , were obtained by following the decay of the  $R_1^{(t)}/R_1^{(0)}$  determined by proton relaxometry. Excellent agreement with literature values was found for *ex*TEMPO-TMV and other macromolecules ( $k^\prime=17\times10^{-4}\pm0.1~\text{s}^{-1}$ ) as shown in the Table S4.  $^{33,35,39}$  Conversely, the oxidation of the reduced *ex*TEMPO-TMV hydroxylamine with an excess of KO<sub>2</sub> also showed a quite extensive reaction with a  $k^\prime=16\times10^{-4}\pm0.1~\text{s}^{-1}$ .

In Vitro MR Imaging. In support of these experimental observations, we have performed phantom imaging on several samples containing exTEMPO-TMV, inTEMPO-TMV, reduced exTEMPO-TMV, oxidized exTEMPO-TMV, and standards. T2-weighted MR imaging at 9.4 T showed that among the 10 tubes, only those containing the nonreduced TMV-TEMPOs showed enhanced  $T_2$  contrast, whereas the free TEMPO (Figure 4d, wells 5 and 6) do not show any contrast difference even when we doubled the concentration of free TEMPO sample (Figure 4d, well 6). While contrast enhancement is evident in the  $T_1$ -weighted images (Figure 4c, wells 8 vs 2), the greatest contrast at this field strength comes from the  $T_2$ -weighted images (Figure 4d, wells 8 vs 2). These results suggest that these TEMPO macromolecular derivatives can detect changes in superoxides at millimolar concentrations. It also shows the potential of using exTEMPO-TMV as a high-



**Figure 6.** Bimodal imaging characteristics of the *in*FITC-*ex*TEMPO-TMV redox probe. (a) Protein crystal structure of a single coat protein of TMV showing the attachment of FITC and TEMPO to the inner (glutamate) and outer (tyrosine) surfaces, respectively. (b) Time-dependent emission behavior of *in*FITC-*ex*TEMPO-TMV redox probe upon addition of 100 eq of ascorbate in 0.1 M pH 7.4 KP buffer. (c) EPR spectra of *in*FITC-*ex*TEMPO-TMV redox probe before (A) and 60 min after (B) addition of 100 eq of ascorbate in 0.1 M pH 7.4 KP buffer. An identical experiment on Cy5 was conducted, and details are in Figure S13.

field  $(T_2)$  contrast agent. This is particularly relevant, because most of clinical MR scanners operate at low fields (0.4–3 T), though implementation of clinical high-field scanners is progressing.

inFITC-exTEMPO-TMV for Bimodal Molecular Imaging. After confirming that exTEMPO-TMV exhibits better  $T_1$ and  $T_2$  relaxation characteristics compared to in TEMPO-TMV, this compound was used for all additional studies. A second goal of using TMV as a platform was to install a fluorophore for bimodal imaging in a position where it would not be quenched by the TEMPO radicals (Scheme 2). The order of the bioconjugation reactions to the TMV virus surface is important to minimize the redox chemistry on the attached TEMPO. We first functionalized the interior surface with FITC using standard CuAAC conditions and then functionalized the exterior surface with TEMPO radicals using our modified procedure. After each bioconjugation reaction, the integrity of the particle was characterized by SEC and TEM (Figure 5a,b). Moreover, TEMPO radical attachment to the exterior surface was confirmed by EPR spectroscopy (Figure 5c). The attachment of FITC to the interior surface of TMV was confirmed by UV-vis spectroscopy (Figure S14), SDS-PAGE (Figure 5d,e), and fluorescence spectroscopy (Figure 5f). To show this quenching process is not dependent on FITC, we

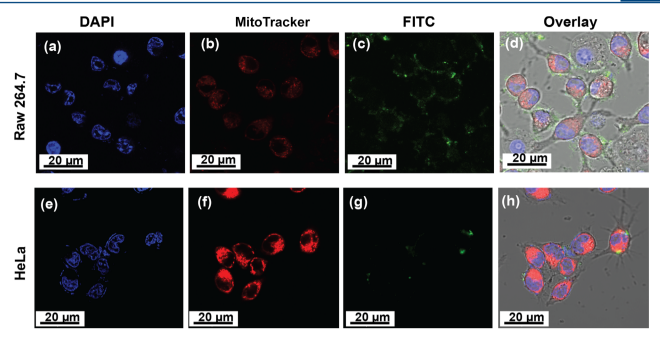
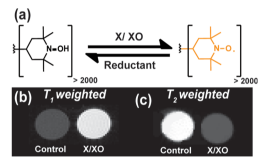


Figure 7. Confocal microscopy images of cellular uptake of *inFITC-red-exTEMPO-TMV* redox probe with (a–d) RAW 264.7 cells and (e,f) HeLa cells. Color code: blue, Hoechst 33342; red, MitoTracker Deep Red; green, *inFITC-red-exTEMPO-TMV*.



**Figure 8.** (a) Schematic showing the redox behavior of *red-ex*TEMPOTMV. (b)  $T_1$ -weighted phantom images and (c)  $T_2$ -weighted phantom images before and after addition of X/XO. The concentration of TMV sample was 5.0 mg mL<sup>-1</sup>, and the reaction mixture consisted of 50% of serum by volume.

attached a cyanine dye (Cy 5) to the interior as well, and we again observed minimal quenching from the radical (Figures S15-S17).

Fluorescein and cyanine dyes, like most fluorophores, are known to be quenched by nitroxyl radicals, <sup>33,42,80</sup> and the tube-like architecture of TMV allowed us to attach the MRI imaging agent and the fluorescent dye far enough apart to prevent quenching of the fluorescence emission of the dye (Figure 6a). To our satisfaction, we found less than 3% variation in fluorescence when shifting between the oxidized radical to the reduced hydroxylamine (Figure 6b) even after an hour following addition of 100 eq of ascorbate. On the other hand, the EPR spectra (Figure 6c) confirms that the radicals are completely reduced by the ascorbate under these same conditions. We again conducted the same experiment with *inCyS-exTEMPO-TMV* and confirmed that the oxidation state of the radical does not affect the emission intensity of Cy 5 (Figure S18), showing the generalizability of this approach with different dyes.

**In Vitro-***in***FITC-***red-ex***TEMPO-TMV Studies.** While the ultimate goal is in vivo imaging, in these early studies, we found

it important to ascertain their behavior in cells and in serum. Cellular uptake could potentially result in trafficking to organelles normally high in superoxide concentrations, such as mitochondria, and could create an unwanted background signal. Further, uptake by macrophages, which are known to efficiently remove macromolecules and nanoparticles, could potentially reduce the circulatory half-life of these particles and may lead to the formation of antibodies against our probe making their clearance faster.

Studies have shown that macrophages as well as HeLa cells interact less effectively with high aspect ratio nanoparticles, and TMV in particular has been extensively studied. 52,81,82 We were curious if we would see uptake with reduced inFITCexTEMPO-TMV (inFITC-red-exTEMPO-TMV) nanoparticles and if this would cause changes in the oxidation state of the ORCA following uptake. We used confocal microscopy on our "quenchless" system to investigate the interaction between inFITC-red-exTEMPO-TMV and macrophages. The confocal microscopy data shows that inFITC-red-exTEMPO-TMV (Figure 7d) associates with macrophages by adhering to the surface with no obvious fluorescence inside the cells and, in particular, the mitochondria, which are stained red in Figure 7b. We also investigated the cellular uptake of inFITC-redexTEMPO-TMV by HeLa cells (Figure 7e-g) and found no significant fluorescence signal coming from cells (Figure 7g). The lack of interaction with mitochondria was corroborated by EPR spectroscopy. To ensure we would observe signal if any trafficking to mitochondria occurs at all, we stimulated the HeLa and macrophages with the superoxide promotor phorbol 12-myristate 13-acetate (PMA)<sup>83–85</sup> and then incubated these cells with red-exTEMPO-TMV for an hour. Cells were collected, and EPR spectra were recorded as a function of time. A lack of an EPR signal over time confirms that redexTEMPO-TMV has not been uptaken by cells (Figure S21). If the red-exTEMPO-TMV had been taken up by the cells and trafficked to an area of ROS activity, we would expect to observe an increase in the EPR signal intensity with time, particularly when the cells were stimulated by PMA.<sup>83</sup> These

results are promising as it suggests that background signal from cellular metabolism is minimal.

red-exTEMPO-TMV Performance in Serum is very rich in a variety of proteins that may interfere with superoxide detection either by (i) directly reducing TEMPO to its EPR silent form or (ii) by covering the probe with a layer of protein—commonly called a protein corona—which would inhibit the "turn-on" response. To ascertain if our sensor functions in a biological environment, which might mimic detection of superoxide from tissue damage, for instance, we tested red-exTEMPO-TMV in serum solution using enzymatically produced superoxide. Xanthine/xanthine oxidase (X/XO) was employed as a superoxide generating agent, as it has been to be a good model of cell-free superoxide generation via EPR/MRI. First, the red-exTEMPO-TMV was made by reduction of exTEMPO-TMV with 0.1 M sodium ascorbate in a suspension of PBS, which was then purged with nitrogen gas to prevent reoxidation. Xanthine (final concentration of 0.05 mM) and red-exTEMPO-TMV (final concentration of 5.0 mg mL<sup>-1</sup>) were mixed in fetal bovine serum (FBS). Finally, xanthine oxidase (final concentration of 0.1 U/ mL) was added to the mixture. EPR spectra were recorded over 24 h, and MR phantom images were obtained from the same samples. Reoxidation of the probe was evident immediately. The  $T_1$ -weighted images and  $T_2$ -weighted images for the redexTEMPO-TMV sample with and without X/XO shows a huge contrast difference (Figure 8b,c) clearly indicating that the serum did not prevent detection of superoxide. These changes were confirmed by EPR, which also showed the regeneration of the TEMPO radical by the superoxide formed from X/XO (Figure S22) in the serum solution.

## CONCLUSIONS

This was a fundamental study of the synthesis, magnetic, and optical characterizations, and in vitro analyses of a new type of a fluorescent/MRI active superoxide sensor that works at both high and low magnetic fields. Our ORCA-conjugated VNPbased nanoparticle probe appears promising for use at both clinical (<3 T) and preclinical (>3 T) MRI fields. Attachment of ORCA to TMV rods resulted in a 10-fold increase in  $r_1$  and a 13-fold increase in  $r_2$  compared to that of the small molecule ORCAs. We have proposed that these increases in relaxivity are ascribed to a decrease in the molecular rotation and diffusional motions brought about by attachment of the ORCA to the large rod-shaped particle. The distinct "inner" and "outer" topology of the TMV permits the creation of a "quenchless" ORCA-based bimodal probe for combined MRI and fluorescence imaging. The probe shows "turn-on" functionality toward enzymatically produced superoxide in the presence of serum, yet does not show appreciable changes in EPR intensity even when endocytosed by cells, suggesting this probe would be ideal for imaging diffuse tissue injury as may be the case in deep tissue diabetic ulceration or coronary heart disease.

## ASSOCIATED CONTENT

#### **S** Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.molpharmaceut.8b00262.

Detailed experimental and synthetic procedures and characterization of small molecules, TMV isolation and

purification protocols, bioconjugation protocols, electron paramagnetic studies, instrumentation (PDF)

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#### **Author Contributions**

\*M.D. and A.F.M. contributed equally to this manuscript. All authors have given approval to the final version of the manuscript. M.D., A.F.M., and J.J.G. designed and conducted the experiments; A.F.M. and A.D.S. helped with MRI studies; Z.C. conducted the cell imaging experiments. M.D., P.M.P., and B.S.P. conducted EPR studies; R.P.W. and S.L. performed TEM imaging; R.P.W. synthesized some precursor compounds. M.A.L. helped with TMV isolations. All authors provided edits to the final manuscript.

#### Notes

The authors declare no competing financial interest.

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