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### Title

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### Permalink

https://escholarship.org/uc/item/5004v69h

Journal Nano Letters, 24(26)

ISSN

1530-6984

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Publication Date 2024-07-03

### DOI

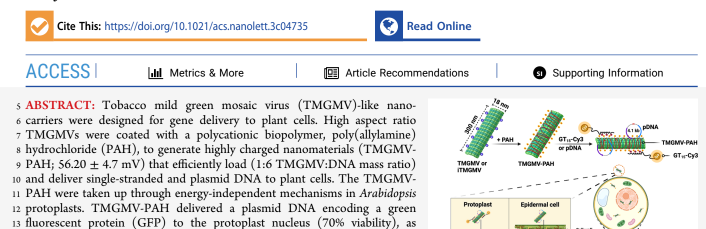
10.1021/acs.nanolett.3c04735

Peer reviewed

pubs.acs.org/NanoLett

## DNA Delivery by Virus-Like Nanocarriers in Plant Cells

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14 evidenced by GFP expression using confocal microscopy and Western blot 15 analysis. TMGMV-PAH were inactivated (iTMGMV-PAH) using UV cross-16 linking to prevent systemic infection in intact plants. Inactivated iTMGMV-17 PAH-mediated pDNA delivery and gene expression of GFP in vivo was

18 determined using confocal microscopy and RT-qPCR. Virus-like nano-

19 carrier-mediated gene delivery can act as a facile and biocompatible tool for advancing genetic engineering in plants.

20 **KEYWORDS**: virus, nanoparticles, gene delivery, protoplasts, plant genetics, agriculture

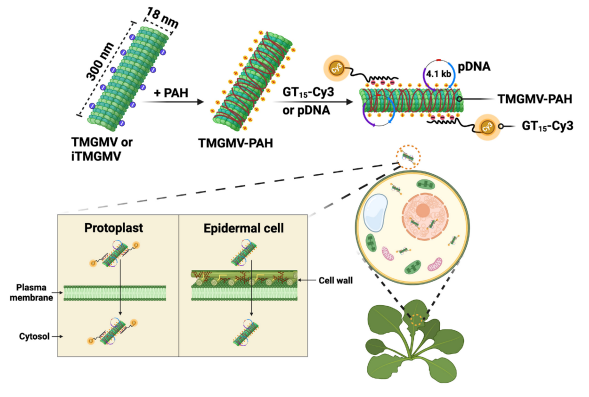
he rapid increase in the human global population is 21 projected to require a 35 to 55% increase in food 23 production by 2050.<sup>1</sup> Addressing this challenge during a 24 changing climate and without sustainable conventional 25 agricultural practices raises concerns about food security.<sup>2</sup> 26 Plant genetic engineering has been widely employed to 27 generate crops with increased yield,<sup>3</sup> improved quality, 28 enhanced resistance to herbicides,<sup>4</sup> insects,<sup>5</sup> diseases,<sup>6,7</sup> and 29 biotic and abiotic stresses.<sup>8,9</sup> Genetically modified plants for 30 biomanufacturing also hold immense potential for synthesizing 31 small-molecule drugs,<sup>10</sup> recombinant protein therapeutics,<sup>11,12</sup>
 32 and vaccines.<sup>13,14</sup> Despite numerous biotechnological advance-33 ments over the past few decades, the genetic transformation of 34 many plant species still poses considerable challenges. The 35 delivery of transgenes into plant species mainly relies on two 36 transformation methods: Agrobacterium tumefaciens-mediated 37 transformation system<sup>15</sup> and particle bombardment.<sup>16</sup> How-38 ever, the Agrobacterium-mediated system has some significant 39 drawbacks such as uncontrollable target gene integration into 40 the host chromosomes causing positional effects on gene 41 expression, and many plant species are inherently resistant to 42 Agrobacterium infection<sup>17</sup> or showed low transformation 43 efficiency (~5% to 33%).<sup>18,19</sup> Biolistics has been utilized in 44 various plant species, as a random gene delivery system into 45 the host nucleus, mitochondria, and chloroplast.<sup>4</sup> Particle 46 bombardment is performed by high-pressure gene gun delivery 47 that damages host genomic DNA and results in random

insertions of multiple copies of the gene<sup>20</sup> The particle 48 bombardment system is also expensive, requires labor-intensive 49 tissue culture and selection, has low transformation efficiency 50 often requiring hundreds of transformation attempts to 51 generate a transgenic line,<sup>20,21</sup> and has not been successfully 52 implemented in diverse plant species.<sup>22</sup> Therefore, there is a 53 pressing need for a versatile, plant-species-independent, and 54 easy-to-use tool for plant genetic transformation, allowing for 55 efficient delivery of exogenous genes.

Recent advancements in nanotechnology have revealed the 57 potential of nanomaterials in facilitating the delivery of genetic  $_{58}$  materials, such as plasmid DNA<sup>23-25</sup> and siRNA,<sup>26,27</sup> as well as  $_{59}$ biomacromolecules like functional proteins,<sup>28</sup> active ingre- 60 dients,<sup>29,30</sup> nutrients,<sup>31</sup> and therapeutics<sup>32</sup> in plants. Single- 61 walled carbon nanotubes (SWCNTs),<sup>23,24,33</sup> mesoporous silica 62 nanoparticles (MSNs),<sup>34,35</sup> layered double hydroxide (LDH) 63 clay nanosheets,<sup>26</sup> and functional peptide-DNA com- 64 plexes<sup>25,36</sup> have demonstrated delivery of functional DNA/ 65 RNA cargoes into plant cells without mechanical assistance. 66

Received: December 4, 2023 Revised: June 10, 2024 Accepted: June 12, 2024





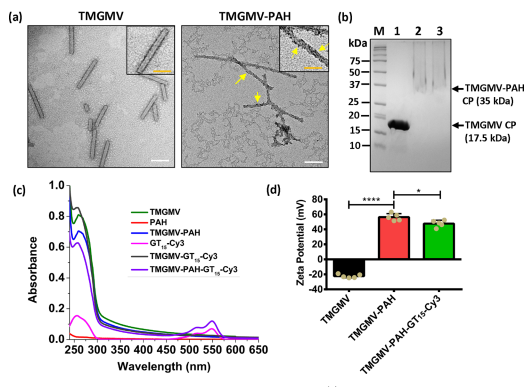
**Figure 1.** Intracellular DNA delivery in *Arabidopsis* plant cells mediated by virus-like nanocarriers. Negatively charged TMGMVs or inactivated (iTMGMVs) were coated with a biopolymer, poly(allylamine) hydrochloride (PAH), imparting them with positive charge (TMGMV-PAH). The TMGMV-PAH were loaded by electrostatics with a DNA oligo ( $GT_{15}$ , 30 bp ssDNA) that was covalently linked to a Cy3 organic dye (TMGMV-PAH-GT<sub>15</sub>-Cy3), or a plasmid DNA (pDNA) encoding a reporter gene of a green fluorescent protein (GFP). The nanocarriers and DNA cargoes spontaneously enter plant cell membranes without mechanical aid through energy-independent uptake mechanisms. Inactivated iTMGMV-PAH mediated the delivery and expression of pDNA in *Arabidopsis* epidermal cells.

67 Several studies have demonstrated the possibility of carbon 68 nanotube-mediated gene delivery in plant nucleus,<sup>23,36</sup> 69 chloroplast,<sup>24,33</sup> and mitochondrial<sup>25</sup> genomes. However, 70 there is a need to develop high aspect ratio nanomaterials 71 for plant transformation that are degradable, biocompatible, 72 and manufactured with controlled aspect ratios on a large scale. 73 We turned toward plant virus nanoparticles as a biodegradable, 74 cost-effective, and easily scalable nanotechnology with tunable 75 surface chemistry.<sup>29,30,37</sup>

Tobacco mild green mosaic virus (TMGMV)<sup>38</sup> is a plant 76 77 virus within the tobamovirus genus, also known as the U2 78 strain of tobacco mosaic virus (TMV), approved by the U.S. 79 Environmental Protection Agency (EPA) for use in bio-80 herbicides.<sup>39</sup> The nucleoprotein components of TMGMV are 81 self-assembled from 2130 identical copies of a coat protein and  $_{82}$  ssRNA to form a 300  $\times$  18 nm soft matter rod-shaped 83 structure with a 4 nm wide hollow interior channel.<sup>29,38,40</sup> The 84 nanocarriers derived from TMGMV are of interest for delivery 85 applications due to their unique physio-chemical properties, 86 such as biodegradability (protein-based particles), the ability to 87 self-assemble into identical and high aspect ratio structures, and large-scale economical production with high purity and 88 89 reproducibly.<sup>29,41</sup> The chemical design space is well understood and TMGMV can be functionalized with cargo through 90 91 covalent chemistry<sup>42</sup> or encapsulation.<sup>29</sup> There are also well-92 established methods of TMGMV RNA inactivation through 93 UV cross-linking or chemical treatments for use in plant <sup>94</sup> species susceptible to infection.<sup>43</sup> TMGMV particles have been 95 utilized as a carrier for active ingredients such as a porphyrin-96 based photosensitizer drugs (500 Zn-porphyrin molecules/

TMGMV) for cancer cell abolition of melanoma and cervical 97 cancer models,<sup>40</sup> as well as ivermectin (10% mass loading 98 efficiency to TMGMV) to treat plants infected with parasitic 99 nematodes.<sup>29,30,44</sup> Plant virus-derived vectors (plasmids with 100 virus genetic elements) have been extensively used for genetic 101 engineering in plants through the mechanical inoculation of 102 plasmid DNA, biolistics, vascular puncture, agroinoculation, or 103 insect-mediated vector delivery.<sup>45,46</sup> These applications 104 focused on delivery of RNA packaged inside the capsid.<sup>47</sup> To 105 date, plant virus coat proteins have not been engineered as 106 carriers for facile plasmid DNA delivery in plant cells.

In this study, we developed native and inactivated TMGMV- 108 based nanomaterials as a platform for the nuclear delivery of 109 DNA in Arabidopsis thaliana protoplasts and intact plants, 110 respectively (Figure 1). Although PEG-mediated protoplast 111 fl transformations achieve high transient transformation efficien- 112 cies (50–90% in viable cells),<sup>48</sup> protoplast systems are crucial 113 for developing genetic transformation tools and understanding 114 nanoparticle-plant cell interaction processes.<sup>23,33,49</sup> Because 115 plant protoplasts lack a cell wall, this study also included DNA 116 delivery analysis in vivo using Arabidopsis leaf epidermal cells. 117 We functionalized TMGMV by covalently coating a poly- 118 cationic biopolymer, poly(allylamine) hydrochloride (PAH), 119 on the TMGMV surface (TMGMV-PAH). The PAH imparts a 120 positive charge to TMGMV-PAH for binding to DNA through 121 electrostatic interactions. PAH has been extensively used for 122 pharmaceutical and drug delivery applications due to its high 123 water-solubility and biodegradable properties.<sup>50,51</sup> To deter- 124 mine whether TMGMV-PAH delivered single-stranded DNA 125 (ssDNA) into protoplast cells without using mechanical aid 126



**Figure 2.** Characterization of TMGMV-PAH coated with single-stranded DNA. (a) Transmission electron microscopy of negative-stained TMGMV and TMGMV-PAH. Yellow arrows indicate PAH coated on the surface of TMGMV. Scale bars 100 nm. (b) Denaturing Nu-PAGE gel electrophoresis under white light followed by Coomassie staining, 1: TMGMV, 2: TMGMV-PAH, 3: TMGMV-PAH-GT<sub>15</sub>-Cy3, M: prestained molecular weight standards. The arrow indicates the position of the TMGMV coat protein (CP) at 17.5 kDa (lower arrow) and PAH conjugated TMGMV-PAH CP at 35 kDa (upper arrow) or higher molecular weight. (c) UV–vis absorbance and (d) zeta potential (10 mM MES, pH 6.0) of TMGMV before and after coating with PAH and GT<sub>15</sub>-Cy3. The data are the means  $\pm$  SD (n = 4). Statistical analysis was performed by one-way analysis of variance (ANOVA) with Tukey's *posthoc* multiple comparison analysis (GraphPad Prism 6); \*P < 0.05; \*\*\*\*P < 0.0001.

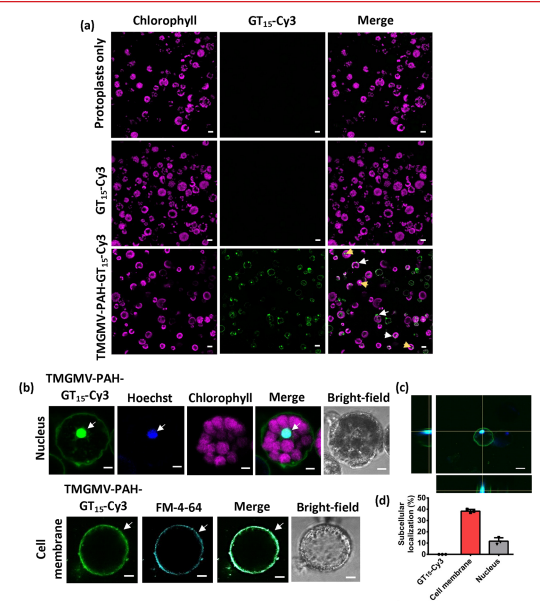
127 while maintaining biocompatibility, we employed confocal 128 microscopy to track the ssDNA cargo covalently bonded to a 129 fluorophore (Cy3) and protoplast bioavailability assays. We 130 also demonstrated the high loading capacity of plasmid DNA 131 (pDNA) onto the TMGMV-PAH, and assessed the pDNA 132 delivery, uptake mechanism, and transgene expression in 133 protoplasts. Finally, we used inactivated iTMGMV-PAH to 134 demonstrate pDNA delivery and expression in *Arabidopsis* leaf 135 epidermal cells *in vivo*. Using virus-like nanocarriers for DNA 136 delivery in plant cells offers a promising solution for plant 137 genetic transformations that is scalable and biocompatible with 138 high manufacturing quality and reproducibility.

The selection of polymer coating for TMGMV focused on 139 140 cationic biopolymers capable of binding electrostatically with 141 negatively charged pDNA. Among various options, PAH, polylysine, and polyarginine were prioritized due to their 142 143 higher  $pK_a$  values (above pH 8) and FDA approval for other 144 applications. TMGMV coated with polylysine and polyarginine 145 were negatively charged, making them unsuitable for pDNA 146 coating (Figure S1). In contrast, PAH TMGMVs were positively charged, and therefore, PAH was chosen as the 147 148 coating for TMGMV in this study. We characterized TMGMV, 149 TMGMV-PAH, and GT<sub>15-</sub>Cy3-loaded TMGMV-PAH 150 (TMGMV-PAH-GT<sub>15</sub>-Cy3) by UV-vis, dynamic light scatter-151 ing (DLS), zeta potential ( $\zeta$ ), transmission electron micros-152 copy (TEM), Nu-PAGE protein analysis, and fluorescence 153 emission spectra. TEM imaging of TMGMV and TMGMV-154 PAH shows high aspect ratio, rod-shaped nanostructures 155 (Figure 2a) consistent with previous studies using TMGMV 156 for pesticide delivery.<sup>29,42</sup> The TMGMV-PAH had a rough

surface, which is different from native TMGMV (Figure 2a), 157 indicating coating of the PAH polymer on the TMGMV 158 surface. We utilized a carbodiimide coupling reaction to 159 covalently bond the amine functional groups of PAH to the 160 carboxyl groups in TMGMV (Figure S2),<sup>42</sup> and the chemical 161 conjugation was confirmed by Fourier-transform infrared 162 spectroscopy (FTIR; Figure S3). Based on TEM analysis, the 163 average lengths of TMGMV and TMGMV-PAH were 164 nonsignificantly different,  $129.9 \pm 57.7$  and  $191.3 \pm 95$  nm, 165 respectively. Notably, broken nanomaterials were also observed 166 in both uncoated TMGMV and TMGMV-PAH, which can 167 occur during preparation or imaging of the TMGMV TEM 168 samples.<sup>29,42</sup> Furthermore, the conjugation of PAH (~17.5 169 kDa) to TMGMV coat protein (CP) was confirmed by 170 denatured Nu-PAGE protein analysis, which indicated the 171 presence of higher molecular weight bands at ~35 kDa, in 172 addition to the TMGMV CP band at ~17.5 kDa (Figure 2b). 173 The smeared protein bands were observed due to the high 174 positive charge of TMGMV-PAH CP (56.20  $\pm$  4.7 mV) that 175 hinders the relative mobility toward the electrode in the Nu- 176 PAGE system. Both TEM and Nu-PAGE analysis indicate that 177 PAH is coated onto the TMGMV-PAH. 178

To investigate DNA delivery by TMGMV-PAH in 179 protoplasts, we used confocal microscopy to track ssDNA 180 oligonucleotide  $(GT)_{15}$  covalently linked to the Cy3 181 fluorescent dye  $(GT_{15}$ -Cy3). Cy3 is bright, photostable, and 182 its emission range does not overlap with chloroplast 183 autofluorescence.<sup>24</sup> GT<sub>15</sub>-Cy3 has been previously employed 184 for coating positively charged carbon nanotubes for determin-185 ing subcellular localization in plants.<sup>24,33,52</sup> The UV–vis 186

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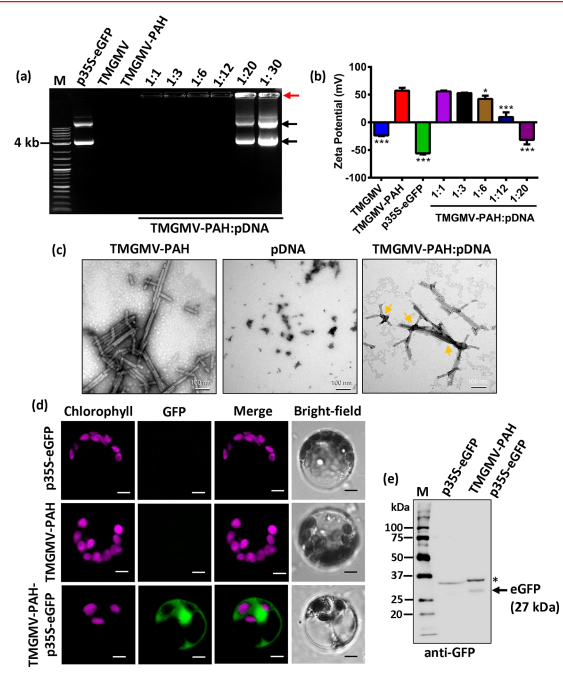


**Figure 3.** Delivery of single-stranded DNA by TMGMV-PAH in plant protoplasts. (a) Confocal images of isolated mesophyll protoplasts with chlorophyll autofluorescence (magenta) exposed to TMGMV-PAH-GT<sub>15</sub>-Cy3 (0.1 mg/mL). The GT<sub>15</sub>-Cy3 was detected in protoplast membranes (white arrows) and nuclei (yellow arrows). Scale bars 30  $\mu$ m. (b) After treatment with TMGMV-PAH-GT<sub>15</sub>-Cy3, protoplasts were stained either with a nuclear marker, Hoechst, or cell membrane staining dye, FM-4-64 for confocal microscopy imaging. Scale bars 5  $\mu$ m. (c) Orthogonal projections from z-stacks of different planes (x/y, x/z, or y/z) of confocal microscopy images indicating localization of GT<sub>15</sub>-Cy3 with Hoechst nuclear marker and FM-4-64 cell membrane dye. The data are means  $\pm$  SD (n = 3).

187 absorbance spectra of TMGMV, TMGMV-PAH, and 188 TMGMV-PAH-GT<sub>15</sub>-Cy3 indicated characteristic absorption 189 peaks at 260 nm (Figure 2c). TMGMV-PAH-GT<sub>15</sub>-Cy3 190 showed distinct absorption peaks at 550 nm that corresponded 191 to the Cy3 dye on TMGMV-PAH (Figure 2c). To validate the 192 binding of GT<sub>15</sub>-Cy3 to TMGMV-PAH and confirm the 193 absence of unbound dye, the sample was purified using a centrifugal filter unit (100 K MWCO). Following the second 194 195 wash step, no absorbance corresponding to GT<sub>15</sub>-Cy3 was 196 detected in the eluent (Figure S4a), whereas TMGMV-PAH-197 GT<sub>15</sub>-Cy3 exhibited fluorescence emission peaks at 567 nm, 198 attributed to the attachment of GT<sub>15</sub>-Cy3 on TMGMV-PAH 199 (Figure S4b). DLS analysis indicated well dispersed nanoma-200 terials with increasing hydrodynamic diameter from  $267 \pm 1.6$ 201 nm for TMGMV to 310  $\pm$  1.3 nm for TMGMV-PAH and 361

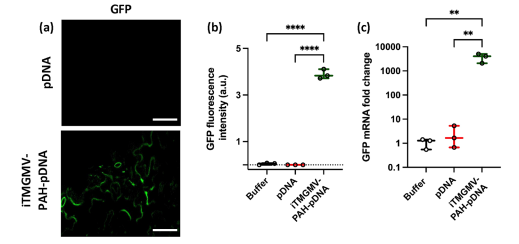
 $\pm$  3.2 nm for TMGMV-PAH-GT<sub>15</sub>-Cy3 (P < 0.005; Figure 202 S4c). We observed a significant change of  $\zeta$  potential after 203 conjugation of PAH from negative charged TMGMV (-22.37 204  $\pm$  2.3 mV) to highly positive charged TMGMV-PAH (56.20  $\pm$  205 4.7 mV; P < 0.0001; 10 mM MES buffer, pH 6.0; Figure 2d), 206 indicating binding of polycationic PAH to the TMGMV 207 surface. As expected, the  $\zeta$  potential for TMGMV-PAH slightly 208 decreased from 56.20  $\pm$  4.7 to 47.69  $\pm$  4.4 mV when loading 209 GT<sub>15</sub>-Cy3 (P < 0.05; Figure 2d) due to the electrostatic 210 bonding between the negatively charged GT<sub>15</sub> and the 211 positively charged TMGMV-PAH. 212

To examine *in vitro* DNA delivery and subcellular local- 213 ization in plant cells using TMGMV-PAH as a nanocarrier, 214 *Arabidopsis* protoplasts were isolated and incubated with 215 TMGMV-PAH coated with GT<sub>15</sub>-Cy3. Protoplasts are model 216



**Figure 4.** Plasmid DNA delivery and expression mediated by virus-like nanocarriers in isolated plant protoplasts. (a) DNA loading analysis by agarose gel electrophoresis of pDNA (p35S-eGFP) bound to TMGMV-PAH at mass ratios 1:1 to 1:30. M: DNA ladder. Black arrows indicate supercoiled (below) and circular (upper) pDNA bands. The red arrow indicates pDNA bound to TMGMV-PAH that prevents its mobility through the gel. (b) Zeta potential measurements of virus-like nanocarriers with or without pDNA (10 mM MES, pH 6.0). Data are means  $\pm$  SD (n = 3-4). Statistical analysis was performed by one-way ANOVA and *Dunnett's* multiple comparisons posthoc test; \*P < 0.01; \*\*\*P < 0.0001. (c) Representative TEM images of TMGMV, TMGMV-PAH, and pDNA-loaded at 1:6 mass ratios to TMGMV-PAH. Scale bar 100 nm. Arrows indicate pDNA attachment to TMGMV-PAH. (d) pDNA delivery and expression mediated by TMGMV-PAH in isolated plant protoplasts determined by confocal microscopy. Scale bar 10  $\mu$ m. (e) GFP expression analysis by Western blotting. The arrow indicates 27 kDa of GFP protein and asterisks indicate nonspecific bands. M, protein ladder.

<sup>217</sup> systems for gene expression analysis that have been used in <sup>218</sup> numerous plant nanoparticle studies of uptake and gene <sup>219</sup> delivery.<sup>23,33,49</sup> To assess the delivery of  $GT_{15}$ -Cy3 bound to <sup>220</sup> TMGMV-PAH and their subcellular localization using <sup>221</sup> confocal microscopy, isolated protoplasts (Figure S5) were <sup>222</sup> incubated with 0.1 mg/mL of TMGMV-PAH-GT<sub>15</sub>-Cy3 at <sup>223</sup> room temperature for 2 h before imaging. Confocal <sup>224</sup> fluorescence microscopy images indicated a significant level of GT<sub>15</sub>-Cy3 fluorescence signal in protoplast cell membranes, <sup>225</sup> and nuclei when treated with TMGMV-PAH-GT<sub>15</sub>-Cy3 <sup>226</sup> (Figure 3a). In contrast, control confocal images of protoplasts <sup>227</sup> f3 treated with GT<sub>15</sub>-Cy3 did not show GT<sub>15</sub>-Cy3 fluorescence <sup>228</sup> signal indicating that GT<sub>15</sub>-Cy3 alone cannot be taken up by <sup>229</sup> protoplasts under these exposure conditions (Figure 3a). To <sup>230</sup> confirm TMGMV-PAH-GT<sub>15</sub>-Cy3 interaction with protoplast <sup>231</sup> cell membranes and GT<sub>15</sub>-Cy3 nuclear delivery by TMGMV- <sup>232</sup>



**Figure 5.** Plasmid DNA delivery and expression mediated by iTMGMV-PAH-pDNA in *Arabidopsis* leaves. Green fluorescence protein (GFP) (a) confocal microscopy images and (b) and fluorescence intensity (n = 3) indicating GFP expression in leaf epidermal cells infiltrated with iTMGMV-PAH-pDNA. Three-week-old *Arabidopsis* leaves were abaxially infiltrated with (1:6) 0.1 mg/mL iTMGMV-PAH: 0.6 mg/mL pDNA and analyzed 2 days post infiltration (n = 3). Scale bars 30  $\mu$ m. One-way ANOVA with Tukey's *posthoc* multiple comparison analysis; \*\*\*\*P < 0.0001. (c) RT-qPCR analysis of GFP mRNA expression levels after 2 days of iTMGMV-PAH-pDNA infiltration in *Arabidopsis* leaves. Statistical analysis was performed by one-way ANOVA with Tukey's *posthoc* multiple comparison analysis; \*\*P < 0.005 (n = 3).

233 PAH, protoplasts were stained with a cell membrane marker 234 FM-4-64 and a nuclear staining marker Hoechst. The GT<sub>15</sub>-235 Cy3 fluorescence was observed localized with FM-4-64 and 236 Hoechst fluorescence signals in protoplasts cell membrane and 237 nucleus, respectively (Figures 3b and S6). Orthogonal 238 projections from Z-stacks of different planes (x/y, x/z, or y/z)239 z) of the confocal microscope images confirmed nuclear 240 uptake of GT<sub>15</sub>-Cy3 using TMGMV-PAH as shown by the 241 colocalization with Hoechst fluorescence dye (Figure 3c). 242 Quantitative subcellular localization analysis indicated that 243 approximately  $38\% \pm 1.5$  of the GT<sub>15</sub>-Cy3 fluorescence signal 244 was observed in protoplast cell membranes, while  $11\% \pm 3.0$ 245 localized with a nuclear marker (Hoechst; Figure 3d). 246 Together, our results indicate that high aspect ratio and highly 247 positive charged TMGMV-PAH allow penetration through 248 plant cell membranes and facilitate ssDNA delivery (GT15-249 Cy3) into the nucleus, similar to inorganic high aspect ratio 250 nanomaterials with positive charge.<sup>23</sup>

To elucidate the mechanism of DNA delivery into plant cells 251 252 by TMGMV-PAH, we conducted a cell uptake assay with 253 TMGMV-PAH-GT<sub>15</sub>-Cy3 at 4 °C to inhibit energy-dependent 254 uptake mechanisms, including endocytosis.<sup>53</sup> We observed a similar percentage of protoplasts with GT<sub>15</sub>-Cy3 delivery by 255 256 TMGMV-PAH at 4 °C (10%  $\pm$  1.6) and 25 °C (11%  $\pm$  3.2) (Figure S7). Thus, DNA delivered by TMGMV-PAH passively 257 258 traverses the protoplast membrane by an energy-independent 259 mechanism. This is consistent with previous studies demon-260 strating that highly charged inorganic nanomaterials sponta-261 neously penetrate plant cells, by creating temporary pores in 262 their lipid membranes.<sup>23,24,33,54,55</sup> To determine the specific 263 endocytosis pathways involved in nanoparticle uptake, a variety 264 of endocytosis inhibitors can be employed.<sup>56</sup> However, 265 temperature dependent assays block all endocytosis pathways, 266 thus giving unequivocal evidence that the nanocarriers are not taken up through energy dependent mechanisms. 267

268 We investigated the TMGMV-PAH loading of pDNA, 269 encoding a green fluorescent protein (GFP) in a transient 270 expression vector (p35S-eGFP) (Figure S8), and delivery in 271 *Arabidopsis* protoplasts. The TMGMV-PAH-pDNA were 272 loaded at various concentrations of pDNA (TMGMV- PAH:pDNA mass ratios 1:1 to 1:20 w/w). The gel 273 electrophoresis of pDNA mobility shift assay (EMSA) showed 274 no unbound or free pDNA running into the agarose gel at a 275 mass ratio of TMGMV-PAH/pDNA = 1:1 to 1:12 (w/w), 276 meaning that pDNA loading was 100% up to a 1:12 (w/w) 277 mass ratio (Figure 4a). The 1:12 TMGMV-PAH to pDNA 278 f4 mass loading ratio is multiple times higher than the 1:2 and 279 10:1 nanomaterial/pDNA loading ratio reported in previous 280 studies using inorganic nanomaterials for DNA delivery in 281 plant cells.<sup>23,57</sup> Oversaturated and unbound free pDNA bands 282 were observed at TMGMV-PAH/pDNA mass ratios of 1:20 283 (w/w) and higher in EMSA (Figure 4a). The loading of pDNA 284 gradually reduced  $\zeta$  potential as the loading ratio of pDNA 285 increased from 1:1 to 1:12 (Figure 4b) due to the electrostatic 286 bonding between the negatively charged pDNA and the 287 positively charged TMGMV-PAH. The highest decrease in  $\zeta$  288 potential was observed after pDNA loading to TMGMV-PAH 289 at a mass ratio of 1:12, dropping from the initial  $+57.53 \pm 5.2$  290 mV for TMGMV-PAH to  $+9.57 \pm 10.6$  mV (P < 0.0001; 291 Figure 4b). At the loading mass ratio of 1:20, the  $\zeta$  potential 292 became negative,  $-31.17 \pm 6.4$  mV, representing the 293 oversaturation of the nanocarriers and free pDNA in the 294 suspension. This finding indicates maximum pDNA loading at 295 a 1:12 mass ratio and is consistent with our EMSA analysis. We 296 confirmed morphological integrity of TMGMV-PAH loaded 297 with pDNA from 1:1 to 1:12 mass ratios by TEM (Figures 4c 298 and S9). In addition, we also assessed pDNA stability by an in 299 vitro pDNA degradation assay using DNase I (nuclease), which 300 showed that pDNA molecules, when loaded onto TMGMV- 301 PAH, were protected from DNase I nuclease activity (Figure 302 S10). 303

To demonstrate pDNA delivery and expression in plant 304 cells, we incubated isolated protoplasts with TMGMV-PAH- 305 pDNA complexes at 1:6 mass ratio having a high positive 306 charge (+42.16  $\pm$  5.1 mV) and loading of pDNA (Figure 4b) 307 to promote uptake through lipid membranes<sup>49</sup> and increase the 308 amount of pDNA delivery, respectively. We used 25  $\mu$ g of 309 pDNA for TMGMV-PAH-mediated protoplast transformation, 310 a standard concentration of pDNA (5–30  $\mu$ g) established for 311 PEG-mediated protoplast transformation.<sup>58</sup> Therefore, we 312

313 adjusted the TMGMV-PAH concentration to 0.04 mg/mL to 314 keep a 1:6 mass ratio of the pDNA loading. Protoplasts were 315 incubated with TMGMV-PAH-pDNA, and gene expression 316 was determined after 24 h by confocal fluorescence microscopy 317 imaging. We observed GFP expression in protoplasts when 318 incubated with TMGMV-PAH-pDNA (Figure 4d) at a 16%  $\pm$ 319 3.0 (P < 0.001) transformation efficiency. This transformation 320 efficiency is lower than what is reported for PEG-mediated 321 transformation in Arabidopsis plant protoplasts (50% to 90% in 322 viable cells).<sup>48</sup> However, this demonstrates that virus-like 323 nanocarriers can be engineered to deliver DNA to the plant 324 nuclear genome. Further optimization of plant virus type or the 325 nanocarrier charge, size, and aspect ratio properties may result 326 in higher transformation efficiencies. Nevertheless, GFP expression was observed using TMGMV-PAH-pDNA, but 327 328 was not detected when protoplasts were incubated with pDNA 329 alone and TMGMV-PAH alone (Figure 4d). To further 330 confirm GFP expression in protoplasts treated with TMGMV-331 PAH-pDNA, we performed a Western blot analysis on total 332 soluble protein using an anti-GFP antibody, which detected an  $\sim$ 27 kDa GFP-specific protein band (Figure 4e). 333

For GFP expression analysis in vivo, we inactivated TMGMV 334 335 to prevent plant infection using UV light exposure as reported 336 previously.<sup>43</sup> The TEM size of inactivated iTMGMV (110.73  $_{337} \pm 30.22$  nm) is similar to those of active TMGMV (129.9  $\pm$ 338 57.7 nm) (P > 0.05) (Figure S11). In contrast, the zeta 339 potential of iTMGMV is more negative  $(-36.29 \pm 4.23 \text{ mV})$ 340 compared to that of active TMGMV ( $-22.4 \pm 2.3 \text{ mV}$ ) (10 341 mM MES Buffer, pH 6.0) (P < 0.0001). This resulted in  $_{342}$  iTMGMV-PAH-pDNA having a higher zeta potential (58.53  $\pm$ 343 0.50 mV) than TMGMV-PAH-pDNA (42.16  $\pm$  5.1 mV; P < 344 0001). We abaxially infiltrated the inactivated iTMGMV-PAH 345 coated in pDNA into 3-week-old Arabidopsis leaves at the 346 previously established 1:6 mass loading ratio. Confocal 347 microscopy analysis indicated that 0.1 mg/mL of iTMGMV-348 PAH bound to 0.6 mg/mL of pDNA can enable GFP 349 expression into leaf epidermal cells (Figure 5a). Buffer or 350 iTMGMV-PAH infiltrated leaves did not exhibit GFP 351 fluorescence (Figure S12). Leaves infiltrated with iTMGMV-352 PAH-pDNA had a high GFP fluorescence intensity (Figure 353 5b). RT-qPCR analysis quantifying GFP mRNA fold change 354 expression supported GFP expression mediated by 0.1 mg/mL 355 iTMGMV-PAH:0.6 mg/mL pDNA (Figure 5c). Together, 356 these analyses show that (i)TMGMV-PAHs have the highest 357 pDNA mass loading ratio for nanocarriers reported to date, 358 preserve and protect the pDNA integrity from degradation, 359 and facilitate spontaneous pDNA translocation across the plant 360 plasma membrane and cell wall, enabling transgene expression 361 in the nucleus in vitro and in vivo.

Maintaining cell viability after exposure to nanocarriers with 363 DNA is crucial for enabling biocompatible gene delivery tools 364 for plants.<sup>59</sup> We evaluated protoplast viability of TMGMV-365 PAH coated with  $GT_{15}$ -Cy3 (0.1–0.5 mg/mL) or pDNA (0.04 366 mg/mL) using fluorescein diacetate (FDA),<sup>60</sup> a lipophilic 367 fluorescent dye that is permeable to membranes of living cells. 368 Following endogenous esterase-mediated enzymatic activity, 369 nonfluorescent FDA is transformed to fluorescein, a green 370 fluorescence compound. Broken cells lack esterases, rendering 371 them devoid of fluorescein signal. The FDA-treated protoplast 372 cells were analyzed by confocal microscopy imaging, and viable 373 cell percentages were calculated based on the fluorescein 374 presence. Both TMGMV-PAH-GT<sub>15</sub>-Cy3 or TMGMV-PAH-375 pDNA treated and control (untreated) protoplasts showed

bright green fluorescence characteristic of fluorescein and 376 normal morphology (Figure S13a,b). Approximately 71% ± 377 3.5 of cells remained viable after exposure to TMGMV-PAH- 378  $GT_{15}$ -Cy3 (0.1 mg/mL), while increased concentrations 379 resulted in a gradual reduction in fluorescein signal and 380 increased number of broken cells (Figure S13c). A dramatic 381 reduction in the fluorescein signal in protoplasts was observed 382 after exposure to TMGMV-PAH-GT<sub>15</sub>-Cy3 (0.5 mg/mL), in 383 which almost no viable cells were observed (Figure S13c). For 384 protoplasts exposed to the TMGMV-PAH:pDNA mass ratio 385 (1:3), approximately  $74\% \pm 3.0$  of cells remained viable, which 386 is not significantly different from the viability of untreated 387 protoplasts (Figure S13d). In contrast, when TMGMV-PAH 388 was loaded with pDNA at the mass ratios of 1:6 and 1:12, 389 significant decreases were observed in cell viability, approx- 390 imately  $65\% \pm 5.5$  (*P* < 0.039) at the 1:6 ratio and  $43\% \pm 8.5$  391 (P < 0.0003) at the 1:12 ratio cells were viable when compared 392 to the protoplasts-only cells (Figure S13d). The TMGMV- 393 PAH-pDNA concentration in this protoplast viability assay was 394 kept similar to that used in the transformation analysis (0.04 395 mg/mL). These findings suggest that an increased loading of 396 pDNA onto TMGMV-PAH can affect plant cell viability. 397 Biocompatibility of iTMGMV-PAH-pDNA in Arabidopsis 398 leaves was determined using propidium iodide, a fluorescent 399 dye that stains the nucleus of dead cells (Figure S14). Confocal 400 microscopy images of leaf cells infiltrated with our chosen 401 concentration for GFP expression analysis of 0.1 mg/mL  $_{\rm 402}$ iTMGMV-PAH: 0.6 mg/mL pDNA showed a similar 403 percentage of dead cells  $(4.5 \pm 1.7\%)$  to leaves treated with 404 buffer control (7.9  $\pm$  3.4%; P > 0.5; Figure S14a,b). Higher 405 concentrations of 0.15 mg/mL iTMGMV-PAH: 0.9 mg/mL 406 pDNA significantly increased the percentage of dead cells 407 (15.8  $\pm$  2.2%; P < 0.01). Overall, our results indicate that 408 DNA coated TMGMV-PAH are highly biocompatible with 409 plant cells both in vitro in plant protoplasts and in vivo in leaf 410 cells. 411

We engineered plant virus coat protein nanocarriers 412 (TMGMV-PAH) for facile plasmid DNA delivery into the 413 plant cell nucleus without mechanical or biological aid, with 414 high biocompatibility and the highest loading of DNA 415 nanocarriers for plant cells reported to date. We demonstrated 416 this approach using TMGMV-PAH that spontaneously 417 delivered a transgene (GFP) encoded in an expression vector 418 (pDNA) into plant protoplasts and epidermal cell nuclei. GFP 419 gene delivery and expression in plant cells has been mediated 420 by high aspect ratio carbon nanotubes.<sup>23–25,33</sup> In this work, we 421 used high aspect ratio protein-based nanomaterials, native 422 TMGMV in protoplasts, and inactivated iTMGMVs *in vivo* to 423 prevent plant infection.<sup>43</sup> TMGMV's ability to move across 424 plant cell barriers in numerous plant species<sup>43,61</sup> suggests that 425 these nanocarriers could mediate DNA delivery to protoplasts 426 or leaf cells from different plant species. 427

Future research will assess if pDNA mediated delivery by 428 TMGMV-PAHs in plant cells results in transient expression of 429 transgenes, similar to what has been reported in previous 430 studies about pDNA delivery using inorganic nanomateri- 431 als.,<sup>23,24,33</sup> or enable stable plant transformation and genome 432 editing with higher efficiency compared to current DNA 433 delivery protocols using biological or mechanical aid. TMGMV 434 may prove to be a promising tool for the delivery of genes, 435 small-interfering RNA (siRNA), and clustered regularly 436 interspaced short palindromic repeats (CRISPR) in plants 437 for gene editing applications. Targeted delivery approaches 438 439 could be implemented for TMGMV-mediated gene delivery 440 into plastid genomes including coating with targeting 441 peptides<sup>62</sup> for gene delivery to plant chloroplasts,<sup>24,6</sup> and a 442 mitochondria.<sup>25</sup> Our nanotechnology approach utilizing 443 TMGMV-PAH for DNA delivery paves the way for developing 444 plant virus-based nanocarriers with tunable and well-controlled 445 properties, <sup>41,42,63</sup> cost-effectiveness, scalability, <sup>64,65</sup> degradabil-446 ity,<sup>63</sup> and high biocompatibility,<sup>63,66</sup> which enable more 447 sustainable agriculture and advanced plant bioengineering.

### 448 ASSOCIATED CONTENT

#### 449 **Supporting Information**

450 The Supporting Information is available free of charge at 451 https://pubs.acs.org/doi/10.1021/acs.nanolett.3c04735.

- Detailed experimental procedures, including nanocarrier 452
- synthesis and characterization, microscopy, protoplast 453
- isolation, abaxial infiltration, RT-qPCR, gel electro-454

phoresis, and biocompatibility assays (PDF) 455

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#### Author Contributions

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<sup>#</sup>These authors contributed equally to this work (M.R.I. and 503 M.A.-Y.). J.P.G and N.F.S. conceived the idea and designed 504 experiments with M.R.I. M.R.I performed nanomaterial 505 synthesis and characterization, in vitro DNA loading and 506 delivery, gene expression analysis, cell viability, endocytosis, 507 and confocal microscopy assays. J.P.G designed in vivo 508 experiments with M.A.-Y. who performed inactivated nano- 509 material synthesis, in vivo pDNA delivery and gene expression 510 analysis using RT-qPCR and confocal microscopy, and 511 biocompatibility assays. N.F.S. and A.A.C. designed the 512 iTMGMV formulation for the in vivo studies, which was 513 prepared and characterized for quality control by A.A.C. G.V.L 514 contributed with data analysis. H.K. performed polymer 515 coating design and synthesis of nanocarriers, TEM, zeta 516 potential, and FTIR analysis of nanomaterials. I.G.-G. purified 517 and lyophilized native TMGMV. A.G.M.-B. performed TEM of 518 nanomaterials loaded with plasmid DNA and analysis with 519 I.G.-G. All authors contributed to writing the manuscript. 520

#### Notes

The authors declare the following competing financial 522 interest(s): A pending patent entitled Compositions and 523 Methods for Delivery of Nucleic Acids is based on this work. 524 J.P.G., M.R.I., H.K. (University of California, Riverside), and 525 N.F.S. (University of California, San Diego) are inventors in 526 this patent. Specific aspects of the manuscript covered in the 527 patent disclosure include compositions and methods for 528 delivery of DNA in plant cells. N.F.S. is a cofounder of, has 529 equity in, and has a financial interest in Mosaic ImmunoEngi- 530 nering Inc. N.F.S. is a cofounder and serves as manager of 531 Pokometz Scientific LLC, under which she is a paid as a 532 consultant to Mosaic ImmunoEngineering Inc., Flagship 533 Laboratories 95 Inc., and Arana Biosciences Inc. The other 534 authors declare no potential conflict of interest. 535

#### ACKNOWLEDGMENTS

This material is based on work mainly supported by the 537 National Science Foundation under Grant FMSG: Bio: 538 2134535. M.A.Y. received funding from NSF National 539 Research Traineeship Program Grant DBI-1922642. A.G.M.- 540 B. was supported by a CONAHCYT doctoral studies 541 scholarship (CVU 1062156). A.A.C. was supported through 542 NIFA-2022-67012-36698. We also thank partial support from 543 UC San Diego Materials Research Science and Engineering 544 Center (MRSEC) through NSF Grant DMR-2011924 to 545 N.F.S. The authors thank the University of California San 546 Diego - Cellular and Molecular Medicine Electron Microscopy 547 Core (UCSD-CMM-EM Core, RRID:SCR 022039) for 548 equipment access and technical assistance. The UCSD- 549 CMM-EM Core is supported in part by the National Institutes 550 of Health Award Number S10OD023527. 551

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