1	Single-component multilayered self-assembling protein nanoparticles
2	displaying extracellular domains of matrix protein 2 as a pan-influenza A
3	vaccine
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18 ABSTRACT (150 words)

19 The development of a cross-protective pan-influenza A vaccine remains a significant challenge. 20 Here, we designed and characterized single-component, self-assembling protein nanoparticles (1c-21 SApNPs) presenting the conserved extracellular domain of matrix protein 2 (M2e) from influenza 22 A viruses of human and other hosts. Vaccination with tandem repeats of M2e (M2ex3) displayed 23 on 1c-SApNPs demonstrated higher survival and lower weight loss compared to the soluble M2ex3 24 antigen against lethal challenges of H1N1 and H3N2 in mice. The mechanism of vaccine-induced 25 adaptive immunity was also investigated in mice. Compared with the soluble M2ex3 antigen, the 26 M2ex3 I3-01v9a 1c-SApNP formulated with a squalene-based adjuvant showed 672 times longer 27 follicular retention, 31 times greater exposure within follicular dendritic cell networks, and up to 28 2.5 times stronger germinal center reactions in lymph nodes. By inducing robust and durable M2e-29 specific functional antibody and T cell responses, the M2ex3-presenting I3-01v9a 1c-SApNP 30 provides a promising pan-influenza A vaccine candidate.

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32 ONE-SENTENCE SUMMARY (125 characters)

33 Protein nanoparticles displaying tandem M2e elicit robust and durable immunity that may protect
34 against influenza A viruses of diverse origins.

35 INTRODUCTION

36 Influenza (flu) is a respiratory disease caused by influenza viruses of the Orthomyxoviridae family 37 (1-5). Influenza viruses are enveloped negative-sense, single-stranded RNA viruses (6) that can be 38 classified as type A, B, C, or D, with influenza A and B viruses (IAVs and IBVs) posing a major 39 threat to human health. The most abundant surface glycoprotein, hemagglutinin (HA), binds the 40 host cell receptors and facilitates cell entry (7, 8). Under the host's immune selection pressure, HA 41 can acquire amino acid substitutions that lead to escape mutants (8). Another surface glycoprotein, 42 neuraminidase (NA), aids in the release of viral particles through cleavage of residues on the host 43 cell's surface (9, 10). Matrix protein 1 (M1) is involved in virus budding, while matrix protein 2 44 (M2) functions as a proton channel to facilitate the maintenance of pH during viral entry and viral 45 replication in host cells (11). IAVs can be classified into subtypes based on the antigenic properties 46 of HA and NA (4), with H1N1 and H3N2 being responsible for most human infections (8). IBVs 47 have a single HA/NA subtype, which can be classified into two lineages, Victoria and Yamagata 48 (1, 2). IAVs can infect many hosts, whereas IBVs are restricted to humans (12).

49 Seasonal flu vaccines have been used as a cost-effective public health tool since the 1940s 50 (13-15). Current flu vaccines are typically quadrivalent, covering two IAV subtypes (H1N1 and 51 H3N2) and two IBV lineages (Victoria and Yamagata), and are produced in chicken eggs (16). As 52 a result, current flu vaccines mainly generate strain-specific neutralizing antibodies (NAbs) and 53 may not protect against mismatched seasonal strains or more distinct strains generated through 54 "antigenic drift", during which HA and NA accummulate small mutations over time. Occasionally, 55 IAVs have the potential to cause global pandemics through "antigenic shift", in which HAs and 56 NAs from different host species recombine to form novel IAV strains against which the human 57 population lacks pre-existing immunity (17). Viral reassortment resulting in highly pathogenic

avian influenza (HPAI) acquiring animal-to-human transmissibility has been on the rise in recent years. As of 2021, there have been 863 cases of HPAI H5N1 and 66 cases of H5N6 in humans, with a > 50% fatality rate (*18*). Therefore, there is an urgent need for cross-protective flu vaccines (*16*), especially for potential pandemic strains originating from diverse animal reservoirs (*19*).

62 Various antigen and vaccine strategies have been explored to develop a universal influenza 63 vaccine (20-27). One strategy targets conserved internal proteins, such as nucleoprotein and M1, 64 to induce influenza-specific T cell responses (28). A second strategy aims to generate broadly 65 neutralizing antibodies (bNAbs) to the conserved regions of HA, such as the stem and parts within 66 the head domain (29-32), and of NA (33). Notably, the highly conserved ectodomain of the M2 67 protein (M2e) presents an attractive target for universal IAV vaccine development (30, 34-37) 68 because of the sequence conservation across IAVs and functional importance of the M2 proton 69 channel to virus fitness and replication. Although M2e is small (~23 aa) and poorly immunogenic, 70 it can be conjugated to large carriers to elicit antibody responses that effectively reduce viral 71 replication (38). Unlike HA and NA, M2e-specific antibodies protect via FcyR-dependent mechanisms, such as antibody-dependent cellular cytotoxicity (ADCC) and phagocytosis (ADCP), 72 73 rather than direct neutralization (37, 39-41). Various carriers have been used to increase the 74 immunogenicity of M2e vaccines, including hepatitis B core protein (HBc) (39), tobacco mosaic 75 virus (TMV) coat protein (42), keyhole limpet hemocyanin (KLH) (38), rotavirus NSP4 (43), 76 GCN4 (44), bacterial flagellin (45), and liposomes (46). Early human trials confirmed the 77 immunogenicity and tolerance of M2e vaccines but also revealed several weaknesses. While an 78 adjuvanted M2e-HBc fusion protein (39) induced a short-lived anti-M2e antibody response, an 79 M2e-flagellin fusion vaccine (45) caused undesirable side effects at higher doses. A vaccine 80 combining M2e with cytotoxic T lymphocyte (CTL) epitopes (47) induced strong cellular

immunity, but the response was narrow and slow, making it unsuitable for effectively mitigating
a future influenza pandemic. These clinical trials highlight the challenges facing M2e-based
vaccine development (*35, 48*), as well as the importance of vaccine carriers, adjuvants, balanced
antibody and T cell responses, and durability.

85 In recent studies, we have demonstrated a rational vaccine strategy that combines antigen 86 optimization and nanoparticle (NP) display. This strategy was inspired by the success of virus-like 87 particles (VLPs), which can be used as vaccines against cognate viruses or as carriers to improve 88 the immunogenicity of subdominant antigens (49-56). Due to their inherent complexity, VLPs can 89 be challenging to produce with yield, purity, and quality acceptable for clinical use. However, 90 protein NPs can be constructed to mimic the size and antigen display of VLPs, with the capability 91 of displaying diverse antigens. We have previously designed multilayered single-component self-92 assembling protein nanoparticles (1c-SApNP) based on bacterial proteins E2p and I3-01 that can 93 self-assemble into 60-mers of 22-25 nm (57-62). Genetic fusion of an antigen to the N-terminus 94 and a locking domain (LD) and pan-reactive T cell epitope (PADRE) to the C-terminus of an NP 95 subunit creates a "vaccine plasmid" encoding a single polypeptide, which will form a protein shell 96 with an array of antigens on the outside, a stabilizing inner LD layer, and a hydrophobic PADRE 97 core. These 1c-SApNPs can be expressed in Chinese hamster ovary (CHO) cells with high yield 98 and purity, and have been applied successfully to vaccines targeting HIV-1 (57, 61, 62), HCV (60), 99 Ebola virus (EBOV) (58), and SARS-CoV-1/2 (59, 63). Notably, these 1c-SApNPs can remain in 100 lymph nodes for weeks to interact with immune cells and generate robust germinal center (GC) 101 reactions, whereas individual soluble antigens are cleared within a few hours.

In this study, we rationally designed M2e-presenting 1c-SApNPs as cross-protective, pan influenza A vaccine candidates and characterized them both in vitro and in vivo. We first designed

104 I3-01v9a, a new version of the I3-01v9 1c-SApNP, for optimal presentation of monomeric antigens. 105 We then displayed human M2e (hM2e) on ferritin (FR), E2p, and I3-01v9a 1c-SApNPs, along 106 with a trimeric scaffold, for initial assessment. Following detailed in vitro characterization, these 107 hM2e immunogens were tested in mice, which were sequentially challenged with mouse-adapted 108 H1N1 and H3N2 after a two-dose vaccination. Based on the results, we next displayed tandem 109 copies of M2e (human, avian and swine), termed M2ex3, on the same carriers and characterized 110 these immunogens following a similar protocol. M2ex3 presented on 1c-SApNPs elicited 111 significantly higher M2e-specific antibody and T cell responses in immunized mice compared with 112 the soluble M2ex3 antigen. As a result, mice immunized with M2ex3-presenting 1c-SApNPs 113 showed higher survival against lethal heterosubtypic challenges. In the mechanistic analysis, the 114 M2ex3 1c-SApNPs exhibited prolonged retention (8 weeks) in lymph node follicles and generated 115 robust GC reactions, which may explain the vaccine-induced immunity and protection. Therefore, 116 the tandem M2e presented on an optimized I3-01v9a 1c-SApNP may provide an effective vaccine 117 candidate for durable cross-protection against seasonal and pandemic influenza A viruses.

118 **RESULTS**

119 Rational design of an I3-01v9a NP scaffold for presenting monomeric antigens

In our early studies, we utilized 24-mer ferritin (FR) and two 60-mers, E2p and I3-01, to display HIV-1 and HCV antigens (*57*, *60*). Locking domains (LDs) and a CD4 T-helper epitope (PADRE) were later incorporated into E2p and I3-01 (and its variant I3-01v9) to generate "multilayered" NP carriers, which were successfully used to display stabilized EBOV glycoprotein (GP) trimers (*58*), SARS-CoV-1/2 spikes (*59*, *63*), and HIV-1 envelope (Env) trimers (*57*) for vaccine development. Notably, the I3-01 NP scaffold appeared to be particularly amendable to structural modification, with multiple design variants tested in our previous studies (*57-61*). In this study, we rationally

127 optimized the I3-01v9 NP scaffold to improve the surface display of monomeric protein antigens 128 (fig. S1). The N-termini of I3-01v9 forms a wide triangle of 50.5 Å, making it more suitable than 129 E2p for displaying monomeric antigens. However, the first amino acid (the antigen anchoring site) 130 is below the NP surface, and as a result, a long flexible peptide linker must be used to connect the 131 antigen to the I3-01v9 N-terminus, leading to the increased structural instability of the antigen-NP 132 fusion constructs. Here, we hypothesized that extending the I3-01v9 N-terminal helix would allow 133 its first residue to reach the NP surface, and consequently, a short peptide linker between the 134 antigen and NP backbone would be sufficient. A computational procedure was devised to facilitate 135 rational design (fig. S1). Briefly, the backbone of a helix (residues 953-982) from a c-MYC 136 transcription factor protein (PDB ID: 6G6L) was grafted onto an I3-01v9 subunit by using residues 137 Glu2 and Glu3 of I3-01v9 for structural fitting. The extended N-terminal helix was then truncated 138 to 11 residues so that its first residue would be just above the NP surface. Then, a protein structure 139 sampling program, CONCOORD (64), was used to generate 1,000 slightly perturbed 140 conformations for the modified I3-01v9 subunit. Next, an ensemble-based protein design program 141 that was previously used to optimize HIV-1 Env trimers (57) and HCV E2 cores (60) was 142 employed to predict the sequence for the first 9 residues of the 11-residue segment using C_{α} and 143 C_{β} -based RAPDF scoring functions (65). The final design, termed I3-01v9a, was determined by 144 combining results from predictions using both scoring functions (fig. S1).

145 Human M2e (hM2e) on multilayered 1c-SApNPs as human influenza A vaccines

The M2 protein from IAVs is a highly conserved proton channel, with a small ectodomain of 24 amino acids in length (M2e) (*35*). Anti-human M2e (hM2e) antibodies have been shown to reduce viral replication, thus decreasing clinical symptoms and severity of the disease. In the immunogen design, M2e is hereafter defined as residues 2-24 excluding the first amino acid (M1).

150 Previously, we rationally redesigned viral antigens and engineered antigen-presenting 151 SApNPs for in vitro and in vivo characterization (57-62). Following a similar strategy, we 152 designed an hM2e scaffold and three SApNP constructs. The crystal structures of hM2e in complex 153 with antibodies mAb65 and mAb148 (66, 67) indicate that hM2e is flexible and can adopt different 154 conformations upon antibody binding. MAb65 recognizes a short turn of Pro10 to Asn13, whereas 155 mAb148 binds to an N-terminal epitope (Ser2-Glu8). We first utilized a capsid-stabilizing protein 156 of lambdoid phage 21, SHP (PDB ID: 1TD0), as a trimeric scaffold to present hM2e. With a 5GS 157 linker, two hM2e epitopes would span ~9.1 nm (measured at Pro10) when all three 1TD0-attached 158 hM2e segments were in a fully open conformation (Fig. 1A). We then displayed hM2e on FR 24-159 mer and reengineered E2p and I3-01v9a 60-mers, all with a 5GS linker (Fig. 1A). Molecular 160 modeling revealed well-spaced hM2e peptides on the particle surface, with diameters of 20.9 nm, 161 29.1 nm, and 32.4 nm for FR, E2p, and I3-01v9, respectively (Fig. 1A). Following a similar 162 terminology, the "multilayered" E2p and I3-01v9a are named E2p-LD4-PADRE (or simply E2p-163 L4P) and I3-01v9a-LD7-PADRE (or simply I3-01v9a-L7P), respectively. One hM2e scaffold and 164 three hM2e-presenting SApNPs were subjected to in vitro characterization.

165 All four hM2e constructs (fig. S2A) were transiently expressed in 25 ml ExpiCHO cells, 166 purified by immunoaffinity chromatography (68) using mAb65 or mAb148 columns, and analyzed 167 by size exclusion chromatography (SEC) (Fig. 1B). The SEC profiles indicated high yield for 168 hM2e-5GS-1TD0 and hM2e-5GS-FR and, in contrast, a notably lower yield for hM2e-5GS-I3-169 01v9a-L7P, as shown by the ultraviolet absorbance at 280 nm (UV₂₈₀). Among the four constructs, 170 hM2e-5GS-E2p-L4P had the lowest yield. Of note, all three SApNPs showed two SEC peaks at 8-171 9 ml and 13-14 ml, corresponding to different NP species. Sodium dodecyl sulfate-polyacrylamide 172 gel electrophoresis (SDS-PAGE) under reducing conditions showed bands for hM2e-5GS-1TD0

173 (13.9 kDa), FR (21.0 kDa), E2p-L4P (38.9 kDa) and I3-01v9a-L7P (33.3 kDa) that were consistent 174 with their calculated molecular weights (MW) (Fig. 1C, left). Blue native-polyacrylamide gel electrophoresis (BN-PAGE) confirmed the high purity of SApNP samples after IAC using an 175 176 mAb148 column, displaying a single high-MW band for each SApNP with no sign of unassembled 177 species (Fig. 1C, right). The structural integrity of IAC-purified SApNPs was validated by 178 negative-stain electron microscopy (nsEM), which showed distinct morphologies for three hM2e-179 presenting SApNPs (Fig. 1D). Notably, hM2e-presenting SApNPs appeared to form "clusters" in 180 solution, which likely correspond to the high-MW peak (8-9 ml) in their SEC profiles (Fig. 1B). 181 Analysis of mAb148-purified SApNPs by dynamic light scattering (DLS) revealed larger-than-182 expected "particle" size for FR (57.2 nm), E2p-L4p (69.6 nm), and I3-01v9a-L7P (46.1 nm) (Fig. 183 **1E**), consistent with the nsEM results. Interestingly, DLS analysis of the SEC fraction (13-14 ml) 184 of an mAb148-purified hM2e-5GS-FR sample still showed three particle size populations, 185 suggesting that cluster formation is an intrinsic feature of hM2e-presenting SApNPs (fig. S2B). 186 Differential screening calorimetry (39) was used to quantify the thermostability of these hM2e 187 constructs. Thermograms were obtained for hM2e-5GS-1TD0 and hM2e-5GS-FR, which showed 188 a melting temperature (T_m) of 72.5-72.8°C and a similar T_{onset} of 57.9-58.7°C (Fig. 1F). For the 189 two large 60-mers, heating, enzyme-linked immunosorbent assay (ELISA), and nsEM were 190 combined to estimate thermostability. Briefly, SApNP samples were heated to 50°C, 60°C, and 191 70°C for 15 min prior to ELISA analysis against mAb148 and mAb65 (Fig. 1G, fig. S2C) and 192 nsEM (fig. S2D). While antibody binding, measured by half maximal effective concentration 193 (EC₅₀), largely remained comparable within the temperature range of 4-70 $^{\circ}$ C, nsEM images 194 showed signs of irregular particle shapes at 70°C, suggesting that the melting points for hM2e-195 5GS-E2p-L4P and I3-01v9a-L7P may be between 60 and 70°C. Lastly, we performed bio-layer

interferometry (BLI) to quantify antibody binding kinetics for the four hM2e immunogens.
Although the three SApNPs outperformed the trimeric hM2e scaffold regardless of the antibody
tested (Fig. 1H, fig. S2E), mAb148 and mAb65 showed different profiles with stronger binding
observed between mAb65 and the two large 60-meric SApNPs (fig. S2E).

In summary, hM2e can be successfully displayed on all three SApNPs, consistent with our previous studies where stabilized HIV-1 (*57, 61, 62*), HCV (*60*), EBOV (*58*), and SARS-CoV-1/2 (*59, 63*) antigens were displayed on the NP surface. Extensive biochemical, biophysical, structural, and antigenic characterizations provided detailed in vitro profiles of hM2e-presenting 1c-SApNPs, thus allowing evaluation of these vaccine immunogens in vivo.

205 In vivo evaluation of a scaffolded hM2e trimer and hM2e-presenting 1c-SApNPs in mice

206 The immunogenicity and protective efficacy of hM2e trimer and hM2e-presenting SApNPs were 207 evaluated in BALB/c mice. First, mouse-adapted A/Puerto Rico/8/1934 (PR8) H1N1 and A/Hong 208 Kong/1/1968 (HK/68) H3N2 were grown in MDCK cells, and various dilutions of the propagated 209 viruses were used to challenge mice to establish a 50% lethal dose in mice (LD₅₀) (Fig. 2A). 210 Plaque-forming units (PFU) of the virus were measured via plaque assay. Using survival rates of 211 mice at various viral dilutions, the Reed-Muench and Spearman-Karber methods (69, 70) were used to calculate the 50% endpoint titers for survival. For a stock of 3.8×10^5 PFU/ml PR8 (H1N1). 212 213 the LD₅₀ was determined to be 12 PFU/ml, with the LD₅₀ \times 10 calculated to be 120 PFU/ml. For a stock of 5.5×10^4 PFU/mL of HK/68 (H3N2), the LD₅₀ was determined to be 1.2×10^4 PFU/ml, 214 215 with the $LD_{50} \times 10$ determined to be 1.2×10^5 PFU/ml.

Groups of BALB/c mice were immunized at weeks 0 and 3 with hM2e vaccines adjuvanted with aluminum phosphate (AP) and challenged at weeks 6 and 10 with mouse-adapted H1N1 and H3N2, respectively (**Fig. 2B**). Survival and weight loss were measured after sequential challenges

219 with $LD_{50} \times 10$ of PR8 (H1N1) and HK/68 (H3N2) (Fig. 2C). For the PR8 H1N1 challenge, all 220 naïve mice succumbed to the challenge by day 8. In the hM2e-5GS-1TD0 (trimer) group, 80% of 221 mice died by day 9, with 20% of mice surviving the challenge. In contrast, 100% of mice survived 222 in the FR, E2p, and I3-01v9a SApNP groups. Similar trends were observed in the average peak 223 weight loss, with naïve mice losing the most weight $(22.1 \pm 1.3\%)$. Similarly, 1TD0 mice lost 19.1 224 \pm 3.3% of their total body weight on average. Among the hM2e SApNP groups, FR mice lost 12.1 225 \pm 8.4% of their total body weight on average, E2p mice lost 16.4 \pm 3.9%, and I3-01 lost the least 226 weight with an average of $10.6 \pm 4.5\%$. In the positive control group, mice immunized with 227 inactivated PR8 H1N1 lost the least weight in the strain-matched challenge, with an average loss 228 of $6.5 \pm 3.8\%$. For the second challenge with HK/68 H3N2, the lowest survival was observed for 229 the inactivated PR8 group, with only 56% of mice surviving the heterologous challenge. The two 230 surviving 1TD0 mice from the previous challenge and all SApNP mice survived the HK/68 H3N2 231 challenge. Following a similar trend, the highest body weight loss was observed for the inactivated 232 PR8 group with an average peak weight loss of $16.9 \pm 4.9\%$. All hM2e vaccine groups showed 233 lower peak weight loss with $10.8 \pm 2.4\%$, $4.4 \pm 4.0\%$, $6.2 \pm 3.4\%$, and $7.9 \pm 2.8\%$ for 1TD0, FR, 234 E2p, and I3-01v9a, respectively. The hM2e-binding antibody response in mouse serum was 235 assessed by ELISA using an hM2e-5GS-foldon trimer (Fig. 2D, fig. S3A). The hM2e SApNP 236 groups demonstrated superior serum binding, measured by EC_{50} titers, at all the time points tested, 237 with the greatest fold increase observed at week 5 for FR (31.2), E2p (52.3), and I3-01v9a (42.8) 238 compared to the 1TD0 group (Fig. 2D). The hM2e vaccine groups showed the highest antibody 239 EC₅₀ titers 4 weeks after the H3N2 challenge at week 14.

In summary, hM2e SApNPs significantly outperformed the soluble hM2e trimer in the vaccination/viral challenge experiment, showing higher survival and lower weight loss that were

well-correlated with the heightened M2e-specific serum antibody titers. These hM2e SApNPs also
demonstrated cross-protection against H1N1 and H3N2 challenges, whereas the inactivated PR8
H1N1 vaccine only protected against the strain-matched challenge.

245 Tandem M2e (M2e×3) on multilayered 1c-SApNPs as pan-influenza A vaccines

The effectiveness of seasonal influenza vaccines range between 10 and 60% as estimated by the U.S. Flu Vaccine Effectiveness Network (*15*). In addition to antigenic drift in circulating human influenza virus strains, the unanticipated emergence of novel strains from swine and avian hosts often causes outbreaks with increased mortality and morbidity. Thus, a broadly protective M2ebased influenza vaccine strategy must incorporate M2e from diverse species.

251 Following the same strategy for hM2e, we designed a trimeric scaffold and three SApNPs 252 to present tandem M2e repeats as vaccine immunogens. Briefly, the hM2e, avian/swine M2e, and 253 human/swine M2e sequences were linked in tandem with short G4 linkers. A structural model of 254 M2ex3 was generated from the crystal structure of hM2e in complex with mAb65 (PDB ID: 4N8C) 255 (Fig. 3A, left). Although this compact structural model may not represent M2ex3 conformations 256 in solution, it may facilitate rational design of the M2ex3 orientation on various carrier scaffolds. 257 For the M2ex3-5GS-1TD0 trimer, the two outmost hM2e epitopes would span ~9.4 nm (measured 258 at Pro10 of hM2e) when the scaffolded M2ex3 segments adopt an extended conformation (Fig. 259 **3A**, middle). For the three SApNPs, molecular modeling yielded diameters of 23.2 nm, 32.4 nm, 260 and 36.2 nm for M2ex3-5GS-FR, E2p-L4P, and I3-01v9a-L7P, respectively, measured at Pro10 261 of hM2e (Fig. 3A, right three). The display of tandem M2ex3 increased not only the particle size 262 but also the number of M2e epitopes, from 60 to 180, for enhanced immune recognition.

263 The four M2ex3 constructs (fig. S4A) were expressed and purified using the same strategy
264 as for their hM2e counterparts (Fig. 3B). Overall, M2ex3 immunogens showed a similar pattern

265 of expression yield in ExpiCHO cells, with the ranking of 1TD0 > FR > 13-01v9-L7P >> E2p-L4P. 266 Of note, the SEC profiles showed a less pronounced peak at 9 ml for FR and I3-01v9-L7P, 267 suggesting a reduced tendency to form NP clusters. Reducing SDS-PAGE showed bands on the 268 gel consistent with MW calculated for M2ex3-5GS-1TD0 (19.3 kDa), FR (26.3 kDa), E2p-L4P 269 (44.4 kDa), and I3-01v9a-L7P (38.7 kDa) (Fig. 3C, left). However, a second band was observed 270 on the gel for M2ex3-5GS-1TD0, FR, and E2p-L4P under reducing conditions. While the extra 271 bands for M2ex3-5GS-1TD0 and E2p-L4P may indicate higher-MW species that are resistant to 272 the reducing agents, the lower band noted for M2ex3-5GS-FR likely suggests degradation during 273 processing. Nonetheless, BN-PAGE confirmed particle assembly and purity for three M2ex3 274 SApNPs (Fig. 3C, right). Similarly, nsEM micrographs demonstrated well-formed, homogeneous 275 particles for all three M2ex3 SApNP samples following mAb148 purification (Fig. 3D). In the 276 DLS profiles (Fig. 3E), M2ex3-5GS-FR exhibited a two-peak distribution with the majority peak 277 showing single particles, indicated by an average size of 28.3 nm. Similarly, M2ex3-5GS-I3-278 01v9a-L7P yielded a homogenous distribution consistent with single particles. In contrast, M2ex3-279 5GS-E2p-L4P formed clusters, as seen in nsEM images and indicated by the DLS-derived particle 280 size distribution. The tandem design appeared to cause a reduction in thermostability for M2ex3-281 5GS-FR, with lower T_m (60.9°C) and T_{onset} (45.7°C) values. The melting point was estimated for 282 the two 60-mer SApNPs using the alternative approach devised for hM2e SApNPs. Similar 283 antibody binding affinity, as indicated by the EC_{50} value, was observed for M2ex3 scaffolds and 284 SApNPs across the whole temperature range (4-70°C) (Fig. 3G, fig. S4B), although irregular 285 particle shapes were noted at 70°C in EM micrographs (fig. S4C). The interactions of M2ex3 286 immunogens with two human antibodies (mAb65 and mAb148) were assessed by BLI (Fig. 3H, 287 fig. S4D). The advantage of particulate display was exemplified by the higher binding signals,

similar to the case of hM2e SApNPs. Interestingly, the binding profiles (both on-rate and signals)
notably improved for M2ex3-5GS-I3-01v9a-L7P (Fig. 3H, rightmost), suggesting that antigenicity
may be affected by both the epitope number and spacing on a particular NP scaffold. In summary,
tandem M2ex3 SApNPs exhibit greater homogeneity and antigenicity than hM2e SApNPs, while
sharing similar yield, structure, and thermostability.

293 Protection against influenza A virus challenge by tandem M2ex3 vaccines in mice

294 BALB/c mice were immunized with tandem M2ex3 vaccines adjuvanted with AP or AddaVax (a 295 squalene-based nanoemulsion adjuvant; AV) at weeks 0 and 3 and thereafter challenged at weeks 296 6 and 10 with H1N1 and H3N2, respectively (Fig. 4A). Survival and weight loss were recorded 297 for both challenges (Fig. 4B). After an $LD_{50} \times 10$ challenge with PR8 (H1N1), all naïve mice died 298 by day 8. In mice immunized with AP-adjuvanted tandem M2ex3 vaccines, 50% of 1TD0 (trimer) 299 mice died by Day 9. Paired with the same AP adjuvant, M2ex3 SApNPs demonstrated higher 300 survival rates: 88% of FR, 100% of E2p, and 100% of I3-01v9a mice survived the H1N1 challenge. 301 In terms of peak weight loss, naïve mice lost the most weight with an average of $21.7 \pm 2.9\%$, and 302 1TD0 (trimer) mice lost the second highest amount of weight (19.4 \pm 7.3%), as expected for the 303 small soluble M2e antigen. In general, M2ex3 SApNP groups showed lower peak weight loss: FR 304 $(15.5 \pm 8.1\%)$, E2P $(11.2. \pm 3.8\%)$, and I3-01v9a $(15.5 \pm 5.3\%)$. Several AddaVax-adjuvanted 305 M2ex3 vaccine groups outperformed their AP counterparts, demonstrating both higher survival 306 and lower peak weight loss. The AddaVax-adjuvanted M2ex3 trimer group showed a slightly 307 higher survival rate at 63% compared to its AP-adjuvanted counterpart (50%). All AddaVax-308 adjuvanted M2ex3 SApNPs achieved 100% survival, with an improvement noted for the M2ex3 309 FR group (12% higher survival compared to its AP counterpart) and no difference in the E2p and 310 I3-01v9a groups between the two adjuvants (all 100% survival). For peak weight loss, amongst

311 AddaVax-adjuvanted groups, the M2ex3 trimer showed the highest weight loss with $18.9 \pm 5.9\%$. 312 Most M2ex3 SApNPs adjuvanted with AddaVax showed lower peak weight loss compared with 313 their AP-adjuvanted counterparts: $13.1 \pm 7.2\%$ (FR), $12.5. \pm 7.7\%$ (E2p), and $10.0 \pm 4.2\%$ (I3-314 01v9a). Inactivated PR8 mice lost the least weight against the strain-matched challenge, with an 315 average of $4.4 \pm 5.3\%$. Next, protection against the second challenge with HK/68 H3N2 was 316 assessed (Fig. 4B). While all mice in the naïve group died by day 5, inactivated PR8 showed 317 limited protection with a survival rate of 63% against the non-strain-matched challenge. All 318 M2ex3-immunized mice that survived the prior PR8 challenge, adjuvanted with AP or AddaVax, 319 survived the HK/68 challenge. Similar to the survival rates, the highest peak weight loss was seen 320 in the naïve group with $19.5 \pm 2.9\%$, followed by the inactivated PR8 group with $15.7 \pm 4.9\%$. 321 M2ex3 trimer and FR, E2p, and I3-01v9a SApNPs adjuvanted with AP showed lower weight loss 322 with $8.9 \pm 5.2\%$, $4.7 \pm 3.2\%$, $6.5 \pm 1.9\%$, and $4.4 \pm 3.0\%$, respectively. Similarly, M2ex3 trimer 323 and SApNP groups adjuvanted with AddaVax showed minimal weight loss: trimer with 4.3 ± 2.1 , 324 FR with $4.3 \pm 1.0\%$, E2p with $5.9 \pm 3.4\%$, and I3-01v9a with $4.9 \pm 3.3\%$. Amongst all M2ex3 325 vaccine formulations, M2ex3 I3-01v9a/AddaVax demonstrated the most effective protection 326 against sequential H1N1 and H3N2 challenges in a mouse model.

The viral load in lungs of mice at day 5 post-PR8 challenge was used as another metric to evaluate the effectiveness of vaccine protection (**Fig. 4C**). Mice were immunized and challenged as described above and sacrificed at day 5 post-infection. Lungs were collected, mechanically disaggregated, and centrifuged to pellet cells. PFUs were measured in the supernatants via plaque assay. Overall, naïve mice had the highest virus load, $6.0 \times 10^5 \pm 3.3 \times 10^5$ PFU/ml. The M2ex3 trimer, FR, E2p, and I3-01v9a groups showed significantly lower viral titers, with 2.9, 7.5, 24.3, and 18.1-fold-lower virus load than the naïve group, respectively. Compared to M2ex3 trimer, FR,

E2p, and I3-01v9a showed 2.5, 8.3, and 6.3-fold-lower virus in lungs, although this difference was not statistically significant. The mice immunized with inactivated PR8 adjuvanted with AddaVax (positive control) did not show any detectable viral loads in lungs. Based on the criteria of survival, weight loss, and viral load in lungs, M2ex3 I3-01v9a SApNP/AddaVax appeared to be the most effective vaccine among all the NP/adjuvant formulations tested.

339 Evaluation of M2e×3 vaccine-induced antibody responses

340 Both E2p and I3-01v9a SApNP groups demonstrated superior serum binding to an M2ex3-5GS-341 foldon probe at all time points compared to the trimer group (Fig. 5A, fig. S5A). The greatest fold 342 difference between SApNP and trimer groups was obtained at week 2, suggesting a rapid onset of 343 anti-M2ex3 antibody response elicited by SApNPs. When paired with AP, E2p and I3-01v9a 344 SApNPs yielded 31.6-fold and 83.8-fold higher EC₅₀ titers than the 1TD0 trimer, whereas an even 345 greater fold difference, 47.6 and 102.4 respectively, was noted when these two SApNPs were 346 adjuvanted with AddaVax. The highest EC₅₀ titers were observed for the SApNP groups at week 347 5, where E2p and I3-01v9a showed 6.7/5.7-fold and 4.9/4.7-fold higher EC₅₀ values than the 348 M2ex3 trimer, respectively, when adjuvanted with AP/AddaVax. Interestingly, FR paired with 349 either adjuvant significantly underperformed E2p and I3-01v9a after the second dose, and trimer 350 at the later time points. Notably, AddaVax groups always outperformed AP groups with higher 351 EC_{50} titers, highlighting the importance of a potent adjuvant for eliciting M2e-specific antibody 352 responses. Importantly, we also confirmed that the incorporation of two non-human M2e epitopes 353 into the hM2e constructs does not reduce hM2e-specific EC₅₀ titers in the M2ex3-immune sera 354 compared to the hM2e-immune sera, as indicated by the hM2e-5GS-foldon probe (Fig. S5B). In 355 fact, mouse sera induced by the M2ex3 trimer and I3-01v9a SApNP formulated with AddaVax 356 showed similar or higher EC₅₀ titers compared to their AP-adjuvanted hM2e counterparts.

357 The recognition of homotetrameric M2e, which represents the "native" M2e conformation 358 during IAV infection, by M2ex3-immune sera was assessed for the AddaVax-adjuvanted 1TD0, 359 FR, and I3-01v9 groups (Fig. 5B, fig. S6). In this experiment, ELISA was performed to test serum 360 binding to M2e presented on MDCK cells infected with pandemic or seasonal H1N1 and H3N2 361 strains. The I3-01v9a group demonstrated significantly higher serum binding to M2e expressed on 362 MDCK cells infected with challenge strain A/Puerto Rico/8/1934 (H1N1) and challenge/pandemic 363 strain A/Hong Kong/1/1968 (H3N2). Additionally, the I3-01v9a group also showed higher serum 364 binding to pandemic A/California/04/2009 (H1N1) and other H1N1 and H3N2 strains: A/Solomon 365 Islands/2/2006 (H1N1), A/Brisbane/10/2007 (H3N2), and A/Aichi/2/1968 (H3N2). As a negative 366 control, serum binding was also assessed against two IBV strains, which express an M2e that is 367 shorter than and antigenically distinct from IAV M2e. As expected, the 1TD0, FR, and I3-01v9a 368 groups showed minimal serum binding to IBV strains B/Brisbane/60/2008 (Victoria Lineage) and 369 B/Florida/4/2006 (Yamagata Lineage). A human M2e antibody in the immunoglobulin form (71), 370 termed mAb148 (66), was used as a positive control in serum ELISA against IAVs. Based on the 371 in vitro and in vivo evaluation, M2ex3 I3-01v9a SApNP adjuvanted with AddaVax was selected 372 as the lead M2ex3 vaccine candidate for further analysis.

373 Distribution, trafficking, and retention of M2ex3 trimers and SApNPs in lymph nodes

Following a similar protocol that was used to analyze HIV-1 and SARS-CoV-2 SApNP vaccines (57, 63), we studied the in vivo behavior of the M2ex3 1TD0 trimer and two SApNPs (FR and I3-01v9a) to achieve a better understanding of their interaction with immune cells in lymph nodes and their ability to induce adaptive immune responses. To mount an effective humoral response, these vaccines must be transported through lymphatics and accumulate in lymph node follicles. The immunogens will then be presented to B cells to generate a robust antibody response through

380 crosslinking of B cell receptors (BCRs) (72-75). We first studied the transport and distribution of 381 M2ex3-presenting I3-01v9a SApNPs in lymph nodes. We injected a single dose of the immunogen 382 intradermally through the footpads (4 footpads, 10 µg/footpad) and isolated the sentinel brachial 383 and popliteal lymph nodes from both sides of the mouse's body at 48 h post-immunization. 384 Immunostaining of the excised lymph node sections was carried out using mAb148 and mAb65 385 (66, 67) to detect M2ex3 presented on I3-01v9a SApNPs (Fig. 6A). Immunohistological images 386 obtained after staining with both antibodies demonstrated a similar distribution of M2ex3 I3-01v9a 387 SApNPs in the center of lymph node follicles (Fig. 6A, images on the left; Fig. 6B, schematics on 388 the right). This intra-lymph node distribution pattern is consistent with the results obtained from 389 previous studies assessing SARS-CoV-2 spike SApNPs (63), HIV-1 Env SApNPs (57) and 390 ovalbumin-conjugated gold NPs (76, 77). Due to the better signal-to-noise ratio, mAb148 was used 391 hereafter to examine the trafficking of three M2ex3 immunogens in lymph nodes.

392 We next studied the trafficking and retention patterns of the M2ex3 trimer and two SApNPs 393 in lymph node follicles up to 8 weeks after a single-dose injection (4 footpads, 10 μ g/footpad) 394 (Fig. 6C). The histological images showed that all M2ex3 immunogens were transported into 395 lymph nodes and accumulated in the subcapsular sinus within 2 h (Fig. 6C). The M2ex3 trimer 396 was trafficked into follicles within 2 h and completely cleared by 12 h. In contrast, the M2ex3 FR 397 SApNP began to be present in follicles at 12 h, reached peak accumulation at 1 week, and remained 398 detectable up to 5 weeks. The M2ex3 I3-01v9a SApNP demonstrated the most superior follicular 399 retention with a period of 8 weeks. The mAb148-stained area was quantified in a time-dependent 400 manner, showing a ~672-times longer retention for the I3-01v9 SApNP compared to the M2ex3 401 trimer (Fig. 6C-D). The area under the curve (78) suggested that the exposure of M2ex3 presented 402 on SApNPs in follicles would be 14-31 times higher than the soluble M2ex3 trimer (Fig. 6E).

403 Additionally, the FR and I3-01v9a SApNPs also showed 45-86 times greater accumulation 404 compared with the M2ex3 trimer at 1 week (Fig. 6F). These results are consistent with our previous 405 findings (57, 63, 76), in which small particles (< 15 nm) can be cleared from lymph node follicles 406 within 48 h, whereas large particles (30-100 nm) remain for weeks. Importantly, I3-01v9a SApNPs 407 displaying M2ex3 antigens or BG505 trimers showed significantly longer follicular retention than 408 those presenting SARS-CoV-2 spikes (~8 weeks vs. ~2 weeks), suggesting a correlation between 409 antigen retention and antigen thermostability (a T_m value of 65°C or greater for M2ex3 and BG505 410 Env vs. 48°C for SARS-CoV-2 spike). Surprisingly, the M2ex3 I3-01v9a SApNP with a low glycan content exhibited identical trafficking and retention patterns to the highly glycosylated 411 412 BG505 Env SApNP (57), suggesting a minimal impact of glycan content on vaccine transport and 413 retention in lymph node follicles. Next, we studied the accumulation and retention patterns of these 414 three M2ex3 immunogens at 2 and 5 weeks using a prime-boost regimen (injected into 4 footpads 415 at weeks 0 and 3, 10 µg/footpad) (Fig. 6G). A similar pattern to the single-dose injection was 416 observed. Interestingly, M2ex3 trimers were detected in follicles at 5 weeks after the boost and 417 showed improved retention compared to the single dose, consistent with our previous SARS-CoV-418 2 study (63). Improvement in accumulation and retention after boost were also observed for the 419 FR and I3-01v9a SApNPs. Overall, displaying tandem M2ex3 on the I3-01v9a SApNP showed 8-420 fold greater accumulation in follicles compared to the soluble trimer 2 weeks after the boost (Fig. 421 **6H**). Prolonged retention of the M2ex3 I3-01v9a SApNP vaccine in lymph node follicles may 422 suggest improved longevity of vaccine-induced immunity.

Follicular dendritic cells (FDCs) located in the center of lymph node follicles are essential
for retention and presentation of the native-like antigens to stimulate B cell responses (72, 74, 75).
FDCs can collect and align soluble antigens, large particles such as immune complexes, viruses,

426 and bacteria on their surfaces or dendrites through a complement receptor-dependent mechanism 427 to generate and maintain GC reactions (73-76, 79, 80). Our previous studies of ovalbumin-428 conjugated gold NPs (76), and SARS-CoV-2 and HIV-1 antigen-presenting SApNPs (57, 63) 429 suggest that FDC networks may be the key cellular component to retain M2ex3 SApNPs. To test 430 this possibility, we collected lymph nodes at the peak of SApNP accumulation (1 week) and other 431 timepoints (48 h to 8 weeks) following a single-dose injection (Fig. 6I, fig. S7A-D). Lymph node 432 tissue was stained with mAb148 (white) for M2ex3, anti-CD21 antibodies (green) for FDCs, and 433 anti-CD169 antibodies for subcapsular sinus macrophages. The signals from both M2ex3 FR and 434 I3-01v9 SApNPs (anti-M2e mAb148) showed colocalization with FDCs (CD21⁺) at 1 week (Fig. 435 6I), confirming the retention of M2ex3 SApNPs in FDC networks.

436 Characterization of GC reactions induced by M2ex3 trimers and SApNPs

437 In GCs, B cells undergo somatic hypermutation, selection, affinity maturation, and class switching, 438 eventually becoming memory or antibody-secreting plasma cells (73, 81-84). FDC networks and 439 T follicular helper ($T_{\rm fh}$) cells support GC formation and maintenance (85, 86). Here, we 440 hypothesized that M2ex3 SApNPs retained by FDC networks could induce more robust and long-441 lived GC reactions in lymph node follicles compared to the soluble M2ex3 trimer. First, we 442 examined whether M2ex3 I3-01v9 SApNPs can induce strong GC reactions after single-dose 443 injections (4 footpads, 10 µg/footpad). Vaccine-induced GC B cells (GL7⁺, red) and T_{fh} cells 444 (CD4⁺Bcl6⁺, co-labeled with cyan and red) were characterized by immunohistology. We observed 445 large GCs attached to FDC networks (CD21⁺, green) with organized dark zone (DZ) and light zone 446 (LZ) compartments in follicles (B220⁺, blue) (Fig. 7A, left). In addition to antigen retention and 447 presentation by FDC networks, T_{fh} cells appear in the LZ of GCs to support B cell stimulation 448 (Fig. 7A, right). Next, we performed immunohistological analysis on the M2ex3 trimer and two

449 SApNPs at 2, 5, and 8 weeks after a single-dose injection (Fig. 7B, fig. S8A-C) and at 2 and 5 450 weeks after the boost (Fig. 7C, fig. S8D-E). Following a previously established protocol (57, 63), we quantified GC reactions using two metrics: GC/FDC ratio (the frequency of GC formation 451 452 associated with FDC networks) and size of GCs. The M2ex3 trimer and both SApNPs induced 453 robust GCs, with the I3-01v9a SApNP showing the largest GCs at 2 weeks after a single-dose 454 injection (Fig. 7B, fig. S8A). As expected, the GC/FDC ratio and GC size declined over time in 455 all groups. Notably, while the GC/FDC ratio for the M2ex3 trimer group decreased to \sim 50%, the 456 I3-01v9a SApNP generated strong and durable GC reactions that lasted for 8 weeks (Fig. 7B and 457 7D, fig. S8C). GCs were restored for all vaccine groups after the boost, but the GC/FDC ratio for 458 the trimer group decreased again significantly at 5 weeks after the boost. Overall, the M2ex3 I3-459 01v9a SApNP generated GCs 2.5 times the size after one dose (Fig. 7B and 7D) and 1.7 times the 460 size after the boost (Fig. 7C and 7E) of GCs elicited by the soluble trimer at 8 weeks.

461 The GCs were further analyzed by flow cytometry. We collected sentinel lymph nodes at 462 2, 5, and 8 weeks after a single-dose injection (Fig. 7F, and fig. S9), and at 2 and 5 weeks after the 463 boost (Fig. 7G) (4 footpads, $10 \mu g/injection$) following the prime-boost regimen. The lymph node 464 tissues were disaggregated into a single cell suspension and stained with an antibody cocktail. The 465 percentage and number of GC B cells and T_{fh} cells were used to quantify the GC reactions, which 466 correlate with the immunohistological results (Fig. 7A-E). Flow cytometry indicated that M2ex3 467 I3-01v9 SApNP elicited the largest GC B cell and T_{fh} cell populations after a single-dose injection 468 (Fig. 7F). GC reactions peaked at 2 weeks for all tested groups and declined over time. While the 469 M2ex3 trimer failed to maintain the GC reactions, the M2ex3 I3-01v9a SApNP induced durable 470 GC reactions lasting for 8 weeks (Fig. 7F). Both the frequency and number of GC B cells and $T_{\rm fh}$ 471 cells could be improved by a boost injection (Fig. 7G). Interestingly, a significant expansion of T_{fh} cells was noted for the FR and I3-01v9a SApNPs 2 weeks after the boost. Overall, the M2ex3
I3-01v9a SApNP elicited 5.7/1.1 times more GC B cells and 7.0/1.3 times more T_{fh} cells compared
with the soluble trimers at 8 weeks after the single-dose/prime-boost injections, respectively (Fig.
7F and 7G). In summary, our immunological analysis suggests that M2ex3 I3-01v9a SApNPs can
generate long-lived GC reactions in lymph nodes more effectively than individual M2ex3 trimers,
resulting in potent and long-lasting M2ex3-specific humoral immunity.

478 Antibody-dependent cell cytotoxicity and functional T cell responses

479 The non-neutralizing M2e-specific antibody responses were evaluated for functional activity using 480 a surrogate antibody-dependent cell cytotoxicity (ADCC) assay (see Methods). The M2ex3 I3-481 01v9a-immune sera elicited significantly higher relative light units (RLUs) in Jurkat effector cells 482 compared with the naïve, M2ex3 trimer, and M2ex3 FR groups, indicating detection of M2e 483 antibodies by mouse Fcy receptor IV (mFcyRIV) expressed on Jurkat cells (Fig. 8A). The largest 484 difference in ADCC activity between the I3-01v9a SApNP and other groups was observed at the 485 lowest serum dilution tested, 20×, with I3-01v9a SApNP showing 7.2-fold, 5.9-fold, and 2.1-fold 486 higher RLU than naïve, trimer, and FR groups, respectively. Thus, this assay confirmed that 487 M2ex3 immune sera binding to homotetrameric M2e on influenza-infected cells has the potential 488 to activate ADCC pathways, which is an important mechanism for M2e-mediated protection.

While antibody-mediated neutralization plays a key role in blocking virus infection, T cellmediated cellular immunity effectively reduces disease severity, hospitalization, and death rate (*87*, *88*). Enzyme-linked immunosorbent spot (ELISpot) analysis demonstrated that the M2ex3 I3-01v9a SApNP group produced significantly higher spot formation in bulk IFN-γ-secreting splenocytes stimulated with the M2ex3-5GS-foldon trimer probe compared to the naïve (> 28fold), M2ex3 trimer (7-fold), and M2ex3 FR (2.5-fold) groups per 8 × 10⁵ splenocytes (**Fig. 8B**).

Similarly, the M2ex3 I3-01v9a group also produced, on average, more spots in IL-4-secreting
splenocytes compared to the naïve (21.8-fold), M2ex3 trimer (1.7-fold), and M2ex3 FR (1.7-fold)
groups, although the findings were not statistically significant.

498 For analysis of specific T cell subsets, T cells from splenic tissue were divided into several 499 subsets, including CD4⁺ helper cells, which have multiple central roles in orchestrating adaptive 500 immune responses, and CD8⁺ cytotoxic T cells, which control virus infection by killing virus-501 infected cells and producing effector cytokines (89-92). Here, we designed a 13-color panel to 502 analyze functional T cell responses by measuring activation induced marker (AIM) and 503 intracellular cytokine staining (ICS) using flow cytometry. We compared various M2ex3 vaccine 504 constructs for the induction of CD4⁺ and CD8⁺ responses specific to the vaccine antigen at 5 days 505 after the PR8 (H1N1) challenge following a two-dose immunization regimen (Fig. 4A). Mouse 506 splenocytes from the M2ex3 trimer, FR, and I3-01v9a SApNP groups were stimulated with the 507 M2ex3-5GS-foldon trimer probe prior to analysis. All three M2ex3 constructs generated balanced 508 Th1 and Th2 responses and relatively lower Th17 responses (Fig. 8C, fig. S10). Among the three 509 vaccines, M2ex3 I3-01v9a SApNP was most effective in terms of the frequency and number of 510 intracellular cytokine (IFN-γ, TNF-α, IL-2, IL-17 and IL4)-producing CD4⁺ T cells, CD40L⁺CD4⁺ 511 T cells, and CD40L⁺CD4⁺ T cells that produce intracellular cytokines, as well as $T_{\rm fh}$ cells. 512 Importantly, M2ex3 I3-01v9a SApNP induced significantly more IFN-y-producing functional 513 $CD4^+$ T cells, which contained 2.7 times more activated ($CD40L^+$) T cells compared with the 514 M2ex3 trimer, resulting in a Th1-skewed response. Similarly, M2ex3 I3-01v9a SApNP elicited 515 more intracellular cytokine (IFN- γ , TNF- α , IL-2)-producing CD8⁺ T cells, CD69⁺CD8⁺ T cells, 516 and CD69⁺CD8⁺ T cells that produce IFN- γ than naïve, M2ex3 trimer, and M2ex3 FR groups (Fig. 8D). Overall, M2ex3 I3-01v9a SApNP induced stronger functional CD4⁺ and CD8⁺ T cell 517

responses than M2ex3 trimer and FR SApNP, consistent with the higher survival, lower weight
loss and viral loads (Fig. 4), and greater M2e-specific antibody responses (Fig. 5).

520 **DISCUSSION**

521 Since the 1918 Spanish flu, influenza has caused millions of deaths and hospitalizations worldwide 522 and remains a serious public health threat. From 2010 to 2017, seasonal flu caused 9.2 to 35.6 523 million reported cases of influenza and 140,000 to 710,000 hospitalizations. Each year, seasonal 524 flu causes an estimated 3-5 million cases of severe illness worldwide (1). Conventional influenza 525 vaccines, such as inactivated viruses produced in eggs, have been shown to significantly reduce disease burden. However, these vaccines mainly induce NAbs to viral epitopes that are prone to 526 527 antigenic drift, allowing viruses to evade vaccine-induced immune responses (17). As a result, 528 annual updates are necessary for seasonal flu vaccines, yet they still may not offer sufficient 529 protection. Additionally, recent evidence has revealed the negative impact of repeated antigen 530 exposure on vaccine efficacy (93). Therefore, it is important to develop a vaccine that can provide 531 broad and durable protection against diverse influenza viruses.

An effective M2e vaccine covering all IAVs and eliciting durable M2e antibody responses 532 533 may address these two goals for developing a universal pan-influenza A vaccine. M2e antibodies 534 are non-neutralizing but can engage alveolar macrophages and natural killer cells to promote viral 535 clearance via ADCC (94, 95). In principle, a successful M2e vaccine could act as a standalone 536 pan-influenza A vaccine, reducing the severity of disease caused by pandemic strains originating 537 from various animal reservoirs that contain novel HA subtypes against which the majority of the 538 human population lacks pre-existing immunity (17). Alternatively, M2e can be used to 539 complement seasonal influenza vaccines, significantly boosting the breadth of these strain-specific 540 inactivated virus vaccines (96). However, to date, an approved M2e vaccine remains elusive. Here,

we approached M2e vaccine development with a rational strategy, in which newly developed 1cSApNPs were used as multivalent carriers to display 60 copies of an M2e antigen to overcome the
intrinsically poor immunogenicity associated with soluble M2e and increase durability of M2especific immunity. Notably, we have previously developed vaccine candidates for HIV-1 (*57, 62*),
HCV (*60*), SARS-Cov-1/2 (*59*), and EBOV (*58*) based on the 1c-SApNP platform. In this study,

we first tested this strategy using hM2e and, based on the results, presented a tandem M2e antigen,
derived from human, avian, and swine IAVs, on FR and two multilayered 1c-SApNPs, E2p and

548 I3-01v9a. All the M2e vaccine constructs were evaluated in vitro and in vivo.

549 Our approach addresses the two limitations of previous M2e-based vaccine candidates in 550 clinical trials: 1) poor immunogenicity and 2) poor durability. First, the presentation of M2e on 551 the surface of 60-meric 1c-SApNPs significantly improves the immunogenicity of M2e. The large 552 size and high thermostability of the M2ex3-presenting I3-01v9a SApNP allows for long retention 553 in the lymph nodes follicles (8 weeks or longer), resulting in robust and prolonged GC reactions 554 compared to those elicited by the scaffolded M2e trimer. Combining the M2ex3 I3-01v9a SApNP 555 with commonly used adjuvants such as AP or AddaVax resulted in potent M2e-specific antibody 556 and functional T cell responses that likely conferred protection against sequential H1N1 and H3N2 557 challenges. In contrast, the inactivated PR8 (H1N1) virus vaccine and a strain-matched challenge 558 offered minimal protection against a follow-up heterologous challenge from a different subtype 559 (H3N2). Overall, our result suggest that the adjuvanted M2ex3 I3-01v9 SApNP can be developed 560 into a potent, durable, and cross-protective influenza vaccine that may overcome the limitations of 561 current marketed influenza vaccines, including the lack of protection against antigenically drifted 562 seasonal or novel pandemic strains (97). Additionally, this vaccine may have advantages over 563 previous M2e vaccine candidates in terms of effectiveness and durability.

564 Future studies will focus on several fronts. First, the lead candidate identified in this study, 565 M2ex3 I3-01v9a SApNP, may be formulated with more potent adjuvants to improve vaccine 566 efficacy. In a recent study, we tested the effect of various adjuvants on a SARS-CoV-2 spike 1c-567 SApNP vaccine and found that toll-like receptor 9 (TLR9) and stimulator of interferon genes 568 (STING) agonists significantly increased NAb titers compared to other adjuvants, including AP 569 and AddaVax (63). Second, an M2e/inactivated virus combination vaccine concept may warrant 570 further evaluation. While M2ex3 1c-SApNP can be used as a standalone vaccine, it may also be 571 combined with a seasonal flu vaccine to boost protection against non-vaccine-matched circulating 572 strains and potential pandemic strains. Third, an M2ex3 1c-SApNP vaccine may protect against 573 highly pathogenic H5N1 and H7N9 influenza strains. Lastly, less commonly explored influenza B 574 M2e (in tandem) can be incorporated onto influenza A M2ex3 1c-SApNP to potentially yield a 575 truly universal, single component influenza vaccine against both influenza A and B viruses. Given 576 the sequence conservation of M2e across HA subtypes, the M2e vaccines developed in this study 577 may protect against diverse IAVs, which account for $\sim 70\%$ of all influenza cases annually, thereby 578 significantly reducing the deaths and hospitalizations associated with influenza worldwide.

579 METHODS

580 Computational design of I3-01v9a for optimal nanoparticle display of monomeric antigens

We redesigned the N-terminus of I3-01v9 for optimal surface display of monomeric antigens, such as M2e, without using long linkers. Because I3-01v9 and I3-01 share nearly identical NP structures (57), the I3-01 structure (PDB ID: 5KP9) was used here for all modeling purposes. We manually extended the I3-01 N-terminal helix as an initial model. Briefly, an α -helix (residues 953-982) of the transcription factor protein c-MYC (PDB ID: 6G6L) was grafted onto an I3-01 subunit by using Glu2 and Glu3 of I3-01 for fitting. The grafted N-terminal helix was truncated to 11 residues

587 so that the first residue would be just above the surface after particle assembly. The CONCOORD 588 suite (64) was used to sample 1000 structures for the modified I3-01 subunit to facilitate ensemble-589 based protein design. Using default OPLS-UA parameters and a *damp* parameter of 0.1, geometric 590 constraints were generated by the program dist as input for the program disco to generate slightly 591 perturbed conformations (64). An ensemble-based design method used in our previous studies (57, 592 60) was employed to predict the first 9 of the 11 residues in the extended N-terminal helix using 593 C_{α} and C_{β} -based RAPDF scoring functions (65). Given a scoring function, Monte Carlo simulated 594 annealing minimization (57, 60) was performed to predict the amino acid composition for each of 595 the 1000 CONCOORD-derived conformations. For each position of the 9-residue helical segment, 596 the frequency of each amino acid type was calculated from the entire ensemble. The final design, 597 I3-01v9a, was determined manually by combining results from both scoring functions.

598 Expression and purification of various M2e immunogens

599 Rationally designed hM2e and tandem M2ex3 scaffolds and SApNPs were characterized in vitro 600 and in vivo. Scaffolded trimers and SApNPs were transiently expressed in ExpiCHOTM cells 601 (Thermo Fisher) using a previously described protocol (57, 58). Briefly, ExpiCHOTM cells were 602 thawed and incubated with ExpiCHOTM Expression Medium (Thermo Fisher) in a shaker incubator at 37°C, 135 rpm, and 8% CO₂. When cells reached a density of 10×10⁶ cell/ml, 603 ExpiCHOTM Expression Medium was added to reduce cell density to 6×10⁶ cell/ml for 604 transfection. ExpiFectamineTM CHO/plasmid DNA complexes were prepared for 100-ml 605 606 transfections in ExpiCHO cells following the manufacturer's instructions. For all constructs tested in this study, 100 µg of antigen plasmid and 320 µl of ExpiFectamineTM CHO reagent were mixed 607 608 in 7.7 ml of cold OptiPRO[™] medium (Thermo Fisher). After the first feed on day 1, ExpiCHO 609 cells were cultured in a shaker incubator at 32°C, 120 rpm, and 8% CO₂ following the Max Titer

610 protocol with an additional feed on day 5 (Thermo Fisher). Culture supernatants were harvested 611 13-14 days after transfection, clarified by centrifugation at 4000 rpm for 20 min, and filtered using 612 a 0.45 µm filter (Thermo Fisher). For all constructs, the M2e trimer or SApNP was extracted from 613 the culture supernatants by using mAb148 or mAb65 antibody columns. The bound protein was 614 eluted three times by 5 ml of glycine buffer (0.2M glycine, pH2.2) and neutralized by adding 0.375 615 ml Tris-base Buffer (2M Tris, pH9.0). Elutions were pooled and buffer exchanged via ultra-616 centrifugal filtration to phosphate buffer saline (PBS). The size of trimers and SApNPs was 617 analyzed by size exclusion chromatography using AKTA pure 25 (Cytiva). Trimer was purified 618 on a Superdex 75 Increase 10/300 GL column (GE Healthcare), whereas SApNPs were 619 characterized on a Superose 6 10/300 GL column. Protein concentration was determined using 620 UV₂₈₀ absorbance with theoretical extinction coefficients.

621 SDS-PAGE and BN-PAGE

622 The trimer and SApNPs were analyzed by sodium dodecyl sulfate-polyacrylamide gel 623 electrophoresis (SDS-PAGE) and blue native-polyacrylamide gel electrophoresis (BN-PAGE). 624 The proteins were mixed with loading dye and added to either a 10% Tris-Glycine Gel (Bio-Rad) or a 4-12% Bis-Tris NativePAGETM gel (Life Technologies). For SDS-PAGE under reducing 625 626 conditions, the proteins were first treated with dithiothreitol (DTT, 25 mM) and boiled for 5 min 627 at 100°C. SDS-PAGE gels were loaded with 1 µg of the sample and BN-PAGE gels were loaded 628 with 4 µg of the sample. SDS-PAGE gels were run for 20 min at 250 V using SDS running buffer (Bio-Rad), and BN-PAGE gels were run for 2-2.5 h at 150 V using NativePAGETM running buffer 629 630 (Life Technologies) according to the manufacturer's instructions. SDS-PAGE gels were stained 631 using InstantBlue (Abcam) and BN-PAGE gels were stained using Coomassie Brilliant Blue R-632 250 (Bio-Rad) and de-stained using a solution of 6% ethanol and 3% glacial acetic acid.

633 Differential scanning calorimetry

Thermal melting curves of trimer and SApNPs following mAb148 or mAb65 purification were obtained from a MicroCal PEAQ-DSC Man instrument (Malvern). Briefly, the purified SApNP protein was buffer exchanged into 1×PBS buffer and concentrated to 0.5-3 μ M prior to the analysis. Melting was probed at a scan rate of 60°C·h⁻¹ from 20°C to 100°C. Data processing, including buffer correction, normalization, and baseline subtraction, was conducted using MicroCal PEAQ-DSC software. Gaussian fitting was performed using Origin 9.0 software.

640 **Dynamic light scattering (DLS)**

Particle size distributions of hM2e and M2ex3 based on three NP platforms (FR, E2p-L4P, and I3-01v9a-L7P) were obtained from a Zetasizer Ultra instrument (Malvern). MAb148/SEC-purified NPs from ExpiCHO cells were diluted to 0.2 mg/ml using 1×PBS buffer, and 30 µl of the prepared NP sample was added to a quartz batch cuvette (Malvern, catalog no. ZEN2112). Particle size was measured at 25 °C in back scattering mode. Data processing was performed on the Zetasizer, and the particle size distribution was plotted using Origin 9.0 software.

647 Negative stain EM analysis

The initial evaluation of various SApNP samples was performed by the Core Microscopy Facility at The Scripps Research Institute. All SApNPs samples were prepared at the concentration of 0.005-0.02 mg/ml. Carbon-coated copper grids (400 mesh) were glow-discharged and 8 µl of each sample was adsorbed for 2 min. Excess sample was wicked away and grids were negatively stained with 2% uranyl formate for 2 min. Excess stain was wicked away and the grids were allowed to dry. Samples were analyzed at 120 kV with a Talos L120C transmission electron microscope

(Thermo Fisher) and images were acquired with a CETA 16M CMOS camera. All SApNP samples
purified by mAb148 were validated under 52,000 × magnification before further use.

656 **Bio-layer interferometry (BLI)**

657 The kinetics profiles of both hM2e and M2ex3 trimers and SApNPs were measured using an Octet 658 RED96 instrument (FortéBio, Pall Life Sciences) against mAb148 and mAb65 antibody. All 659 assays were performed with agitation set to 1000 rpm in FortéBio 1× kinetic buffer. The final 660 volume for all solutions was 200 µl per well. Assays were performed at 30°C in solid black 96-661 well plates (Geiger Bio-One). For all trimers and SApNPs, 5 µg/ml antibody in 1× kinetic buffer 662 was loaded onto the surface of anti-human Fc Capture Biosensors (AHC) for 300 s. A 60 s 663 biosensor baseline step was applied prior to the analysis of the association of the antibody on the 664 biosensor to the antigen in solution for 200 s. A two-fold concentration gradient of antigen, starting 665 at 25 nM for the hM2e trimer/SApNPs and 22 nM for the M2ex3 trimer/SApNPs, was used in a 666 titration series of six. The dissociation of the interaction was followed for 300 s. The correction of 667 baseline drift was performed by subtracting the mean value of shifts recorded for a sensor loaded 668 with antibody but not incubated with antigen, and for a sensor without antibody but incubated with 669 antigen. Octet data were processed by FortéBio's data acquisition software v.8.1. Peak signals at 670 the highest antigen concentration were summarized in a matrix to facilitate comparisons between 671 different vaccine platforms.

672 **Propagation of influenza viruses**

673 For challenge in mice, the following reagents were obtained through BEI Resources, NIAID, NIH:

674 Influenza A Virus, A/Puerto Rico/8/1934-Mouse Adapted (H1N1), NR-28652 and Influenza A

675 Virus, A/Hong Kong/1/1968-1 Mouse-Adapted 12 (H3N2), NR-28621. In brief, 4.4 × 10⁶ Madin-

676 Darby canine kidney (MDCK) cells (CCL-34TM; ATCC[®]) were plated overnight in 100 mm cell

culture dishes. The next day, cells were washed with PBS and incubated with a multiplicity of infection (MOI) of 1 for PR8 or HK/68 in serum free media for 1 hour. Next, cells were washed and 10 ml of serum-free DMEM containing 0.2% bovine serum albumin (BSA; VWR international) and 1 μ g/ml L-1-tosylamido-2-phenylethyl chloromethyl ketone (TPCK)-trypsin (Sigma Aldrich) was added to the dishes. Cells were incubated for 65 h after which the supernatant was collected, centrifuged at 4000 RPM for 10 min, aliquoted, and frozen at -80°C until use.

683 Immunoplaque assay to quantify influenza viruses

684 Virus PFU/ml of grown viruses was quantified using an immunoplaque assay. In brief, MDCK 685 cells were plated in 96-well plates at 25,000 cells/well. The cells were then washed with PBS and 686 infected with 10-fold serially diluted virus stocks. The inoculum was then removed, and cells were 687 overlayed with 0.7% low-melt agarose (Axygen) in serum-free DMEM containing 0.2% w/v BSA 688 and 1 µg/ml TPCK-trypsin. Twenty hours later, cells were fixed with 100 µl of 3.7% wt 689 formaldehyde for 1 h. Cells were then permeabilized with 50 µl ice-cold methanol for 20 min. 690 Fixed cells were then washed with deionized water and incubated with 50 µl of FluA-20 (non-691 pandemic H1N1 strains), 2D1 (CA 09 H1N1), or F045-092 (pandemic or seasonal H3N2 strains) 692 IgG (10 μ g/ml) for 1 h. Plates were washed and 50 μ l of 1:5000 diluted HRP-goat anti-human IgG 693 was added to the wells. Plates were then placed on a shaker at 225 rpm for 1 h. Cells were then 694 washed and 50 µl of TrueBlueTM Peroxidase Substrate (SeraCare) was added to wells and 695 incubated for 10-15 min for the development of plaques. Lastly, plates were washed with deionized 696 water and left to dry overnight. Plaques were quantified using a Bioreader[®] 7000 (BIOSYS 697 Scientific®).

698 Mouse immunization and challenge

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699 Six-to-eight-week-old female BALB/c mice were purchased from The Jackson Laboratory and 700 housed in ventilated cages in environmentally controlled rooms at The Scripps Research Institute, 701 in compliance with an approved IACUC protocol and Association for Assessment and 702 Accreditation of Laboratory Animal Care (AAALAC) international guidelines. Institutional 703 Animal Care and Use Committee (IACUC) guidelines were followed for all mice studies. Mice 704 were immunized at weeks 0 and 3, with 80 μ l of antigen/adjuvant mix containing 10 μ g of hM2e 705 or M2ex3 antigen in 40 µl PBS and 40 µl of adjuvant: aluminum phosphate (alum phosphate) or 706 AddaVax (InvivoGen). For the hM2e study, alum phosphate was used. For the M2ex3 study, both 707 alum phosphate and AddaVax were evaluated. The following reagent was obtained through BEI 708 Resources, NIAID, NIH: Influenza A Virus, A/Puerto Rico/8/1934 (H1N1), BPL-Inactivated, NR-709 19325, and used as a positive control for the first strain-matched challenge (3 μ g/mouse without 710 adjuvant). Each immunization dose was split amongst 4 footpads (20 µl/each). To assess the 711 protectivity and efficacy of the hM2e and M2ex3 constructs, immunized mice were sequentially 712 challenged with $LD_{50} \times 10$ of mouse-adapted A/Puerto Rico/8/1934 (PR8) H1N1 and A/Hong 713 Kong/1/1968 (HK/68) H3N2 at weeks 6 and 10, respectively. To establish the lethal dose of 50% 714 (LD₅₀) in mice for PR8 H1N1 and HK/68 H3N2, various dilutions of grown virus stock were 715 administered to mice (25 µl/nostril) after light anesthetization with isoflurane. Survival, weight 716 loss, and morbidity were monitored for 14 days. Mice that exhibited > 25% weight loss or showed 717 visible signs of distress were euthanized. Next, Reed-Muench and Spearman-Karber methods were 718 used to determine the 50% endpoint titer for both PR8 H1N1 and HK/68 H3N2 in mice (69, 70). 719 For first challenge at week 6, mice immunized with hM2e or M2ex3 constructs were lightly 720 anesthetized with isoflurane and $LD_{50} \times 10$ of PR8 H1N1 (50 µl) was administered to each mouse 721 (25 µl/nostril). Five mice from M2ex3 + AddaVax groups were sacrificed at Day 5 to assess the

722 viral load in lungs. In brief, mice were euthanized, and lungs were isolated. The lung tissue in PBS 723 was then crushed and spun down at 1200 rpm for 10 min. The lung supernatant was then aliquoted 724 and frozen at -80°C for future analysis. Viral loads were evaluated in lung supernatant using the 725 immunoplaque assay mentioned previously. Remaining mice were monitored for survival and 726 weight loss for 14 days post-challenge. At week 10, surviving mice were lightly anesthetized and 727 challenged with $LD_{50} \times 10$ HK/68 H3N2. Mice that exhibited > 25% weight loss or showed visible 728 signs of distress were euthanized. Blood of immunized mice was collected 2 weeks after each 729 immunization or challenge (weeks 2, 5, 10, and 14). All bleeds were performed through the facial 730 vein. Blood was spun down at 14,000 rpm for 10 min to separate serum from the rest of the whole 731 blood. The serum was then heat-inactivated at 56°C for 30 min and spun down at 8,000 rpm for 732 10 min to remove precipitates.

733 Enzyme-linked immunosorbent assay (ELISA)

734 For assessing hM2e-specific binding of hM2e-immune sera, 50 µl hM2e-5GS-foldon trimer probe 735 was coated on the surface of half-well 96-well high-binding polystyrene plates at a concentration 736 of 0.1 µg/well. Plates were kept at 4°C overnight. The next day, plates were washed 5× with PBS 737 containing 0.05% v/v Tween 20® (PBST). Plates were then blocked with 150 µl of 4% w/v nonfat 738 milk (Bio-Rad) for 1 h. Plates were then washed and 50 μ l of hM2e-immune sera was added to 739 each well for 1 h. Serum was diluted 20× in 4% nonfat milk followed by seven 10-fold dilutions. 740 M2e antibodies mAb148 and mAb65 were used as positive controls. Next, plates were washed and 741 50 µl of 1:3000 dilution horseradish peroxidase (HRP)-conjugated goat anti-mouse IgG in PBST 742 was added to the wells and incubated for 1 h. Plates were then washed and 50 ul of 1-StepTM 743 3,3',5,5'-tetramethylbenzidine (TMB; Thermo Fisher) substrate was added to each well and 744 incubated for 3 min. Plates were washed 6× and 50 µl of 2.0 N sulfuric acid (Aqua Solutions, Inc.)

745	was added to each well. Plates were then immediately read on a plate reader (BioTek Synergy)
746	using a wavelength of 450 nm. An identical ELISA method was used for M2ex3-specific binding
747	of M2ex3-immune sera, except for the use of M2ex3-5GS-foldon trimer probe.

748 Cell-based ELISA

749 For cell-based ELISA, the following 8 reagents were obtained through BEI Resources, NAID, 750 NIH: 1) Influenza A Virus, A/Puerto Rico/8/1934 (H1N1; NR-348), 2) Influenza A Virus, 751 A/California/04/2009 (H1N1; NR-136583), 3) Influenza A Virus, A/Solomon Islands/3/2006 752 (H1N1; NR-41798), 4) Influenza A Virus, A/Hong Kong/1/1968 (H3N2) (mother clone), ,NR-753 28620, 5) Influenza A virus, A/Brisbane/10/2007 (H3N2; NR-12283, 6) Influenza A virus, 754 A/Aichi/2/1968 (H3N2; NR-3177), 7) Influenza B Virus, B/Florida/4/2006 (Yamagata Lineage; 755 NR-41795), and 8) Influenza B Virus, B/Brisbane/60/2008 (Victoria Lineage; NR-42005). The 756 viruses were grown in MDCK cells using the same method mentioned previously for propagating 757 challenge strains. For cell-based ELISA, MDCK cells were plated overnight in 96-well cell culture 758 plates at a density of 18,000 cells/well. The next day the cells were washed and infected with 100 759 µl of 1 of the 8 viruses at a MOI of 1. Twenty hours later, the virus was removed, and cells were 760 washed before being fixed with 100 µl of 3.7% wt formaldehyde. Cells were then washed, and the 761 previous ELISA protocol was used except with an incubation step with TMB for 5 min.

762 Histology, immunostaining, and imaging

To study vaccine distribution, trafficking, retention, cellular interaction, and GC reactions in lymph nodes, M2ex3 trimer and FR and I3-01v9a SApNP immunogens formulated with AddaVax adjuvant were injected intradermally into four mouse footpads using 29-gauge insulin needles. Mice were anesthetized with 3% isoflurane in oxygen during immunization. Similar protocols of mouse injection, lymph node collection and tissue analysis were utilized from our previous study

768 (57, 63). The injection dose was 80 µl of antigen/adjuvant mix containing 40 µg of vaccine 769 immunogen per mouse or 10 µg per footpad. Mice were euthanized at 2 h to 8 weeks after a single-770 dose injection or 2 and 5 weeks after the boost, which occurred at 3 weeks after the first dose. 771 Brachial and popliteal sentinel lymph nodes were collected for immunohistological study. Fresh 772 lymph nodes were isolated and merged into frozen section compound (VWR International, catalog 773 no. 95057-838) in a plastic cryomold (Tissue-Tek at VWR, catalog no. 4565). Tissue samples were 774 frozen in liquid nitrogen and stored at -80°C before shipping to The Centre for Phenogenomics in 775 Canada for tissue processing, immunostaining, and imaging. Lymph node tissue sections were 776 sliced 8 µm thick on a cryostat (Cryostar NX70) and placed on a charge slide. Next, tissue slides 777 were fixed in 10% neutral buffered formalin and permeabilized in PBS buffer that contained 0.5% 778 Triton X-100 before immunostaining. The slides were blocked with protein Block (Agilent) to 779 prevent nonspecific antibody binding. Primary antibodies were applied on tissue slides and 780 incubated overnight at 4°C. After washing with tris-buffered saline with 0.1% Tween-20 (TBST), 781 secondary antibodies that we conjugated with either biotin or a fluorophore were utilized and 782 incubated at 25°C for 1 hour. Lymph node tissues were stained with anti-human Ab148 or Ab65 783 (1:200), and biotinylated goat anti-human secondary antibody (Abcam, catalog no. ab7152, 1:300), 784 followed by streptavidin-horseradish peroxidase (HRP) reagent (Vectastain Elite ABC-HRP Kit, 785 Vector, catalog no. PK-6100) and diaminobenzidine (ImmPACT DAB, Vector, catalog no. SK-786 4105).

To study cellular interactions between M2ex3 trimer immunogens and cell components in lymph nodes, FDCs were labelled using anti-CD21 primary antibody (Abcam, catalog no. ab75985, 1:1800), followed by anti-rabbit secondary antibody conjugated with Alexa Fluor 555 (Thermo Fisher, catalog no. A21428, 1:200). Subcapsular sinus macrophages were labeled using

791 anti-sialoadhesin (CD169) antibody (Abcam, catalog no. ab53443, 1:600), followed by anti-rat 792 secondary antibody conjugated with Alexa Fluor 488 (Abcam, catalog no. ab150165, 1:200). B 793 cells were labeled using anti-B220 antibody (eBioscience, catalog no. 14-0452-82, 1:100), 794 followed by anti-rat secondary antibody conjugated with Alexa Fluor 647 (Thermo Fisher, catalog 795 no. A21247, 1:200). GC reactions induced by M2ex3 trimers and SApNPs were studied by 796 immunostaining. GC B cells were labeled using rat anti-GL7 antibody (FITC; BioLegend, catalog 797 no. 144604, 1:250). T_{fh} cells were labeled using anti-CD4 antibody (BioLegend, catalog no. 798 100402, 1:100), followed by anti-rat secondary antibody conjugated with Alexa Fluor 488 799 (Abcam, catalog no. ab150165, 1:1000). GC forming cells were stained using Bcl6 antibody 800 (Abcam, catalog no. ab220092, 1:300), followed by anti-rabbit secondary antibody conjugated 801 with Alexa Fluor 555 (Thermo Fisher, catalog no. A21428, 1:1000). Nuclei were labeled using 802 4',6-diamidino-2-phenylindole (DAPI) (Sigma-Aldrich, catalog no. D9542, 100 ng/ml). The 803 immunostained lymph node tissues were scanned using an Olympus VS-120 slide scanner with a 804 Hamamatsu ORCA-R2 C10600 digital camera. The vaccine transport and induced GC reactions 805 in lymph nodes were quantified through bright-field and fluorescent images using ImageJ 806 software.

807 Lymph node disaggregation, cell staining, and flow cytometry

GC reactions in terms of frequency and numbers of GC B cells ($GL7^+B220^+$) and T_{fh} cells ($CD3^+CD4^+CXCR5^+PD-1^+$) were characterized using flow cytometry (**fig. S9**). Mice were euthanized at 2, 5, and 8 weeks after a single-dose injection or 2 and 5 weeks after the boost, occurring at 3 weeks after the first dose (4 footpads, 10 µg/footpad). Fresh axillary, brachial, and popliteal sentinel lymph nodes were collected for GC study. Lymph node tissues were disaggregated mechanically and merged in enzyme digestion solution in an Eppendorf tube with
814 958 µl of Hanks' balanced salt solution (HBSS) buffer (Thermo Fisher Scientific, catalog no. 815 14185052), 40 µl of 10 mg/ml collagenase IV (Sigma-Aldrich, catalog no. C5138), and 2 µl of 10 816 mg/ml DNase (Roche, catalog no. 10104159001) and incubated at 37°C for 30 min. The lymph 817 node tissue was then filtered through a 70 µm cell strainer to obtain a single cell suspension. Cell 818 samples were spun down at 400 \times g for 10 min and the cell pellets were resuspended in HBSS 819 blocking buffer containing 0.5% (w/v) bovine serum albumin and 2 mM EDTA. Anti-CD16/32 820 antibody (BioLegend, catalog no. 101302) was added into the Eppendorf tube to block the 821 nonspecific binding on Fc receptors. Cells were kept on ice for 30 min and transferred to 96-well 822 V-shaped-bottom microplates with pre-prepared cocktail antibodies, including Zombie NIR 823 live/dead stain (BioLegend, catalog no. 423106), Brilliant Violet 510 anti-mouse/human 824 CD45R/B220 antibody (BioLegend, catalog no. 103247), FITC anti-mouse CD3 antibody 825 (BioLegend, catalog no. 100204), Alexa Fluor 700 anti-mouse CD4 antibody (BioLegend, catalog 826 no. 100536), PE anti-mouse/human GL7 antibody (BioLegend, catalog no. 144608), Brilliant 827 Violet 605 anti-mouse CD95 (Fas) antibody (BioLegend, catalog no. 152612), Brilliant Violet 421 anti-mouse CD185 (CXCR5) antibody (BioLegend, catalog no. 145511), and PE/Cyanine7 anti-828 829 mouse CD279 (PD-1) antibody (BioLegend, catalog no. 135216). Cells were mixed with antibody 830 cocktail and placed on ice for 30 min. Cell samples were spun down at $400 \times g$ for 10 min and the 831 cell pellets were resuspended in HBSS blocking solution for washing one more time. Cells were 832 then fixed with 1.6% paraformaldehyde (Thermo Fisher Scientific, catalog no. 28906) in HBSS 833 and placed on ice for 30 min. Cells were spun down at $400 \times g$ for 10 min and placed in HBSS 834 blocking buffer at 4°C before test. Sample events were acquired using a 5-laser AZE5 flow 835 cytometer (Yeti, Bio-Rad) with Everest software at the Core Facility of The Scripps Research 836 Institute. The data were analyzed using FlowJo 10 software.

837 ADCC (antibody-dependent cell cytotoxicity) surrogate assay

838 The potential for M2e-specific antibodies to induce antibody-dependent cell cytotoxicity (ADCC) 839 was evaluated using a mouse FcyRIV ADCC Reporter kit (Promega). In brief, MDCK cells were 840 plated in white 96-well plates overnight at 18,000 cells/well. The next day, cells were washed with 841 PBS and infected with PR8 H1N1 at a MOI of 1. Twenty hours later, the cells were washed and 842 25 µl of M2ex3-immune sera was added to the wells for 1 h (sera was diluted 20× followed by 10-843 fold dilutions). Next, as per the kit's instructions (Promega), 75,000 mouse FCyRIV Jurkat cells 844 were added to each well. The plates were incubated for 6 h at 37°C. Lastly, 75 µl of Bio-Glo[™] 845 Reagent were added to the well. Relative light units were measured using a plate reader after a 5-846 min incubation of each plate.

847 Splenocyte isolation

848 At week 14, mice were anesthetized using isoflurane and euthanized using cervical dislocation. 849 Spleens were harvested from mice and kept in PBS on ice. Next, spleens were crushed with the 850 back of a syringe and resuspended in 20 ml of PBS. Cells were centrifuged at 1200 rpm for 10 851 min. Next, supernatant was discarded, and 3 ml of ACK lysis buffer (Lonza) was added to the cells 852 and incubated for 5 min. Next 12 ml of PBS was added to the tubes, which were then centrifuged 853 at 1200 rpm for 5 min. Supernatant was then discarded, and cells were resuspended in 1 ml PBS. 854 Cells were passed through a 70-µm cell strainer. Cells were then centrifuged for 5 min. Lastly, 855 cells were resuspended in 10% DMSO in FBS, transferred into a -80°C freezer overnight, and then 856 stored in the vapor phase of liquid nitrogen until analysis.

857 Enzyme-linked immunosorbent spot (ELISpot) assay

858 For analyzing IFN-γ and IL-4-secreting splenocytes of mice immunized with M2ex3, ELISpot was

859 used. First, Multiscreen® filter plates (Millipore Sigma) were coated with capture IFN-γ or IL-4

860 (BD Biosciences) at a 1:200 dilution. The plates were incubated at 4 °C overnight. The next day, 861 plates were washed and blocked with 200 µl of complete RPMI 1640 (10% FBS, 1% Penn-Strep, 862 1% L-glutamine; Gibco) medium. Next, the M2ex3-5GS-foldon trimer probe was prepared at 50 863 µg/ml in complete RPMI. Concanavalin A (10 µg/ml; BD Biosciences) was prepared as a positive 864 control. Next, spleen samples were thawed, resuspended in RPMI and centrifuged at 1200 RPM 865 for 10 min. Cells were then counted using an automated cell counter (Countess II; Thermo Fisher) 866 and suspended to reach a concentration of 1.6×10^7 cells/ml. Next, RPMI was discarded from the 867 filter plates and 50 µl of antigen was added to the well. Next, 50 µl of cell suspension was added 868 to each well, producing a final cell concentration of 8×10^5 splenocytes/well. After addition of 869 cells, the final concentrations of the M2ex3-5GS-foldon trimer probe and Concanavalin A were 870 2.5 µg/well and 0.5 µg/well, respectively. Cells without antigen were used as a negative control. 871 Plates were incubated for 48 h at 37°C. Next, cells were washed 2× with deionized water followed 872 by 3 washes with PBST. Next, 50 µl detection IFN-y or IL-4 antibodies (BD Biosciences) diluted 873 1:250 in dilution buffer (10% FBS in PBS) were added to corresponding wells. Plates were then 874 incubated for 2 h at room temperature (RT). Next, plates were washed 3× with PBST and 50 µl 875 Streptavidin-HRP (BD Biosciences) diluted 1:100 in dilution buffer was added to the well and 876 incubated for 1 h. Plates were then washed 4× with PBST and 2× with PBS. AEC Final Substrate 877 (BD Biosciences) was then added to the wells for 15-20 min for the development of spots. Plates 878 were kept in the dark overnight and read using a Bioreader[®] 7000.

879 **T cell culture and stimulation**

Functional M2e-specific T cells responses were characterized by measuring activation induced marker (AIM) and intracellular cytokine staining (ICS) using flow cytometry (**fig. S10**). Mouse splenocytes were isolated from naïve or vaccinated mice at 5 days after prime-boost immunizations

883 and followed by a H1N1 virus challenge. Cryopreserved splenocytes were thawed by diluting cells 884 in 10 ml pre-warmed complete RPMI media with 10% deactivated fetal bovine serum (FBS) and 885 1% penicillin/streptomycin (P/S). Cells were spun down at $400 \times g$ for 10 min and cell pellets were 886 resuspended in RPMI media. The numbers of splenocytes were counted and adjusted to 10 million 887 cells/ml. One million splenocytes for each mouse were placed into 96-well U-shaped-bottom 888 microplates. Cells were cultured in the presence of the M2ex3-5GS-foldon trimer probe (1 µg per 889 well) at 37°C for a total of 24 h. After 20 h, Brefeldin A Solution (BioLegend, catalog no. 420601) 890 was added in the culture to enhance intracellular cytokine staining signals by inhibiting the protein 891 transport processes in the rough endoplasmic reticulum and Golgi apparatus. After 4 h, cells were 892 then spun down at $400 \times g$ for 10 min and cell pellets were resuspended in HBSS blocking buffer. 893 Anti-CD16/32 antibody (BioLegend, catalog no. 101302) was added for 30 min on ice to block 894 nonspecific binding to Fc receptors. Cells were then transferred to 96-well V-shaped-bottom 895 microplates with pre-prepared cocktail antibodies for surface marker staining, including 896 LIVE/DEAD Fixable Blue Dead Cell Stain Kit (Thermo Fisher Scientific, catalog no. L34962), 897 FITC anti-mouse CD3 antibody (BioLegend, catalog no. 100204), Alexa Fluor 700 anti-mouse 898 CD4 antibody (BioLegend, catalog no. 100536), BUV737 Anti-Mouse CD8a antibody (BD 899 Bioscience, catalog no. 612759), APC anti-mouse CD154 antibody (BioLegend, catalog no. 900 106510), Brilliant Violet 421 anti-mouse CD69 Antibody (BioLegend, catalog no. 104527), 901 APC/Fire 810 anti-mouse CD279 (PD-1) antibody (BioLegend, catalog no. 135251), and Brilliant 902 Violet 605 anti-mouse CD185 (CXCR5) Antibody (BioLegend, catalog no. 145513). Splenocytes 903 were mixed with antibody cocktail and placed on ice for 30 min. Cells were spun down at $400 \times g$ 904 for 10 min and the cell pellets were resuspended in HBSS blocking solution, then washed once 905 more. Cells were fixed with 1.6% paraformaldehyde (Thermo Fisher Scientific, catalog no. 28906)

906 in HBSS and placed on ice for 30 min. Cell were then washed two times with intracellular staining 907 permeabilization wash buffer (BioLegend, catalog no. 421002) and stained with previously 908 prepared antibody cocktail for intracellular staining, including PE anti-mouse IFN-γ antibody 909 (BioLegend, catalog no. 505808), Brilliant Violet 785 anti-mouse TNF-α antibody (BioLegend, 910 catalog no. 506341), PE/Cyanine5 anti-mouse IL-2 antibody (BioLegend, catalog no. 503824), 911 PE/Cyanine7 anti-mouse IL-4 antibody (BioLegend, catalog no. 504118), and Brilliant Violet 711 912 anti-mouse IL-17A Antibody (BioLegend, catalog no. 506941). Cells were mixed with 913 intracellular antibodies and placed on ice for 30 min. Cells were then washed again with 914 intracellular staining permeabilization wash buffer. Cells were stored in HBSS blocking buffer at 915 4°C prior to analysis. Sample events were acquired using a Cytek Aurora spectral analytical flow 916 cytometer with SpectroFlo software at the Flow Cytometry Core Facility of The Scripps Research 917 Institute. The data were analyzed using FlowJo 10 software.

918 Statistical analysis

Data was collected from 8-13 mice per group in the immunization studies, challenge experiments, and sera binding assays. For the vaccine transport and GC study in lymph nodes and T cells in spleens, 5 mice per group with different vaccine constructs were compared using one-way ANOVA, followed by Tukey's multiple comparison *post hoc* test. Statistical significance is indicated as the following in the figures: ns (not significant), *p < 0.05, **p < 0.01, ***p < 0.001. ****p < 0.0001. The graphs were generated using GraphPad Prism 9.3.1 software.

925 SUPPLEMENTARY MATERIALS

- 926 Supplementary material for this article is available at XXX.
- 927 Supplementary Figures 1-10.

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1165 Figure Legends

1166 Figure 1. Design and in vitro characterization of hM2e immunogens. (A) Structural models of

1167 human M2e (hM2e), hM2e-5GS-1TD0 trimer, and three hM2e-presenting nanoparticles (NPs).

1168 Left: Amino acid sequence and ribbons/surface model of hM2e (from PDB ID 4N8C). Middle:

- 1169 Ribbons/surface model of hM2e-5GS-1TD0 trimer, in which a trimeric viral capsid protein SHP
- 1170 (PDB ID: 1TD0) is used to display hM2e. Right: Surface models of hM2e on 24-meric ferritin
- 1171 (FR) and 60-meric E2p-L4P and I3-01-L7P 1c-SApNPs. The SApNP size is indicated by diameter
- 1172 (in nm). (B) SEC profiles of hM2e-5GS-1TD0 trimer and hM2e-presenting FR, E2p-L4P and I3-
- 1173 01-L7P SApNPs. The hM2e trimer was processed on a Superdex 75 10/300 increase GL column,
- 1174 while three SApNPs were processed on a Superose 6 increase 10/300 GL column. (C) SDS-PAGE
- 1175 under reducing conditions (left) and BN-PAGE (98) of hM2e-presenting FR, E2p-L4P and I3-

1176 01v9a-L7P SApNPs. Notably, hM2e-5GS-1TD0 is included on the SDS gel for comparison. (D)

1177 Negative-stain EM micrographs of mAb148-purified FR, E2p-L4P and I3-01v9a-L7P SApNPs.

- 1178 (E) DLS profiles of mAb148-purified FR, E2p-L4P and I3-01v9a-L7P SApNPs. Average particle
- size derived from DLS are labeled. (F) Thermostability of the hM2e-5GS-1TD0 trimer and hM2e-
- 1180 5GS-FR SApNP with Tm, $\Delta T_{1/2}$ and T_{on} measured by DSC. (G) ELISA analysis of the hM2e trimer
- and SApNPs (FR, E2p-L4P and I3-01v9a-L7P) binding to mAb65 (left) and mAb148 (98) after
- 1182 heating to 50 °C, 60 °C and 70 °C for 15 minutes. (H) Antigenic profiles of the hM2e trimer and
- 1183 SApNPs (FR, E2p-L4P and I3-01v9a-L7P) to mAb148 using BLI.

1184 Figure 2. Assessment of hM2e scaffolds and nanoparticles in a mouse challenge model. 1185 (A) Benchmark challenge studies assessing survival and weight loss to establish the 50% lethal 1186 intranasal challenge dose in mice for mouse-adapted A/Puerto Rico/8/1934 (PR8) H1N1 and 1187 A/Hong Kong/1/1968 (HK/68) H3N2; N = 10 mice/group. Mice were monitored for survival, 1188 weight loss, and morbidities for 14 days. (B) Schematic representation of mouse immunization 1189 regimen for hM2e constructs, sequential challenges of $LD_{50} \times 10$ of PR8 and HK/68, blood 1190 collection, and sacrifice; N = 10 mice/group. (C) Survival and weight loss of mice challenged with 1191 $LD_{50} \times 10$ of PR8 (H1N1) followed by an $LD_{50} \times 10$ of HK/68 (H3N2). Mice were monitored for 1192 survival, weight loss, and morbidities for 14 days. (D) ELISA curves showing hM2e-immune sera 1193 binding to the hM2e-5GS-foldon trimer probe and calculated 50% effective concentration (EC₅₀) 1194 values for weeks 2, 5, 10, and 14. The assay was performed in duplicate with a starting serum 1195 dilution of 20x followed by seven 10-fold titrations. Images of mouse immunization, virus 1196 challenge, and blood and organ collection created with BioRender.com.

Figure 3. Design and characterization of tandem M2ex3 immunogens. (A) Structural models
of tandem M2ex3, M2ex3-5GS-1TD0 trimer, and three M2ex3-presenting SApNPs. Left: Amino

1199 acid sequences of human, avian/swine, and human/swine M2e and ribbons/surface model of 1200 tandem M2ex3 (based on hM2e from PDB ID 4N8C). The G4 linker is shown as a dotted line. 1201 Middle: Ribbons/surface model of M2ex3-5GS-1TD0 trimer, in which 1TD0 is a trimeric viral 1202 capsid protein. Right: Surface models of M2ex3 on 24-meric ferritin (FR) and 60-meric E2p-L4P 1203 and I3-01-L7P 1c-SApNPs. The SApNP size is indicated by diameter (in nm). (B) SEC profiles of 1204 M2ex3-5GS-1TD0 trimer and M2ex3-presenting FR, E2p-L4P and I3-01-L7P SApNPs. The 1205 tandem M2ex3 trimer and three SApNPs were processed on a Superdex 75 10/300 increase GL 1206 column and a Superose 6 increase 10/300 GL column, respectively. (C) SDS-PAGE (left) under 1207 reducing conditions and BN-PAGE (98) of tandem M2ex3-presenting FR, E2p-L4P and I3-01v9a-1208 L7P SApNPs. Notably, M2ex3-5GS-1TD0 is included on the SDS gel for comparison. (D) 1209 Negative-stain EM micrographs of mAb148-purified FR, E2p-L4P and I3-01v9a-L7P SApNPs. 1210 (E) DLS profiles of mAb148-purified FR, E2p-L4P and I3-01v9a-L7P SApNPs. Average particle 1211 size derived from DLS are labeled. (F) Thermostability of the M2ex3-5GS-1TD0 trimer and 1212 M2ex3-5GS-FR SApNP with Tm, $\Delta T_{1/2}$ and T_{on} measured by DSC. (G) ELISA analysis of the 1213 M2ex3 trimer and SApNPs (FR, E2p-L4P and I3-01v9a-L7P) binding to mAb65 (left) and 1214 mAb148 (98) after heating to 50 °C, 60 °C and 70 °C for 15 minutes. (H) Antigenic profiles of the 1215 M2ex3 trimer and SApNPs (FR, E2p-L4P and I3-01v9a-L7P) to mAb148 using BLI.

1216 Figure 4. Survival and weight loss assessment of tandem M2ex3 scaffold and nanoparticles

in a mouse challenge model. (A) Schematic representation of mouse immunization regimen for M2ex3 constructs, sequential intranasal challenges of $LD_{50} \times 10$ of mouse-adapted PR8 (H1N1) and HK/68 (H3N2), blood collection, and sacrifice. Groups were as follows: M2ex3 groups adjuvanted with alum phosphate (n = 8), M2ex3 groups adjuvanted with AddaVax (n = 13), and Inactivated PR8 (n = 8). Inactivated PR8 + AddaVax (n = 13) was used as a positive control for

1222 lung viral titers for the PR8 challenge. (B) Survival and weight loss of mice challenged with LD50 1223 \times 10 of PR8 (H1N1) followed by an LD₅₀ \times 10 of HK/68 (H3N2). Mice were monitored for 1224 survival, weight loss, and morbidities for 14 days. (C) Lung viral titers in M2ex3-immunized mice 1225 at Day 5 post-PR8 H1N1 challenge (n = 5). Visual representation of plaques formed from the lung 1226 supernatants of various M2ex3-immunized mice. The highest countable plaques were observed in 1227 naïve mice. The lowest number of plaques were observed in the lung supernatants of E2p- and I3-1228 01v9a NP-immunized mice. The assay was performed in duplicate starting at a lung supernatant 1229 dilution of 1x followed by 10-fold titrations. Statistical analysis shows significance between 1230 M2ex3 groups compared to naïve mice using One-way ANOVA. The error bars indicate 1231 mean \pm standard deviation; **p < 0.01, ***p < 0.001, and ****p < 0.0001. Images of mouse 1232 immunization, virus challenge, and blood and organ collection created with BioRender.com.

Figure 5. M2ex3-immune sera binding to scaffolded M2ex3 and homotetrameric M2e. (A) 1233 1234 ELISA curves showing M2ex3-immune sera (adjuvanted with alum phosphate or AddaVax) 1235 binding to the M2ex3-5GS-foldon trimer probe and calculated 50% effective concentration (EC₅₀) 1236 values for weeks 2, 5, 10, and 14. N = 8 or 13 at weeks 2 and 5. N = variable based on surviving 1237 mice/group post-challenge for weeks 10 and 14. The assay was performed in duplicate with a 1238 starting serum dilution of 20× followed by seven 10-fold titrations. (B) Serum binding to M2e on 1239 the surface of MDCK cells infected with various influenza A strains. Strains: A/Puerto 1240 Rico/8/1934 (H1N1), A/California/04/2009 (H1N1)pdm09, A Solomon Islands/2/2006 (H1N1), 1241 A/Hong Kong/1/1968 (H3N2), A/Brisbane/10/2007 (H3N2), A/Aichi/2/1968 (H3N2), 1242 B/Brisbane/60/2008 (Flu B, Victoria Lineage B/Florida/4/2006), and (Flu B, Yamagata Lineage). 1243 MAb148 (M2e antibody) was used as a positive control. The assay was performed in duplicate 1244 with a starting serum dilution of 50x followed by five 10-fold titrations. Statistical analysis shows

significance between trimer and NP groups and positive control using Two-way ANOVA. The error bars indicate mean \pm standard deviation; *p<0.05, **p<0.01, ***p<0.001, and ****p<0.0001.

1248 Fig. 6. Prolonged retention of M2ex3-presenting SApNPs in lymph node follicles. (A) 1249 Distribution of I3-01v9a SApNPs displayed M2ex3 trimers in a lymph node at 48 h after a single-1250 dose injection (10 µg/injection, 40 µg/mouse). Anti-M2e Ab148 and Ab65 were used to stain the 1251 lymph node tissues. (B) schematics of M2ex3 trimer presenting SApNP accumulation in lymph 1252 node tissues. (C) Trafficking and retention of the M3ex3 trimer and FR and I3-01v9a SApNPs in 1253 lymph node follicles at 2 h to 8 weeks after a single-dose injection. Scale bar = 50 μ m for each 1254 image. (D) Time-dependent curve and (E) Area under the curve (78) of the Ab148-stained area in 1255 immunohistological images of M3ex3 immunogen retention in lymph node follicles up to 8 weeks. 1256 (F) Quantification of M3ex3 vaccine accumulation in lymph node follicles at 1 week after a single-1257 dose injection. (G) Histological images of the M2ex3 trimer and two SApNP vaccine accumulation 1258 and retention in lymph node follicles at 2 weeks and 5 weeks after the boost, which occurred at 3 1259 weeks after the first dose. (H) Quantification of vaccine accumulation in lymph node follicles at 1 1260 week after the boost. In mouse injection, all vaccine immunogens were mixed with AddaVax 1261 adjuvant. Data were collected from more than 10 lymph node follicles (n = 3-5 mice/group). (I) 1262 Interaction of M2ex3 trimer presenting SApNPs with FDC networks in lymph nodes at 1 week 1263 after a single-dose injection. Both FR and I3-01v9a SApNP immunogens were colocalized with 1264 FDC networks. Immunofluorescent images are pseudo-color-coded (CD21⁺, green; CD169⁺, red; 1265 Ab148, white). Scale bars = 500 and 100 μ m for a complete lymph node and enlarged image of a 1266 follicle, respectively. The data points are expressed as mean \pm SEM for (D) and SD for (E, F and

H). The data were analyzed using one-way ANOVA followed by Tukey's multiple comparison
post hoc test. ***p < 0.001, ****p < 0.0001.

1269 Fig. 7. Induction of robust and long-lived germinal center reactions by M2ex3-presenting 1270 SApNPs. (A) Top image: Immunofluorescent images of M2ex3 trimer presenting I3-01v9a 1271 SApNP vaccine candidate induced germinal centers (GCs) at 2 weeks after a single-dose injection 1272 (10 µg/injection, 40 µg/mouse). Bottom image: robust GC reaction with organized light zone (LZ) 1273 and dark zone (DZ) compartments in lymph node follicles. GC B cells ($GL7^+$, red) attached to 1274 FDCs (CD21⁺, green) and T_{fh} cells located in LZ of GCs. Scale bars = 500 and 100 μ m for a 1275 complete lymph node and the enlarged image of a follicle, respectively. (B) and (C) quantification 1276 of GCs in terms of the GC/FDC ratio and the size of GCs induced by the M2ex3 trimer, and FR 1277 and I3-01v9a SApNP vaccines using immunohistological images at 2, 5, and 8 weeks after a single-dose injection or at 2 and 5 weeks after the boost, which occurred at 3 weeks after the first 1278 1279 dose (n = 5 mice/group). (**D**) and (**E**) representative GC images induced by three M2ex3 vaccine 1280 constructs at 8 weeks using a single-dose or prime-boost regimen. Scale bar = $50 \,\mu\text{m}$ for the image 1281 of an enlarged lymph node follicle. (F) and (G) Quantification of GC reactions in terms of the 1282 percentage and number of GC B cells and T_{fh} cells using flow cytometry after a single-dose or prime-boost immunizations. In mouse immunization, all vaccine constructs were formulated with 1283 AddaVax adjuvant. The data points are shown as mean \pm SD. The data were analyzed using one-1284 1285 way ANOVA followed by Tukey's multiple comparison post hoc test for each timepoint. *p < 1286 0.05, **p < 0.01, ***p < 0.001, ***p < 0.0001.

1287 Figure 8. Innate and T cell responses of M2ex3 scaffolds and M2ex3-presenting SApNPs

1288 (A) ADCC activity measured by RLU of FcyRIV-expressing Jurkat effector cells binding to

1289 M2ex3-immune sera bound to M2e on PR8 (H1N1)-infected MDCK cells. The data is presented 1290 as mean \pm SEM (B) Spot formation of IFN- γ and IL-4-secreting splenoctyes from M2ex3-1291 immunized mice at Day 5 post-PR8 H1N1 challenge. Mouse splenocytes were isolated from 1292 immunized mice with M2ex3 trimer and FR and I3-01v9a SApNP vaccine constructs at 5 days 1293 after prime-boost immunization followed by H1N1 virus challenge (n = 5 mice/group). Splenocytes of naïve mice without immunization but with a H1N1 virus challenge were included 1294 1295 as control samples. Quantification of the percentage and number of vaccine-induced functional (C) 1296 $CD4^+$ and (D) $CD8^+$ T cell responses using flow cytometry. In mouse immunization, all vaccine 1297 immunogens were coupled with AddaVax adjuvant. The data points are shown as mean \pm SD. The 1298 data were analyzed using one-way ANOVA followed by Tukey's multiple comparison post hoc 1299 test for each timepoint. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.

1300 SUPPLEMENTAL LEGENDS

1301 **fig. S1.** Schematic of computational design for I3-01v9a.

- 1302 fig. S2. Construct and in vitro characterization of hM2e-presenting trimer and SApNPs.
- 1303 fig. S3. Serum binding of individual mice immunized with hM2e 1TD0 trimer and SApNPs.
- 1304 fig. S4. Construct and in vitro characterization of M2ex3-presenting trimer and SApNPs.
- 1305 fig. S5. Serum binding of individual mice immunized with M2ex3 1TD0 trimer and SApNPs.
- 1306 **fig. S6.** M2ex3 serum binding of individual mice to cell-surface homotetrameric M2e.
- 1307 fig. S7. Immunohistological images of M2ex3 1TD0 trimer and SApNPs in lymph nodes.
- 1308 fig. S8. Immunohistological analysis of M2ex3 1TD0 trimer and SApNP-induced GCs.
- 1309 fig. S9. Flow cytometry analysis of M2ex3 1TD0 trimer and SApNP-induced GCs.
- 1310 **fig. S10.** Flow cytometry analysis of M2ex3 1TD0 trimer and SApNP-induced T cell responses.

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design

CONCOORD perturbation

(1000 structures)

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DEEISQFCKEKGVFYMPGVMTPTELVKAMKLGHTIL

KLFPGEVVGPQFVKAMKGPFPNVKFVPTGGVNLDNV

CEWFKAGVLAVGVGSALVKGTPVEVAEKAKAFVEKI

RGCTE

fig. S1. Schematic representation of computational design for I3-01v9a. Top left: The molecular surface model (gray) of I3-01v9 NP and a zoomed-in view of the ribbon model of an I3-01v9 trimer (chains A, B, and C colored in sky blue, plum, and light green, respectively), with the N-termini forming a triangle of 50.5 Å. Also shown is the ribbon model of a c-MYC transcription factor protein (PDB ID: 6G6L), from which the backbone of a helix (residues 953-982, orange) was grafted onto an I3-01v9 subunit by using residues E2 and E3 (yellow) of I3-01v9 for fitting. Top right: A side view of the ribbon model of an I3-01v9 trimer with the fitted full helix backbone (30 residues) and truncated helix backbone (11 residues). Bottom right: 1,000 slightly perturbed backbone conformations of the modified I3-01v9 subunit with an extended N-terminal helix generated using CONCOORD, a protein structure sampling program. Bottom middle: Predicted amino acid (frequency, %) for each position of the first 9 residues using C α and C β -based RAPDF scoring functions. Bottom left: The final sequence design, termed I3-01v9a, with the predicted residues of the N-terminal helical extension colored in orange. The anchoring residues E2 and E3 are colored in yellow, and the last remaining turn of the N-terminal helix is colored in red.

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 IMPLEY JOSTILUS MAKKGLGCVULLCGAVFVSPSQEIHARERRGAR<mark>SR</mark>SLLTEVETPIRNEWGCRCNDSSDS<mark>AS</mark>GGGGSEVRIFAGNDPAHTATGSSGISSPTPALTPLMLDEATGKLVVWDGQKAGSAVGILVLPLEGTETALTYYK: TFATEAIHWPESVDEHKKANAFAGSALSHAA

>hM2e-5GS-FR

DAMKRGLCCVLLLCGAVFVSPSQEIHARFRRGAR<mark>SR</mark>SLLTEVETPIRNEWGCRCNDSSD<mark>AS</mark>GGGGS<mark>ASGDIIKLLNEQVNKEMQSSNLYMSMSSWCYTHSLDGAGLFLFDHAABEYEHAKKLIIFLNENNVPVQLT ISAPEHKFEGLTQIFQKAYEHEQHISESINNIVDHAIKSKDHATFNFLQWYVAEQHEEEVLFKDILDKIELIGNENHGLYLAD</mark>

>hM2e-5GS-E2p-LD4-PADRE(hM2e-E2p-L4P)

DAMKRGLCCVLLLCGAVEVSPSQEIHARFRRGAR<mark>SR</mark>SLLTEVETPIRNEWGCRCNDSSD<mark>AS</mark>GGGGSGAAAKPATTEGEFPETREKMSGIRRAIAKAMVHSKHTAPHVTLMDEADVTKLVAHRKKFKAIAAEKGIKLT FLPYVVKALVSALREYPVLNTAIDDETEEIIQKHYYNIGIAADTDRGLLVPVIKHADRKPIFALAQEINELAEKARDGKLTPGEMKGASCTITNIGSAGGQWFTPVINHPEVAILGIGRIAEKPIVRDGEIVAAPML ALSLSFDHRMIDGATAQKALNHIKRLLSDPELLLMGGGGSFSEEQKKALDLAFYFDRRLTPEWRRYLSQRLGLNEEQIERWFRRKEQQIGWSHPQFEKGS<mark>RKFVAAWTLKRAA</mark>

>hM2e-5GS-I3-01v9a-LD7-PADRE(hM2e-I3-01v9a-L7P)

DAMKRGLCCVLLLCGAVEVSSQETHARFRGAR<mark>E</mark>SLITEVETFIRDEWGCRCNDSDAS</mark>GGGSGARLABELCKKMELFKHKIVAVLRANSVEEAKEKALAVFEGGVHLLEITFTVPDADTVIKELSFLKEKG AIIGAGVTSVEQCRKAVESGAEFIVSPHLDAEITVECLEKGVFYMFGVMTPTELVKAMKLGHNILKLFPGEVVGPQFVKAMKGPFPNVKFVPTGGVNLDNVCEWFKAGVLAVGVGSALVKGTPDEVREKAKAFVEK IFGCTEGGGSS PAVDIGDRIDELKALEALSAEDGHDDVGGRLESLIRRNNSRRADSS





fig. S2. Construct and in vitro characterization of hM2e-presenting trimer and SApNPs. (A) Construct sequences of hM2e-presenting FR, E2p-L4P, and I3-01v9-L7P SApNPs, with the gene fragments of leader sequence, restriction site, human Matrix protein 2 extra-virion domain (residue: 2-24), flexible linker, NP-forming subunit, locking domain (LD), and PADRE highlighted in yellow, green, gray, light magenta, cyan, olive green, and red shades, respectively. (B) negative stain EM image for hM2e-presenting SApNPs. (C) ELISA curves of for hM2e-presenting SApNPs binding to mAb148 and mAb65 antibody. (D) Dynamic Light Scatter results for hM2e-FR after SEC purification. (E) Antigenic evaluation of hM2e-presenting SApNPs using BLI for mAb65 antibody. A two-fold concentration gradient of antigen, starting at 5.0 uM for hM2e 1TD0 trimer, 80.0 nM for FR-SApNP, and 20.0 nM for E2p and I3-01v9a SApNPs, was used in a titration series of six.

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fig. S3. Serum binding of individual mice immunized with hM2e 1TD0 trimer and SApNPs. ELISA curves showing hM2e-immune sera (adjuvanted with aluminum phosphate) binding to the hM2e-5GS-foldon trimer probe and calculated 50% effective concentration (EC_{50}) values for weeks 2 and 5 (N = 10). N = variable at weeks 10 and 14 based on surviving mice/group post-H1N1 and H3N2 challenges at weeks 6 and 10, respectively. Serum was tested was from mice immunized with hM2e 1TD0 trimer, and FR, E2p, and I3-01v9a SApNP at weeks 0 and 3. The assay was performed in duplicate with a starting serum dilution of 20x followed by seven 10-fold titrations.

M2ex3-5GS--1TD0

DAMKRGLCCVLLLCGAVFVSPSQEIHARFRRGARSESLLTEVETPIRNEWGCRCNDSSDGGGGSLLTEVETPTRNGWESKSSDSSDGGGGSLLTEVETPTRSEWESRSSGSSD<mark>AS</mark>GGGS<mark>EVRIFAGNDPAHTATGS</mark> SGISSPTPALTPLMLDEATGKLVVWDGOKAGSAVGILVLPLEGTETALTYYKSGTFATEAIHWPESVDEHKKANAFAGSALSHAA

>M2ex3-5GS-FR

<mark>DAMKRGLCCVLLLCGAVFVSPSQEIHARFRRGARSE</mark>SLLTEVETPIRNEWGCRCNDSSDGGGGSLLTEVETPIRNGWESKSSDSSDGGGGSLLTEVETPIRSEWESRSSGSSD<mark>AS</mark>GGGSS<mark>GDIIKLLNEQVNKEMQ</mark>S SNLYMSMSSWCYTHSLDGAGLFLFDHAAEEYEHAKKLIIFLNENNVPVQLTSISAPEHKFEGLT<u>Q</u>IFQKAYEHEQHISESINNIVDHAIKSKDHATFNFLQWYVAEQHEEEVLFKDILDKIELIGNENHGLYLAD

>M2ex3-5GS-E2p-LD4-PADRE (M2ex3-E2p-L4P)

DAMKRGLCCVLLLCGAVFVSPSQEIHARFRRGAR<mark>SR</mark>SLLTEVETPIRNEWGCRCNDSSDGGGGSLLTEVETPTRNGWESKSSDSDGGGGSLLTEVETPTRSEWESRSSGSSD<mark>AS</mark>GGGSG<mark>GAAKPATTEGEFPETF</mark> EKMSGIRRAIAKAWVHSKHTAPHVTLMDEADVTKLVAHRKKKKAIAAEKGIKLTFLPYVVKALVSALREYPVLNTAIDDETEEIIQKHYYNIGIAADTDRGLLVPVIKHADRKPIFALAGEINELAEKARDGKLTPG EMKGASCTITNIGSAGGWFTPVINH PEVAILGIGRIAEKPIVRDGEIVAAPMLALSLSFDHRMIDGATAQKALNHIKRLLSDPELLLMGGGGS<mark>FSEEQKKALDLAFYFDRRLTPEWRRYLSQRLGINEEQIERWFF</mark> RKEQQIGWSHPOFEEGS<mark>MEFVAAWIKAAR</mark>

>M2ex3-5GS-I3-01v9a-LD7-PADRE(M2ex3-I3-01v9a-L7P)

DAMKRGLCCVLLLCGAVFVSPSQEIHARFRRGAR<mark>SR</mark>SLLTEVETPIRNEWGCRCNDSSDGGGGSLLTEVETPTRNGWESKSSDSDGGGGSLLTEVETPTRSEWESRSSGSSD<mark>AS</mark>GGGSAKLAEELQKKMEELFKK HKIVAVLRANSVEEAKEKALAVFEGGVHLIEITFTVPDADTVIKELSFLKEKGAIIGAGTVTSVEQCRKAVESGAEFIVSPHLDAEITVFCLEKGVFYMPGVMTPTELVKAMKLGHNILKLFPGEVVGPQFVKAMKG PFPNVKFVPTGGVNLDNVCEWFKAGVLAVGVGSALVKGTPDEVREKAKAFVEKIRGCTEGGGGS<mark>SPAVDIGDRLDELEKALEALSAEDGHDDVGQRLESLLRRWNSRRAD</mark>GS<mark>AKEVAAWILKAAR</mark>



MZex3	Primary Antibody Binding EC ₅₀ (µg/ml)	
Groups	m A b65	mAb148
M2ex3-5GS-1TD0 4°C	0.005851	0.009314
M2ex3-5GS-FR 4°C	0.003391	0.002564
M2ex3-5GS-FR 50°C	0.003373	0.00249
M2ex3-5GS-FR 60°C	0.002864	0.002555
M2ex3-5GS-FR 70°C	0.003176	0.002486
M2ex3-5GS-E2p-L4P 4°C	0.003277	0.003288
M2ex3-5GS-E2p-L4P 50°C	0.003149	0.003302
M2ex3-5GS-E2p-L4P 60°C	0.003209	0.003274
M2ex3-5GS-E2p-L4P 70°C	0.003013	0.00337
M2ex3-5GS-I3-01v9a-L7P 4*C	0.003325	0.002538
M2ex3-5GS-I3-01v9a-L7P 50°C	0.002818	0.0026
M2ex3-5GS-I3-01v9a-L7P 60°C	0.003039	0.002729
M2ex3-5GS-I3-01v9a-L7P 70°C	0.002895	0.002561

70 °C

60 °C



fig. S4. Design and in vitro characterization of M2ex3-presenting SApNPs. (A) Construct sequences of tandem M2epresenting FR, E2p-L4P, and I3-01v9-L7P NPs, with the gene fragments of leader sequence, restriction site, tandem Matrix protein 2 extra-virion domain (including human, avian/swine and human/swine), flexible linker, NP-forming subunit, locking domain (LD), and PADRE highlighted in yellow, green, gray, light magenta, cyan, olive green, and red shades, respectively. (B) negative stain EM image for M2ex3-presenting SApNPs. (C) ELISA curves of for M2ex3-presenting SApNPs binding to mAb148 and mAb65 antibody. (D) Antigenic evaluation of M2eX3-presenting SApNPs using BLI for mAb65 antibody. A two-fold concentration gradient of antigen, starting at 5 uM for M2ex3 1TD0 trimer, 80 nM for FR SApNP, and 20 nM for E2p and I3-01v9a SApNPs, was used in a titration series of six.

4 °C





fig. S5. Serum binding of individual mice immunized with M2ex3 1TD0 trimer and SApNPs. (A) ELISA curves showing M2ex3-immune sera (adjuvanted with aluminum phosphate or AddaVax) binding to the M2ex3-5GS-foldon trimer probe and calculated 50% effective concentration (EC_{50}) values for weeks 2, 5, 10, and 14. N = 8 or 13 at weeks 2 and 5. N = variable at weeks 10 and 14 based on surviving mice/group post-H1N1 and H3N2 challenges at weeks 6 and 10, respectively. Serum was tested was from mice immunized with M2ex3 1TD0 trimer, and FR, E2p, and I3-01v9a SApNP at weeks 0 and 3. The assay was performed in duplicate with a starting serum dilution of 20x followed by seven 10-fold titrations. (B) ELISA curves showing hM2e (+ AP)- and M2ex3 (+ AV)-immune sera binding to the hM2e-5GS-foldon trimer probe at week 5. N = 10 for hM2e groups. N = 13 for M2ex3 groups. Results indicate that M2ex3-immune sera demonstrates similar or higher binding to hM2e foldon compared to hM2e-immune sera. The assay was performed starting at a serum dilution of 20x followed by seven 10-fold titrations.





fig. S6. M2ex3 serum binding of individual mice to cell-surface homotetrameric M2e. Week 5 serum from mice immunized with M2ex3 1TD0, FR SApNP, and I3-01v9a SApNPs adjuvanted with AddaVax (n = 13) were tested against homotetrameric M2e on various influenza strains: A/Puerto Rico/8/1934 (H1N1), A/California/04/2009 (H1N1)pdm09, A Solomon Islands/2/2006 (H1N1), A/Hong Kong/1/1968 (H3N2), A/Brisbane/10/2007 (H3N2), A/Aichi/2/1968 (H3N2), B/Brisbane/60/2008 (Flu B, Victoria Lineage), and B/Florida/4/2006 (Flu B, Yamagata Lineage). MAb148 (M2e antibody) was used as a IAV positive control. The assay was performed in duplicate with a starting serum dilution of 50x followed by five 10-fold titrations.



B Single-dose - 2 weeks





Single-dose - 8 weeks

D



fig. S7. Immunohistological images of M2ex3 1TD0 trimer and SApNPs in lymph nodes. Immunostaining images of M2ex3 1TD0 and M2ex3-presenting FR and I3-01v9a SApNP interaction with FDC networks in lymph node follicles at (A) 48 hours, (B) 2 weeks, (C) 5 weeks, and (D) 8 weeks after a single-dose injection (10 μ g per injection, totaling 40 μ g per mouse). Immunofluorescent images are pseudo color coded (CD21+, green; CD169+, red; anti-M2ex3 mAb148, white). Scale bars = 500 and 100 μ m for complete lymph node and enlarged image of a follicle, respectively. Data were collected from more than 10 lymph node follicles (n = 3-5 mice/group).
А	Single-dose - 2 w			fig. S
M2ex3 1TD0				
	•	-		
M2ex3 FR SApNP				
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M2ex3 I3-01v9a SApNP				
				(
В	Single-dose - 5 w			
M2ex3 1TD0				
			C	
M2ex3 FR SApNP				
M2ex3 I3-01v9a SApNP				
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Single-dose - 8 w





fig. S8. Immunohistological analysis of M2ex3 1TD0 trimer and SApNP-induced GCs. Immunohistological images of GCs at (A) 2, (B) 5, and (C) 8 weeks after a single-dose injection of M2ex3 1TD0 and M2ex3-presenting FR, and I3-01v9a SApNP vaccines (10 μ g per injection, totaling 40 μ g per mouse), with a scale bar of 500 μ m for each image. Images of GCs at (D) 2 and (E) 5 weeks after the boost, which occurred at 3 weeks after the first dose (n = 5 mice/group).



fig. S9. Flow cytometry analysis of M2ex3 1TD0 trimer and SApNP-induced GCs. Gating strategy for analyzing GCs (GC B cells and T follicular helper cells) using flow cytometry (n = 5 mice/group).



fig. S10. Flow cytometry analysis of M2ex3 1TD0 trimer and SApNP-induced T cell responses. Gating strategy for analyzing CD4+ and CD8+ T cell responses against the M2ex3-5GS-foldon probe using flow cytometry at 5 days after prime-boost immunization and virus challenge (n = 5 mice/group).







А









