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Computationally designed haemagglutinin with nanocage plug-and-display elicits pan-H5 influenza vaccine responses

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ABSTRACT

The increasing spread of highly pathogenic avian influenza (HPAI) A/H5 viruses poses a pandemic threat. Circulating clade 2.3.4.4b viruses have demonstrated rapid transcontinental dissemination, extensive reassortment, epizootic spread and potential sustained mammal-to-mammal transmission, signifying a heightened risk of becoming a human pathogen of high consequence. A broadly protective, future-proof vaccine against multiple clades of H5 influenza is urgently needed for pandemic preparedness. Here, we combine two novel vaccine technologies to generate a Digitally Immune Optimised and Selected H5 antigen (DIOSvax-H5_{inter}) displayed multivalently on the mi3 nanocage using the SpyTag003/SpyCatcher003 conjugation system. Mice immunized with DIOSvax-H5_{inter} Homotypic Nanocages at low doses demonstrate potent, cross-clade neutralizing antibody and T cell responses against diverse H5 strains. DIOSvax-H5_{inter} Homotypic Nanocages provide a scalable vaccine candidate with the potential for pan-H5 protection against drifted or newly emergent H5 strains. This World Health Organization preferred characteristic is essential for prospective strategic stockpiling in the pre-pandemic phase.

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KEYWORDS Avian influenza; universal influenza vaccine; pandemic preparedness; protein engineering; virus-like particles; nanoparticles; computational biology

Introduction

Highly pathogenic avian influenza (HPAI) viruses are characterized by the acquisition of a polybasic cleavage site in their surface glycoprotein haemagglutinin (HA) [1,2]. Recognition and cleavage by ubiquitous cellular proteases permit productive infection beyond the respiratory tract, facilitating systemic infection and increased viral pathogenicity [1,2]. HPAI viruses of the H5 subtype of the A/goose/Guangdong/1/1996 (gs/GD) lineage originated in domestic poultry in southeast Asia [2]. Evolution of the HA gene and multiple reassortment with low pathogenicity avian influenza viruses (LPAI) enabled adaptation for transmission of clade 2 viruses to wild aquatic birds that expanded the geographic range of the virus within the continent [3]. Since 2018, re-emergence of a novel reassortant of clade 2.3.4.4b in migratory bird populations with enhanced genetic plasticity and host adaptation has led to a global panzootic with infections in over 150 avian and 50 mammalian species [4]. Potential sustained mammal-to-mammal transmission and human infections from marine and terrestrial mammals, including food producing dairy cattle, highlight the urgent need for pandemic preparedness [4].

A critical task of pandemic preparation is stockpiling effective vaccines [5]. Based on global surveillance of circulating HPAI A/H5 isolates, the World Health Organization provides an annually updated list of representative candidate vaccine virus (CVV) strains [5]. Strain-specific inactivated vaccines have been traditionally stockpiled for zoonotic influenza viruses

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[6,7]. mRNA [8–10] and viral-vectored vaccine candidates [11] matched to the latest CVV H5 A/Astrakhan/3212/2020 (H5Ast20) also have been developed, which can be manufactured on a reduced timescale in response to a pandemic. However, as recently observed for the coronavirus disease 2019 (COVID-19) pandemic, accelerated evolution of RNA viruses in an expanding host reservoir can lead to drastic antigenic changes, resulting in vaccine mismatch [12]. Adaptive mutations that facilitate humanto-human transmission, such as greater HA stability and binding to human a2,6-linked sialic acids and enhanced viral polymerase function, may be anticipated, but the exact composition of mutations in a specific spillover event cannot be precisely predicted [4]. A broad H5 vaccine offering cross-clade protection would effectively circumvent the need for perfect antigenic match with an emergent strain with pandemic potential.

Here, we combined the Digitally Immune Optimised Synthetic Vaccine (DIOSynVax) antigen design technology [13–17] with the Plug-and-Display Spy-Tag003/SpyCatcher003-mi3 nanoassembly platform [18,19] to generate a computationally designed H5 nanocage vaccine. Using the DIOSynVax computational pipeline, we produced a novel, recombinant H5 immunogen with antigenic determinants representative across different clades of the H5 subtype, DIOSvax-H5_{inter} [16,17]. DIOSvax-H5_{inter} was immunized as a soluble protein or coupled to a self-assembling Homotypic Nanocage and compared to the CVV H5Ast20 as a Homotypic Nanocage in mice. After two doses, only the group immunized with DIOSvax-H5_{inter} Homotypic Nanocages elicited potent neutralization of all 12 tested H5 clades and subclades, as well as robust CD4⁺ T cell responses against two antigenically distant H5 clades that emerged either before or after the antigen was designed. Our results suggest the DIOSvax-H5_{inter} Homotypic Nanocage could be employed as a broadly protective stockpiling vaccine that offers variant-proof protection against the prospective threat of an H5 avian flu pandemic in humans before it begins.

Results

In silico design of pan-H5 antigen

A computationally-optimized H5 antigen, termed DIOSvax-H5_{inter}, was generated through the DIOSynVax phylogeny-based computational pipeline for designing broadly reactive immunogens [14,16,17]. Nucleotide sequences of influenza HA proteins were downloaded from the Global Initiative on Sharing All Influenza Data (GISAID) EpiFlu[™] database [20] (June 2018) and used as input to generate a multiple sequence alignment. The resulting alignment was used as input for phylogenetic tree reconstruction. The resulting tree was further analysed using HyPhy [21] to reconstruct the statistically-inferred and phylogenetically-optimized DIOSvax-H5_{inter} design (Figure 1(a) and S1).

Comparison with wild-type viral isolates from 12 major clades and subclades of the H5 subtype shows that DIOSvax-H5_{inter} is positioned phylogenetically between clades 2.1.3.2 and 2.2 (Figure 1(b)). DIOSvax-H5_{inter} has the highest sequence similarity of 98.4% to the clade 2.2 A/whooper swan/Mongolia/244/2005 strain and the lowest sequence similarity of 83.9% to the LPAI American non-gs/GD lineage A/chicken/Mexico/07/2007 strain (Figure S2). The DIOSvax-H5_{inter} polypeptide sequence differs from the circulating clade 2.3.4.4b CVV H5Ast20 strain at 41 residues spanning both head and stem domains (Figure 1(c)).

Preparation and characterization of computationally-designed H5 nanocages

mi3 is a computationally-designed, self-assembling 60-mer protein nanocage, which can be highly efficiently expressed and purified in a scalable fashion from Escherichia coli [24,25]. Up to ~100 mg/L Spy-Catcher003-mi3 can be produced in shaker flask cultures, which can be further increased substantially by industrial-scale fermentation [26]. The nanocage shows high stability against heat and freeze-thaw and can be readily lyophilized for long-term storage [25,27]. Genetic fusion of mi3 to SpyCatcher003 enables high efficiency coupling through spontaneous formation of an isopeptide bond to SpyTag003-antigens [19] of various symmetries [26]. Multivalent display presents antigens in highly repetitive arrays that recapitulate features common to viral surfaces. This arrangement enhances immune potency through mechanisms including enhanced uptake by antigenpresenting cells and efficient cross-linking by B cell receptors [28,29]. This Plug-and-Display nanocage platform has been used to develop influenza vaccines [26] and pan-sarbecovirus vaccines with potent immunogenicity in animal models [30–32].

We genetically fused the DIOSvax-H5_{inter} ectodomain or the H5Ast20 ectodomain to a T4 bacteriophage fibritin foldon [33] flanked by two flexible glycine-serine linkers and C-terminal SpyTag003 and His₆ tags [26] in a pEVAC vector backbone [34] (Figure 1(d)). Fusion to the fibritin foldon domain promotes HA trimerisation to mimic their natural conformation on virion surfaces. The characteristic polybasic cleavage sites of HPAI A/H5 viruses were converted to a monobasic form to stabilize HA in the pre-fusion conformation and prevent uncontrolled cleavage by constitutive proteases present in the mammalian Expi293FTM expression cell line [35].



Figure 1. In silico design and assembly of H5 Nanocage vaccines. (A) Structural models of HA antigens. Conservation of residues between full-length DIOSvax-H5_{inter} (H5DIOS) (blue) and H5Ast20 (green, GenBank accession: UJS29065.1) mapped onto the van der Waals surface of HA homotrimers. Residue substitutions are represented in pink. Structural prediction was performed by AlphaFold 3 [22] and displayed using PyMOL. (B) Phylogenetic tree of HA amino acid sequence identity of H5 viruses used in this study. (C) Amino acid sequence alignment of full-length DIOSvax-H5_{inter} and H5Ast20 by Clustal Omega v.1.2.4, numbered according to DIOSvax-H5_{inter} [23]. *indicates a stop codon. The polybasic cleavage sites were converted to monobasic forms in the final vaccine constructs. Major H5 antigenic sites are highlighted: A (purple), B (pink) and D (tan) in the receptor-binding domain; C (red) and E (grey) in the esterase and fusion domains; broadly neutralizing epitopes in the stem domain (green). (D) Genetic organization of DIOSvax-H5_{inter}-SpyTag003 and H5Ast20-SpyTag003 constructs, indicating the HA ectodomain, T4 foldon, tag and linker locations. (E) Schematic of Plug-and-Display nanocage vaccine assembly of DIOSvax-H5_{inter} Homotypic Nanocage.

Spontaneous isopeptide bond formation This involved deletion of residues 341-345 (RRRKK) for DIOSvax-H5_{inter} and residues 341-344 (KRRK) for H5Ast20. Both HA constructs were efficiently expressed, secreted and affinity-purified via the terminal His₆ tag.

Purified HA-SpyTag003 were coupled to Spy-Catcher003-mi3 (Figure 1(e)) by mixing the two proteins at different molar ratios for 24 h at 4°C. Minimal residual unconjugated HA-SpyTag003 were observed for 1:1 and 3:2 coupling ratios following SDS-PAGE/Coomassie (Figure 2(a and b)), indicating that coupling was occurring predominantly through all three SpyTag003 moieties. The diffuse HA-Spy-Tag003 bands were consistent with the natural heterogeneity of HA glycosylation. This was confirmed by digestion with Peptide N-Glycosidase F (PNGase F), which removed N-linked glycans and caused a decrease in apparent molecular weight and sharpening of the bands (Figure 2(c)). Dynamic light scattering (DLS) demonstrated an increase in hydrodynamic



Figure 2. Production and characterization of H5 Nanocage vaccines. (A) Titration of coupling of HA-SpyTag003 to SpyCatcher003mi3 Nanocage at different molar Nanocage:HA ratios, analysed by SDS–PAGE/Coomassie. (B) Validation of the DIOSvax-H5_{inter} (H5DIOS) Homotypic Nanocage and H5Ast20 Homotypic Nanocage vaccine lots by SDS–PAGE/Coomassie. (C) Analysis of HA-Spy-Tag003 with SDS–PAGE/Coomassie, with or without PNGase F de-glycosylation. Molecular weight markers are in kDa. (D) Dynamic light scattering of SpyCatcher003-mi3 nanocages. The mean hydrodynamic diameter is shown ± 1 s.d.; n = 3. (E) Transmission electron micrographs of negative-stained SpyCatcher003-mi3 nanocages. 50,000× magnification, scale bar = 20 nm.

diameter for SpyCatcher003-mi3 coupled to either HA immunogen, relative to the nanocage on its own (Figure 2(d)). Transmission electron microscopy (TEM) confirmed the integrity of both coupled nano-cages (Figure 2(e)).

To characterize the antigenicity of vaccine immunogens, we measured binding using an enzyme-linked immunosorbent assay (ELISA) to immobilized H5 soluble proteins or Homotypic Nanocages by the broadly neutralizing anti-HA monoclonal antibody CR9114 [36] and by matched or closely matched mouse sera generated in-house from previous vaccination studies (Figure S3). CR9114 is a broadly neutralizing antibody targeting a conserved, 3-dimensional epitope in the HA stem of all influenza A viruses [36]. Matched or closely matched mouse sera should contain a mixture of antibodies that are head- or stem-specific, subtypespecific or broad, and neutralizing or non-neutralizing. Binding to CR9114 and matched or closely matched sera was observed for all three immunogens, which indicated that the HA proteins were correctly folded and multiple epitopes were exposed and accessible for binding. We noticed a substantial decrease in the maximal OD₄₅₀ and IgG titre area under curve (AUC) for the DIOSvax-H5_{inter} antigen when expressed on Homotypic Nanocages, likely due to reduced accessibility by binding antibodies during multivalent display.

Computationally-designed HA nanocages elicit TNF⁺IFNy⁻CD4⁺ T cell responses against multiple H5 clades

We compared the immunogenicity of DIOSvax- $H5_{inter}$ and H5Ast20 Homotypic Nanocages against DIOSvax- $H5_{inter}$ Soluble protein. Doses were normalized by the number of HA trimers to allow comparison between soluble and multimerised antigens and facilitate an equimolar amount of Spy-Catcher003-mi3 nanocages with similar occupancy for both Homotypic Nanocages. Groups of six mice were immunized intramuscularly at weeks 0 and 4 with 0.5 µg of HA-SpyTag003 antigen adjuvanted 1:1 with AddaVax. Terminal bleed and splenocyte harvest were performed at week 8 (Figure 3(a)).

Cross-reactive T cell responses targeting multiple conserved viral proteins have been shown to play important roles in mitigating influenza infection-associated pathology and protecting against severe disease [37,38]. Typically, CD8⁺ T cells are responsible for direct killing of infected cells, while CD4⁺ T cells predominantly augment CD8⁺ T cell functions and antibody responses during viral infections [37,38]. Interestingly, influenza-specific CD4⁺ cells have also been shown to mediate viral clearance [37,38]. HPAI A/H5 HAspecific CD4⁺ and CD8⁺ T cells were measured by

intracellular cytokine staining for tumour necrosis factor (TNF) and interferon gamma (IFNy). Following ex vivo stimulation with commercially-available peptide pools of the clade 2.1.3.2 A/Indonesia/CDC835/2006 and the clade 2.3.4.4b A/Aves/Guanajuato/CPA-18539-23/2023 strains, all three vaccine groups induced significant TNF⁺IFN γ ⁻CD4⁺ cells to both clades, compared to the Uncoupled Nanocage control (Figure 3(b)). Interestingly, mice immunized with DIOSvax-H5_{inter} Soluble antigens induced significantly higher percentages of TNF⁺IFNy⁻CD4⁺ than the other two vaccine groups, indicating superior performance as a T cell immunogen. Antigen-specific T cells were not induced above background levels for all other assessed phenotypes (Figure S4). TNF has been associated with potent anti-influenza activity, including increased production of cytokines and chemokines [39] and stronger inhibition of viral replication than IFNy [40] by lung epithelial cells. Thus, the vaccine-induced cytokines could play functionally important roles as a critical part of the host responses that contribute to overall protection.

Computationally-designed HA nanocages neutralize multiple H5 clades

Neutralizing antibodies that block virus entry into host cells are well established as the primary immune correlate of protection for influenza vaccine effectiveness [41]. In recent years, pseudotype-based neutralization assays have become widely accepted as a reliable surrogate for microneutralisation assays using live viruses [34,42]. We tested neutralization of boosted sera at terminal bleed against a panel of H5 lentiviral pseudotypes [34,42] from the 12 selected clades and subclades (Figure 4). H5Ast20 Homotypic Nanocages elicited potent neutralization against the matched clade 2.3.4.4b and the closely related 2.3.4.4a and 2.3.4.4c pseudotypes, but failed to induce significant neutralization against several of the other tested pseudotypes when compared to the Uncoupled Nanocage control (Figure 4 and compiled in Figure 5).

In contrast, DIOSvax-H5_{inter} Homotypic Nanocages induced significant neutralization against all of the tested pseudotypes. Every mouse immunized with the DIOSvax-H5_{inter} Homotypic Nanocage demonstrated neutralization of every tested H5 pseudotype with $ID_{50} > 2 \times$ the limit of detection (Figure 4). DIOSvax-H5_{inter} Soluble antigens elicited high neutralization comparable to DIOSvax-H5_{inter} Homotypic Nanocage against pseudotypes from clade 1, 2.1.3.2, 2.2, 2.3.2.1a and the LPAI lineage EU non-gs/GD, but failed to consistently neutralize the circulating 2.3.4.4b and its closely related 2.3.4.4a, 2.3.4.4c and 2.3.4.4h strains (Figure 4). Our results show that the



Figure 3. Broad CD4⁺ T cell response against antigenically distant H5 clades. (A) Schedule of immunization and blood sampling of mice. (B) HPAI A/H5 HA-specific T cell responses quantified by TNF⁺IFN γ ⁻CD4⁺T cells. Splenocytes were collected from mice immunized with DIOSvax-H5_{inter} Soluble (dark blue), DIOSvax-H5_{inter} Homotypic Nanocage (light blue), H5Ast20 Homotypic Nanocage (green) or Uncoupled Nanocage (grey). Splenocytes were stimulated ex vivo with peptide pools for the HA of either the clade 2.1.3.2 A/Indonesia/CDC835/2006 or the clade 2.3.4.4b A/Aves/Guanajuato/CPA-18539-23/2023 strain. Cytokine production was measured by intracellular cytokine staining and flow cytometry. Each dot represents one mouse. The mean is denoted by a bar \pm 1 s.d.; *n* = 6. Statistical significance was calculated by ANOVA, followed by Tukey's multiple comparison post hoc test of % gated cells and plotted as **P* < 0.05, ***P* < 0.01, ****P* < 0.001.

DIOSvax-H5_{inter} Homotypic Nanocages, developed by combining two synthetic vaccine technologies, induced significantly broader neutralizing antibody responses than either the multivalent display (H5Ast20 Homotypic Nanocage) or the computational design (DIOSvax-H5_{inter} Soluble) alone (Figure 5). Potent neutralization by DIOSvax-H5_{inter} Homotypic Nanocages across all three main lineages of influenza HPAI and LPAI A/H5 viruses provides validation of unprecedented pan-H5 reactivity by our forward-predicting vaccine design technology.

Discussion

The ongoing panzootic of clade 2.3.4.4b HPAI A/H5 viruses has propelled global efforts in pre-pandemic vaccine design and stockpiling [43]. Historically stock-piled, U.S.-licensed inactivated clade 1 and clade 2.1.3.2 vaccines demonstrated limited cross-reactivity against H5Ast20, with 25-64% of tested vaccinee sera reaching protective haemagglutination inhibition titres after multiple adjuvanted high-dose immunisations [6]. Recent development of clade 2.3.4.4b inactivated, recombinant protein [7] and high-dose [8,9]



Figure 4. Broad neutralizing antibody response against diverse H5 clades. Neutralization of influenza pseudotypes by boosted mouse sera. Clade 1, 2.1.3.2, 2.2, 2.3.2.1a, 2.3.4.4a, 2.3.4.4b, 2.3.4.4c, 2.3.4.4h, 4, 7.1, AM non-gs/GD and EU non-gs/GD pseudotypes were tested for neutralization by terminal bleed sera from mice immunized with DIOSvax-H5_{inter} (H5DIOS) Soluble (dark blue), DIOSvax-H5_{inter} Homotypic Nanocage (light blue), H5Ast20 Homotypic Nanocage (green) or Uncoupled Nanocage (grey). Each dot represents one mouse, showing the serum dilution giving 50% inhibition of infection (ID₅₀). Dashed horizontal lines represent the limit of detection of $\log_{10}(ID_{50}) = 1.9$. The mean is denoted by a horizontal line ± 1 s.d.; n = 6. Statistical significance was calculated by ANOVA, followed by Tukey's multiple comparison post hoc test of $\log_{10}(ID_{50})$ values and plotted as *P < 0.05, **P < 0.01, ***P < 0.001, ***P < 0.001.

or self-amplifying [10] mRNA vaccines conferred homologous protection. Although cross-clade binding antibodies were detected, the absence of neutralizing antibodies rendered sterilizing immunity unlikely [8–10]. Multivalent mRNA-lipid nanoparticles designed as universal influenza priming vaccines are under clinical development, which are not expected to induce protective levels of neutralizing antibody responses against heterologous viruses [44]. Even with the streamlined manufacturing process of mRNA vaccines, these strain-specific approaches are only applicable post hoc upon the precise



Figure 5. Further comparison of H5 neutralization. (A) Neutralization of influenza pseudotypes by boosted mouse sera grouped by immunogen. Mice were immunized with DIOSvax-H5_{inter} (H5DIOS) Soluble, DIOSvax-H5_{inter} Homotypic Nanocage, H5Ast20 Homotypic Nanocage or Uncoupled Nanocage. Clade 1 (pink), 2.1.3.2 (orange), 2.2 (purple), 2.3.2.1a (brown), 2.3.4.4a (blue), 2.3.4.4b (lime), 2.3.4.4c (grey), 2.3.4.4h (aquamarine), 4 (yellow), 7.1 (violet), AM non-gs/GD (light blue) and EU non-gs/GD (magenta) pseudotypes. The mean is shown ± 1 s.d.; n = 6. Statistical significance is presented in Figure 4. (B) Percent of mice that responded to the specified H5 pseudotype. Response was defined as $ID_{50} > 2 \times$ the limit of detection.

characterization of a pandemic strain. Hence, vaccines based on virus isolates of the past remain vulnerable to rapid and inevitable antigenic mismatches due to the highly mutagenic nature of influenza viruses.

To mitigate the uncertainty of zoonotic spillover in the pre-pandemic phase, next-generation vaccine technologies deploy future-proofing immunogens that offer broader protection. Antigenic cartography analysis of subclades 2.3.4.4a-h identified a recombinant strain, denoted H5-Re11 Q115L/R120S/A156 T, that was antigenically representative at clade-level only [45]. Multi-clade protection against matched virus challenge has also been achieved in chickens with a virus-like particle vaccine without evidence of broader protection [46]. More recently, viral-vectored vaccines based on parainfluenza virus 5 [11] and Herpesvirus of Turkeys have demonstrated limited mismatched, cross-clade protection within the gs/GD lineage. However, pan-H5 breadth, particularly against more distantly related 4 and 7.1 clades and the LPAI lineages, has not been reported [11,45,46].

Here, our integrated approach combined computational antigen design and multivalent nanocage

delivery, both of which individually elicited consistent, broadly protective responses against different rapidly evolving RNA viruses in multiple animal models [13–15,30–32]. A pan-sarbecovirus vaccine developed using the DIOSynVax technology [14] and a human cytomegalovirus vaccine developed using the Spy-Tag/SpyCatcher technology have demonstrated adequate safety in rabbit toxicology screening and are currently in clinical trials in the UK [47]. Leveraging these established platforms, we have created a forward-designed and retrospective-validated avian influenza vaccine candidate, which induced robust pan-H5 humoral and cellular immunogenicity against diverse H5 clades. The DIOSvax-H5_{inter} Homotypic Nanocage represents a scalable, dose-sparing vaccine candidate for pre-pandemic stockpiling, which provides a proactive, robust response to newly emergent H5 strains with pandemic potential. As the virus continues to evolve, novel H5 strains are incorporated in the iterative DIOSynVax pipeline of computational design and immune optimization for updating H5 vaccine candidates. Importantly, due to the modular nature of the SpyTag003/SpyCatcher003-mi3 platform, updated DIOSvax-H5 sequences can be rapidly engineered for multivalent nanocage delivery to provide potent and durable protection against the changing antigenic landscape of H5 viruses.

Materials and methods

Phylogenetic analysis

Influenza HA nucleotide sequences were downloaded from the GISAID database [20] and employed for multiple sequence alignment using MUSCLE [48]. This alignment then provided the input for phylogenetic tree reconstruction using IQ-TREE [49]. The phylogenetic tree was then analysed with HyPhy [21], allowing reconstruction of the DIOSvax-H5_{inter} design.

Plasmids and cloning

Cloning was performed via standard PCR methods using the Q5 High-Fidelity PCR kit (New England Biolabs), Gibson assembly using the Gibson Assembly[®] Master Mix (New England Biolabs), and site-directed mutagenesis using the Q5[®] Site-Directed Mutagenesis Kit (New England Biolabs). The genes of interest were validated by Sanger sequencing (Source Bioscience).

pET28a-SpyCatcher003-mi3 (GenBank MT945417, Addgene 159995) was previously described [26]. pEVAC-H5Ast20-SpyTag003-His₆ (GenBank PV751237) was created by inserting influenza A/Astrakhan/3212/2020 H5 ectodomain (GISAID EPI1846961, residues 1-521), GSGSGSPGS linker, T4 bacteriophage fibritin foldon domain (GYIPEAPRDGQAYVRKD-GEWVLLSTFL) [33], (GGS)₃G linker, SpyTag003 and His₆ tag into the pEVAC plasmid [34], followed by deletion of residues 341-344 (KRRK) to remove the polysite. pEVAC-DIOSvax-H5_{inter}basic cleavage SpyTag003-His₆ was created by replacing H5Ast20 ectodomain with DIOSvax-H5_{inter} ectodomain (residues 1-522), followed by deletion of amino acids 341-345 (RRRKK) to remove the polybasic cleavage site.

Bacterial protein expression of SpyCatcher003mi3

Bacterial protein expression of the SpyCatcher003mi3 nanocage was previously described [32]. *E. coli* BL21(DE3) cells (Agilent) were transformed with pET28a-SpyCatcher003-mi3 and grown on LB-Agar plates containing 50 µg/mL kanamycin for 16 h at 37 °C. One colony from this plate was inoculated in 10 mL LB containing 50 µg/mL kanamycin and grown for 16 h at 37 °C with shaking at 200 rpm. This starter culture was added to 1 L LB containing 50 µg/mL kanamycin and incubated at 37 °C with shaking at 200 rpm until optical density (OD)₆₀₀ was 0.6. Cultures were induced with 0.5 mM isopropyl β - D-1-thiogalactopyranoside and grown for 16 h at 22 $^{\circ}$ C with shaking at 200 rpm, before centrifugation for 30 min at 4,000 g and 4 $^{\circ}$ C.

Purification of SpyCatcher003-mi3

Purification of the SpyCatcher003-mi3 nanocage was previously described [32]. Cell pellets were resuspended in 20 mL 20 mM Tris-HCl, 300 mM NaCl, pH 8.5, that was supplemented with 0.1 mg/mL lysozyme, 1 mg/mL cOmplete mini EDTA-free protease inhibitor (Roche) and 1 mM phenylmethanesulfonyl fluoride (PMSF). The lysate was incubated for 45 min at 4 °C with end-over-end mixing. An Ultrasonic Processor equipped with a microtip (Cole-Parmer) was used for sonication on ice (four times for 60 s, 50% duty-cycle). Centrifugation was performed for 45 min at 35,000 g and 4 °C to clear cell debris. 170 mg of ammonium sulfate was added for every mL of lysate. The mixture was incubated for 1 h at 4 °C, with mixing at 120 rpm to precipitate the particles. Centrifugation for 30 min at 30,000 g and 4 °C was performed. The pellet was resuspended in 10 mL mi3 buffer (25 mM Tris-HCl, 150 mM NaCl, pH 8.0) at 4 °C. The solution was filtered sequentially through 0.45 μ m and 0.22 μ m syringe filters (Starlab). The filtrate was dialysed for 16 h against 1,000-fold excess of mi3 buffer. The dialysed particles were centrifuged for 30 min at 17,000 g and 4 °C and subsequently filtered through a 0.22 µm syringe filter. The purified SpyCatcher003-mi3 was loaded onto a HiPrep Sephacryl S-400 HR 16-600 column (GE Healthcare), which had been equilibrated with mi3 buffer using an ÄKTA Pure 25 system (GE Healthcare). The proteins were separated with collections at each 1 mL elution fraction. The fractions containing the purified particles were pooled and concentrated using a Vivaspin 20 100 kDa molecular weight cut-off (MWCO) centrifugal concentrator (GE Healthcare). The final nanocage was stored at -80 °C.

Mammalian protein expression of HA-SpyTag003

Mammalian expression of all HA constructs was performed in Expi293FTM cells (Thermo Fisher). Cells were cultured in Expi293TM Expression Medium (Thermo Fisher) in suspension at 37 °C, 8% (v/v) CO_2 , \geq 80% relative humidity and rotating at 125 rpm. Transient transfections were performed using the ExpiFectamineTM 293 Transfection Kit (Thermo Fisher). Cells at 3×10^6 cells/mL were transfected with 1 µg of plasmid DNA and 2.7 µL of ExpiFectamineTM 293 per mL of culture. ExpiFectamineTM 293 Transfection Enhancers 1 and 2 were added 20 h posttransfection. Cell supernatants were harvested 5 days post-transfection by centrifuging for 10 min at 4000 g and 4 °C and sterile filtering through a 0.45 μ m filter followed by a 0.22 μ m filter (Merck). cOmplete mini EDTA-free protease inhibitor cocktail was added to the supernatants at 1 mg/mL (Roche).

Purification of HA-SpyTag003

HA-SpyTag003 proteins were purified by nickel-nitrilotriacetic acid (Ni-NTA) affinity chromatography. Ni-NTA agarose (Qiagen) was washed with 2×10 column volumes (CV) of Ni-NTA buffer (50 mM Tris-HCl, 300 mM NaCl, pH 7.8 at 4 °C). Mammalian cell supernatants were supplemented with 10× Ni-NTA buffer (500 mM Tris-HCl, 3 M NaCl, pH 7.8 at 4 °C) at 10% (v/v) and incubated with Ni-NTA affinity matrix for 1 h at 4 °C with rolling. The mixtures were added to an Econo-Pac Chromatography Column (Bio-Rad), passed through by gravity filtration, and washed with 2×10 CV of Ni-NTA wash buffer (10 mM imidazole in Ni-NTA buffer). Ni-NTA elution buffer (200 mM imidazole in Ni-NTA buffer) was added to the column, incubated for 5 min and passed through by gravity filtration to elute proteins. A total of 6×1 CV elutions was performed. Total eluates were dialysed through 3.5 kDa MWCO Spectra-Por® Float-A-Lyzer® G2 (Spectrum Labs) for 16 h against 1,000-fold excess phosphate buffered saline (PBS) (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.7 mM KH₂PO₄, pH 7.4). Dialysed eluates were concentrated in a 20 mL 10 kDa PierceTM Protein Concentrator (Thermo Fisher) by centrifuging for 5 min at 4,000 g and 4 °C.

SDS-PAGE

All fractions collected during transfection harvest and affinity chromatography (supernatant, flow-through, washes 1-2 and elutions 1-6) and purified proteins were analysed by SDS-PAGE under reducing conditions using the CriterionTM Cell electrophoresis system (Bio-Rad). 2× Laemmli Sample Buffer (Bio-Rad) was supplemented with 2-mercaptoethanol at 5% (v/ v) and mixed with samples at 50% (v/v). Diluted samples were heated for 5 min at 95 °C and resolved in a 12.5% Criterion[™] Precast Tris-HCl gel (Bio-Rad). Electrophoresis was performed in Tris/Glycine/SDS buffer (Bio-Rad) for 70 min at 180 V. Gels were stained for 1 h with InstantBlue® Coomassie Stain (Abcam) and destained for 16 h in MilliQ water. Imaging was performed with ChemiDoc XRS + imager and ImageLab software (Bio-Rad).

PNGase F digestion

De-glycosylation of HA was performed using the PNGase F kit (New England Biolabs). $2 \mu M$ of HA was heated with $1 \mu L$ of $10 \times$ Glycoprotein Denaturing Buffer (0.5% (w/v) SDS, 40 mM dithiothreitol) for

10 min at 100 °C. The denatured protein was chilled on ice for 1 min and centrifuged for 10 s at 2000 g. 2 μ L of 10× GlycoBuffer 2 (50 mM sodium phosphate, pH 7.5 at 25 °C), 2 μ L of 10% (v/v) NP-40, 6 μ L of MilliQ water and 1 μ L of PNGase F at 500,000 units/ mL were added and incubated for 1 h at 37°C, before resolving in a 12.5% (w/v) CriterionTM Precast Tris-HCl gel (Bio-Rad) with Coomassie staining.

Endotoxin depletion and quantification

SpyCatcher003-mi3 and HA-SpyTag003 proteins were depleted of endotoxins using Triton X-114 phase separation [50]. Triton X-114 was mixed with the proteins at 1% (v/v). The mixtures were incubated for 5 min on ice, then incubated for 5 min at 37 °C, and centrifuged for 1 min at 16000 g and 37 °C. The top phase was transferred to a fresh tube. A total of three repetitions were performed, followed by a final repetition without the addition of Triton X-114. All samples were assessed for final endotoxin concentrations using PierceTM Chromogenic Endotoxin Quant Kit (Thermo Fisher) and confirmed to be below the accepted <20 Endotoxin Units (EU) per mL for vaccine products [51].

Immunogen preparation

The concentration of SpyCatcher003-mi3 nanocages was determined by A₂₈₀ measurements taken using a NanoDropTM One Spectrophotometer (Thermo Fisher). The concentration of HA-SpyTag003 was measured using the PierceTM Bicinchoninic Acid Assay kit (Thermo Fisher). Doses were normalized by the number of antigens, to facilitate an equimolar amount of SpyCatcher003-mi3 nanocages with similar occupancy in each condition. 1.5 µM HA-SpyTag003 was incubated with 1 µM SpyCatcher003-mi3 for 24 h at 4 °C in PBS, pH 7.2. Uncoupled HA-SpyTag003 was incubated for 24 h at 4 °C in PBS, pH 7.2, without the addition of SpyCatcher003-mi3. Post-coupling, immunogens were analysed by SDS-PAGE/Coomassie, DLS and ELISA. Immunogens were further diluted in PBS, pH 7.2, to the final doses of 0.5 µg of total Spy-Tagged antigens, which relates to 8 pmol of uncoupled HA. Before immunization, immunogens were mixed 1:1 with AddaVax (Invivogen).

Dynamic light scattering (DLS)

Samples were centrifuged for 30 min at 16000 g and 4 °C, before 70 μ L was loaded onto a plastic cuvette. Samples were measured in triplicate at 20 °C using a Zetasizer Nano S (Malvern Panalytical) with 11 scans of 10 s each. Before collecting data, the cuvette was incubated in the instrument for 2 min to allow the sample temperature to stabilize. The size

distribution was determined by the Zetasizer Software v.7.13 (Malvern Panalytical), calculating the mean and s.d. from the multiple scans.

Transmission electron microscopy (TEM)

Samples were centrifuged for 30 min at 16000 g at 4 °C. Carbon 400 mesh copper grids (EM Resolutions) were processed in a Quorum Emitech K100X glow discharger at 25 mA for 2 min. Samples were applied to the grids for 30 s, washed $2 \times$ with distilled H₂O for 10 s and stained with 2% (w/v) uranyl acetate for 10 s. Samples were blotted with filter paper between steps and air-dried after staining. Grids were imaged using a FEI Tecnai G2 80–200-keV transmission electron microscope at 200 keV with a 20 µm objective aperture at the Cambridge Advanced Imaging Centre.

ELISA

NuncTM MaxiSorpTM 96-Well flat-bottom plates (Thermo Fisher) were coated with 50 µL per well of 1 µg/mL coupled or uncoupled HA-SpyTag003 in PBS, pH 7.2, and incubated overnight at 4 °C. Plates were blocked with 3% (w/v) skimmed milk in PBS supplemented with 0.1% (v/v) Tween 20 (PBST) for 2 h at RT. Sera and monoclonal antibodies were serially diluted in 1% (w/v) skimmed milk in PBST using an eight-point, three-fold series starting at 1:200 and $0.5 \,\mu g/mL$, respectively. After removing the blocking buffer, 50 µL per well of sera or broadly neutralizing monoclonal antibodies CR9114 [36], were added and incubated for 2 h at RT with shaking at 350 rpm. Plates were washed $3 \times$ with 200 µL per well of PBST. 50 µl of 1:3000 (v/v) Peroxidase AffiniPureTM Goat Anti-Mouse IgG (H + L) (Jackson ImmunoResearch, 115-035-003) or 1:5000 (v/v) Peroxidase AffiniPureTM Rabbit Anti-Human IgG (H + L) (Jackson ImmunoResearch, 309-035-003) per well was added and incubated for 1 h at RT with shaking at 350 rpm. Plates were washed $3 \times$ with 200 µL per well of PBST. 50 µL per well of 1-Step Ultra TMB chromogenic substrate (Merck) was added to the plates incubated for 3 min at RT, before the chemical reaction was stopped with 50 μ L 1 N H₂SO₄. OD₄₅₀ was measured using BioTek 800 TS Absorbance Reader and Gen6 software (Agilent). Using GraphPad Prism (GraphPad Software v10.3.1), nonlinear regression was performed to plot sigmoidal dose-response curves, from which area under the curve (AUC) values were determined and plotted as means ± 1 s.d. of duplicate measurements.

Mouse immunization and blood sampling

Animal experiments were performed according to the UK Animals (Scientific Procedures) Act 1986, under Project Licence (PP9157246) and approved by the University of Cambridge Animal Welfare and Ethical Review Body. Groups of six 8–10-week-old female BALB/c mice were obtained from Charles River Laboratories. Mice were immunized twice at a 28-day interval. A total volume of 100 μ L of 0.5 μ g of HA-SpyTag003 was administered via the intramuscular route over both hind legs. Blood was sampled from the saphenous vein at 14-day intervals and animals were terminally bled by cardiac puncture under non-recovery anaesthesia on day 56. Whole blood was allowed to clot at 25°C for 1–2 h and centrifuged for 5 min at 10000 g. Clarified sera were transferred to fresh tubes, heat-inactivated for 30 min at 56 °C, and stored at -30 °C.

Lentiviral pseudotype production

Influenza pseudotyped viruses A/Vietnam/1203/2004 (clade 1), A/Indonesia/5/2005 (clade 2.1.3.2), A/whooper swan/Mongolia/244/2005 (clade 2.2), A/Hubei/1/ 2010 (clade 2.3.2.1a), A/Sichuan/26221/2014 (clade 2.3.4.4a), A/Astrakhan/3212/2020 (clade 2.3.4.4b), A/ gyrfalcon/Washington/41088-6/2014 (clade 2.3.4.4c), A/Guangdong/18SF020/2018 (clade 2.3.4.4h), A/ goose/Guiyang/337/2006 (clade 4), A/chicken/Vietnam/NCVD-016/2008 (clade 7.1), A/chicken/Mexico/07/2007 (AM non-GS/GD) and A/mallard duck/ Netherlands/41/2015 (EU non-GS/GD) were produced in HEK293T/17 cells as previously described [34,42]. Cells were cultured in DMEM GlutaMAXTM (Thermo Fisher) supplemented with 10% (v/v) foetal bovine serum (FBS), 100 U/mL penicillin and 100 µg/mL streptomycin at 37 °C, 5% (v/v) CO₂, ≥80% relative humidity. Transient transfections with lentiviral packaging plasmids p8.91-gag-pol [52] and pCSFLW-firefly luciferase [53], protease-bearing plasmid pCMV-TMPRSS4 [54] and glycoprotein-bearing plasmid pEVAC-HA were performed using the FuGENE® HD Transfection Reagent (Promega). Cells at 4×10^5 cells/mL were seeded in 6-well plates and incubated for 24 h at 37 °C until ~70% confluence. Cells were transfected with 125 ng of p8.91, 187.5 ng of pCSFLW, 0-62.5 ng of pCMV-TMPRSS4 and 5-125 ng of pEVAC-HA plasmid DNA per mL of culture with 3 μ L of FuGENE HD per μ g of DNA. Exogenous neuraminidase from Clostridium perfringens (Merck) was added 24 h post-transfection at 0.5 U/mL. Cell supernatants were harvested 48 h post-transfection by sterile filtering through a 0.45 µm filter (Merck) and stored at -80°C. Pseudotypes were titrated on HEK293T/17 cells for 48 h at 37 °C using an eight-point, twofold series starting at 1:2 to measure relative luminescence units (RLU)/mL.

Pseudotype-based microneutralisation assay

Pseudotype-based microneutralisation assays (pMN) were performed as previously described [34,42]. Sera

and the broadly neutralizing monoclonal antibody FI6 [55] were serially diluted in NuncTM MicroWellTM 96-Well flat-bottom plates (Thermo Fisher) using a twelve-point, twofold series starting at 1:80 and 0.5 µg/mL, respectively. Influenza HA-bearing pseudotyped viruses at $1.5-5 \times 10^6$ RLU/mL were added and incubated for 1 h at 37 °C and 5% (v/v) CO₂. HEK293 T/17 cells at 1.5×10^4 cells/well were then added and incubated for 48 h at 37 °C and 5% (v/v) CO₂. After removing the cell supernatants, excess Bright-GloTM (Promega) was added and incubated for 5 min at RT. Luminescence was measured using GloMax[®] Explorer (Promega). Individual data points were normalized to 100% and 0% neutralization controls, corresponding to values derived from uninfected cells and cells infected with pseudotyped virus in the absence of serum, respectively. Nonlinear regression analysis was performed to plot sigmoidal doseresponse curves and determine half-maximal inhibitory dilution (ID₅₀) values in log₁₀ scale. Statistical significance of differences in log₁₀(ID₅₀) values between groups was determined using the ANOVA test, followed by the Tukey's multiple comparison post hoc test. Results were plotted using GraphPad Prism (GraphPad Software v.10.3.1).

Intracellular cytokine staining and flow cytometry

Single-cell suspension of murine splenocytes was prepared as previously described [56]. Peptide pools consisting of 15 amino acid peptides with 11 amino acid overlap spanning the H5N1 A/Indonesia/CDC835/ 2006 (clade 2.1.3.2; Swiss-Prot ID: A1BK62) and A/ Aves/Guanajuato/CPA-18539-23/2023 (clade 2.3.4.4b; Swiss-Prot ID: WPD27583.1) HA proteins were used for stimulation (JPT peptides). Cells were treated by 50 U/mL of Benzonase[®] nuclease (Sigma Aldrich) in cell culture medium (RPMI 1640 Medium GlutaMAXTM (Thermo Fisher) supplemented with 1 mM sodium pyruvate, 50 µM 2-mercaptoethanol, 10% (v/v) FBS, 100 U/mL penicillin and 100 µg/ml streptomycin) to prevent clumping. Cells were resuspended in NuncTM MicroWellTM 96-Well U-bottom plates (Thermo Fisher) at 5×10^5 cells/well in cell culture medium. $2 \mu g/mL$ peptides or $1 \times eBioscience^{TM}$ Cell Stimulation Cocktail (Thermo Fisher) were added and incubated for 2 h at 37°C, 5% (v/v) CO₂, \geq 80% relative humidity. 1× eBioscienceTM Protein Transport Inhibitor Cocktail (Thermo Fisher) was added and incubated for another 4 h. Plates were centrifuged for 3 min at 330 g to remove supernatant. Cells were washed with flow cytometry buffer (PBS supplemented with 2.5% (v/v) FBS and 2 mM EDTA). For surface marker staining, cells were incubated with 1:1,000 (v/v) eBioscienceTM Fixable Viability Dye eFluorTM 780 (Thermo Fisher), 1:100 (v/v) CD3e Monoclonal Antibody (145-2C11) PE-Cyanine5.5 (Thermo Fisher, 35-0031-82), 1:100 (v/v) CD8a Monoclonal Antibody (53-6.7) PE-Cyanine7 (Thermo Fisher, 25-0081-82) and 1:100 (v/v) CD4 Monoclonal Antibody (RM4-5) APC (Thermo Fisher, 17-0042-82) in flow cytometry buffer for 30 min at 4 ° C. Cells were washed twice with flow cytometry buffer, fixed in Fixation/Permeabilization solution (BD Biosciences) for 10 min at RT and washed twice with flow cytometry buffer. For intracellular cytokine staining, cells were incubated with 1:100 (v/v) eBioscienceTM IFN gamma Monoclonal Antibody (XMG1.2) Alexa FluorTM 488 (Thermo Fisher, 53-7311-82) and 1:100 (v/v) eBioscience[™] TNF alpha Monoclonal Antibody (MP6-XT22) PE-eFluor[™] 610 (Thermo Fisher, 61-7321-82) in 1× Perm/Wash buffer (BD Biosciences) for 30 min at RT. Cells were washed twice with BD Perm/WashTM buffer (BD Biosciences) and resuspended in flow cytometry buffer. Plates were analysed using the AttuneTM Nxt Flow Cytometer (Thermo Fisher). Data analysis was performed in Attune Cytometric Software 6.21 (Thermo Fisher). Values less than zero after subtraction of background were assigned a value of zero. Statistical significance of differences in % gated cells between groups was determined using the ANOVA test, followed by the Tukey's multiple comparison post hoc test. Results were plotted using GraphPad Prism (GraphPad Software v.10.3.1).

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Contributions

C.Q.H. conceived the main ideas of the study and acquired funding. C.Q.H. and R.A.H. designed and coordinated the project. L.S.T., M.R.H. and J.L.H. acquired funding and supervised the project. S.D.W.F. developed the computational pipeline and designed the antigen. J.L.H., S.V. and G.W.C. tested, validated and selected the candidate antigen breadth of immunogenicity by gene delivery. C.Q.H. and R.A.H. performed DNA cloning, protein production and purification, prepared immunogens, and completed SDS- PAGE, DLS, TEM and ELISA. C.Q.H., G.W.C., L.O. and P.T. performed mouse immunization and blood sampling. C.Q.H. produced lentiviral pseudotypes with guidance from G.W.C. and N.T.. C.Q.H. and G.W.C. performed pseudotype-based microneutralisation. C.Q.H. and A.C.Y.C. performed splenocyte purification. C.Q.H. and E.T.A. performed intracellular cytokine staining and flow cytometry. S.V. and P.P. contributed to data analysis and visualization. C.Q.H. wrote the manuscript. All authors contributed substantially to discussion of the content, reviewed and/or edited the manuscript before submission.

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Disclosure statement

J.L.H., G.W.C., S.V. and S.D.W.F. developed, tested and validated the H5 antigen DIOSvax-H5_{inter} by gene delivery and are inventors of patent applications on computational vaccine development methods (US20220040284A1) and influenza vaccines (US20230149530A1). J.L.H. and S.D.W.F. are co-founders and shareholders of DIOSynVax Ltd. M.R.H. is a co-founder and shareholder of SpyBiotech. M.R.H. is an inventor on a patent on spontaneous amide bond formation (EP2534484) and a patent on SpyTag003: SpyCatcher003 (UK Intellectual Property Office 1706430.4). S.D.W.F. is an employee of Microsoft.

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Data availability statement

Further information and request for resources and reagents should be directed to and will be fulfilled by the lead contacts, J.L.H. (jlh66@cam.ac.uk), M.R.H. (mh2186@cam.ac.uk) and L.S.T. (lst21@cam.ac.uk).

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