Nucleoside-modified mRNA encoding HSV-2 glycoproteins C, D, and E prevents clinical and subclinical genital herpes

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The goals of a genital herpes vaccine are to prevent painful genital lesions and reduce or eliminate subclinical infection that risks transmission to partners and newborns. We evaluated a trivalent glycoprotein vaccine containing herpes simplex virus type 2 (HSV-2) entry molecule glycoprotein D (gD2) and two immune evasion molecules: glycoprotein C (gC2), which binds complement C3b, and glycoprotein E (gE2), which blocks immunoglobulin G (IgG) Fc activities. The trivalent vaccine was administered as baculovirus proteins with CpG and alum, or the identical amino acids were expressed using nucleoside-modified mRNA in lipid nanoparticles (LNPs). Both formulations completely prevented genital lesions in mice and guinea pigs. Differences emerged when evaluating subclinical infection. The trivalent protein vaccine prevented dorsal root ganglia infection, and day 2 and 4 vaginal cultures were negative in 23 of 30 (73%) mice compared with 63 of 64 (98%) in the mRNA group (P = 0.0012). In guinea pigs, 5 of 10 (50%) animals in the trivalent subunit protein group had vaginal shedding of HSV-2 DNA on 19 of 210 (9%) days compared with 2 of 10 (20%) animals in the mRNA group that shed HSV-2 DNA on 5 of 210 (2%) days (P = 0.0052). The trivalent mRNA vaccine was superior to trivalent proteins in stimulating ELISA IgG antibodies, neutralizing antibodies, antibodies that bind to crucial gD2 epitopes involved in entry and cell-to-cell spread, CD4+ T cell responses, and T follicular helper and germinal center B cell responses. The trivalent nucleoside-modified mRNA-LNP vaccine is a promising candidate for human trials.

INTRODUCTION
Herpes simplex virus type 2 (HSV-2) is a common sexually transmitted infection with 14% prevalence among 15- to 49-year-old persons in the United States and 11% globally (1). Individuals remain infected for life, with periodic clinical recurrences that may be painful or subclinical recurrences that risk transmission of infection to intimate partners and newborns (2). The incidence of neonatal herpes is about 14,000 cases annually worldwide (3) and results in substantial morbidity and mortality because of encephalitis, pneumonia, and hepatitis, even with antiviral therapy (4, 5). Genital herpes increases the risk of acquisition and transmission of HIV by three- to fourfold, further highlighting the need for an effective prophylactic vaccine (6).

Previous HSV-2 prophylactic human vaccine efforts included immunizing with two essential virus entry molecules, HSV-2 glycoproteins B (gB2) and D (gD2) (7), or gD2 alone (8, 9). In the gB2/gD2 trial, vaccine recipients had delayed onset of infection but overall were not protected (7). In the first of two reports using gD2 alone, a subgroup analysis indicated that HSV-1/HSV-2 doubly seronegative women, but not men, were protected against HSV-2 genital lesions (8). A repeat study performed in doubly seronegative women failed to confirm this result but noted 58% protection against genital lesions caused by HSV-1. Sixty percent of the cases of genital herpes in the control group were caused by HSV-1 (9). Enzyme-linked immunosorbent assay (ELISA) antibodies to gD2 correlated with protection against genital HSV-1 infection (10). Maternal and neonatal anti-bodies that neutralize HSV or mediate antibody-dependent cellular cytotoxicity (ADCC) also correlate with protection against severe neonatal herpes (11–13). These studies suggest that antibodies are important for prevention of herpes infection.

Our vaccine strategy is designed to produce potent antibody responses and includes an entry molecule, gD, and two additional HSV glycoproteins, gC and gE. HSV-1 and HSV-2 gC and gE are immune evasion molecules that block the effectiveness of antibody responses (14–16). HSV gC binds complement component C3b to inhibit complement activation, whereas gE binds the immunoglobulin G (IgG) Fc domain of antibodies targeting HSV antigens to block IgG Fc activities, including complement activation and ADCC (17–20). Antibodies produced to the three glycoproteins perform multiple antiviral activities, including neutralizing virus (gC2 and gD2), blocking cell-to-cell spread (gD2 and gE2), and preventing immune evasion from antibody and complement (gC2 and gE2) (16, 21–24).

No genital herpes vaccine has prevented both clinical and subclinical infection. Our previous vaccine studies in mice and guinea pigs used baculovirus-produced gC2, gD2, and gE2 subunit proteins with CpG and alum as adjuvants. The vaccine provided strong protection against clinical lesions in mice (100%) and guinea pigs (98%), but it did not prevent vaginal virus replication on days 2 and 4 after infection in mice and 17 of 36 (47%) guinea pigs developed subclinical infection detected by vaginal shedding of HSV-2 DNA between days 28 and 49 after infection (16, 22). Here, we evaluated whether an immunization strategy that uses nucleoside-modified mRNA to encode the same three HSV-2 glycoproteins improves vaccine efficacy.

Nucleoside-modified mRNA vaccines have recently emerged as highly promising for infectious disease vaccine development (25). Modifications in mRNA increase translation and reduce inflammatory side effects. Modifications include altering the 5′ cap, the 5′ and 3′ untranslated regions, and poly A tails; replacing uridine with

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Characterization of the truncated gc2, gd2, and ge2 proteins produced in cells transfected with nucleoside-modified mRNA-LNP

Nucleoside-modified mRNA was prepared to encode gc2 amino acids 27 to 426, gd2 amino acids 26 to 331, and ge2 amino acids 24 to 405, each truncated before their transmembrane domains. Expression of the glycoproteins was assessed in human embryonic kidney (HEK) 293 cells that were transfected with the mRNAs or with PolyC RNA as a control. The proteins in the mRNA-transfected cells reacted with gc2, gd2, or ge2 antibodies by Western blot and were of the expected size when accounting for different glycosylation patterns in mammalian cells (fig. S1, A to C). Supernatant fluids from the transfected cells contained the glycoprotein antigens when assayed by ELISA (fig. S1, D to F). We conclude that cells transfected with each nucleoside-modified mRNA synthesized and secreted the truncated glycoprotein.

Vaccine efficacy in mice

We infected mice intravaginally with either 5 × 10^3 or 5 × 10^4 plaque-forming units (PFU) of HSV-2 to assess protection over a 10-fold titer range.

Protection against 5 × 10^3 PFU (2750 LD₅₀) of intravaginal HSV-2 after intradermal immunization with nucleoside-modified mRNA-LNP

BALB/c mice were immunized intradermally (ID) twice at a 28-day interval with nucleoside-modified mRNA-LNP consisting of 10 g of gd2 mRNA-LNP; 10 g each of gc2, gd2, and ge2 mRNA combined in an LNP; or 10 g of PolyC RNA-LNP as a control. For comparison, mice were immunized intramuscularly (IM) three times at 2-week intervals with 5 g each of gc2, gd2, and ge2 baculovirus-expressed proteins combined with CpG/alum. These concentrations and dosing interval were chosen on the basis of our previous studies in mice using baculovirus subunit protein antigens and mRNA-LNP studies performed with other pathogen vaccines (21, 22, 29–31). Twenty-eight days after the last immunization, mice were treated with medroxyprogesterone to synchronize and thin the vaginal epithelium and, 5 days later, were infected intravaginally with 5 × 10^3 PFU of HSV-2 strain MS [275 median lethal dose (LD₅₀)]. All mice in the PolyC control group were euthanized by day 10, whereas no other animals died (Fig. 1A).

No animal in the trivalent mRNA, trivalent protein, or gd2 mRNA group developed ruffled fur, hunched posture, abnormal gait, or lethargy as disease indicators. Weights of animals in the trivalent mRNA, trivalent protein, or gd2 mRNA group did not differ significantly from one another, yet each group differed significantly from the PolyC RNA controls (Fig. 1B). No genital disease developed in the trivalent mRNA or trivalent protein group. Mild genital disease occurred in one animal in the gd2 mRNA group. Overall, these three groups did not differ significantly from one another, whereas extensive genital disease developed in the PolyC RNA mice (Fig. 1C).

Day 2 and 4 vaginal cultures measure acute viral infection and represent a stringent test of vaccine efficacy. Notably, 39 of 39 animals in the mRNA trivalent group had negative vaginal cultures on day 2, which was significantly different from 7 of 10 in the trivalent protein group and 16 of 20 in the gd2 mRNA group (Fig. 1D). Significant differences between the trivalent mRNA group and the trivalent protein group or the gd2 mRNA group were also detected on day 4 after infection (Fig. 1D). We assessed day 2 and 4 vaginal HSV-2 DNA copy number by quantitative polymerase chain reaction (qPCR) to further evaluate indicators of virus infection. Approximately 5 log₁₀ HSV-2 DNA was detected in the trivalent mRNA group on day 2 and approximately 4 log₁₀ on day 4 despite recovering no replication-competent virus on either day, which suggests that the virus was coated with neutralizing antibodies and/or degraded (Fig. 1E). By day 4, the HSV-2 DNA copy number was significantly lower in the trivalent mRNA mice than in the trivalent protein or gd2 mRNA alone mice (Fig. 1E).

Peak HSV-2 DNA copy number in dorsal root ganglia (DRG) develops on day 4 or 5 after challenge in naïve mice (22). All 10 mice in the PolyC RNA group had HSV-2 DNA detected in DRG on day 4 compared with 3 of 10 in the gd2 mRNA group, 1 of 19 in the trivalent mRNA group, and 5 of 10 in the trivalent subunit protein group. Differences between trivalent mRNA and trivalent subunit protein were significant, whereas trivalent mRNA and gd2 mRNA differences were not significant (Fig. 1F).

Protection against 5 × 10⁴ PFU (2750 LD₅₀) of intravaginal HSV-2 in mice

As a more stringent test of vaccine efficacy, five mice were immunized ID twice with 10 g each of trivalent mRNA-LNP and infected with a 10-fold higher HSV-2 dose, 5 × 10⁴ PFU of strain MS (2750 LD₅₀). We did not include the trivalent protein group because breakthrough infections occurred at the lower challenge dose. All five mice survived, had no genital disease, had negative vaginal cultures on days 2 and 4, and had no HSV-2 DNA detected in DRG at the end of the experiment on day 28 (Fig. 1G). Therefore, no breakthrough infections occurred even at a 10-fold higher challenge dose.

Vaccine efficacy in mice: Protection against intravaginal HSV-2 after IM immunization with nucleoside-modified mRNA-LNP

Our trivalent subunit protein vaccine studies in mice use IM immunization (22). We evaluated the trivalent mRNA vaccine given IM to compare the two vaccine formulations using the same immunization
route. Mice were immunized as in Fig. 1, except that one additional group received trivalent mRNA-LNP using 2 μg of each immunogen to assess protection at lower mRNA concentrations. Animals were challenged with 5 × 10³ PFU of HSV-2 strain MS (275 LD₅₀). All PolyC control animals were euthanized by day 11, whereas all other animals survived (Fig. 2A). No animal in the 2 or 10 μg trivalent
mRNA or trivalent subunit protein group developed ruffled fur, hunched posture, abnormal gait, or lethargy, and no significant differences in weight loss were detected comparing these three groups, whereas the PolyC controls had extensive weight loss (Fig. 2B). The PolyC RNA group developed severe genital disease, whereas the other groups were totally protected (Fig. 2C). Day 2 vaginal cultures were positive in all 15 mice in the PolyC RNA group, 1 of 10 in the trivalent subunit protein group, 4 of 5 in the 2 μg trivalent mRNA group, and 0 of 20 in the 10 μg trivalent mRNA group (Fig. 2D). All 20 mice in the 10 μg trivalent mRNA group also had negative vaginal cultures on day 4, as noted in 1D immunized animals (Fig. 2D). DRG were harvested at the time of euthanasia in nine mice from the PolyC RNA group or at the end of the experiment on day 28 in the other groups. All 9 animals in the PolyC RNA group had HSV-2 DNA detected in DRG, whereas 0 of 10 were positive in the trivalent subunit protein group, 1 of 5 in the 2 μg trivalent mRNA group, and 0 of 20 in the 10 μg trivalent mRNA group (Fig. 2E). We conclude that dose-response protection was apparent in the trivalent mRNA mice immunized IM based on significantly better protection at 10 μg than 2 μg against vaginal infection on day 2 and slightly better protection of DRG. Although 10 μg trivalent mRNA outperformed trivalent subunit protein when evaluating day 2 and 4 vaginal cultures, differences were not statistically significant.

**Combining the ID and IM results in mice**

We combined the IM and ID results for the 10 μg trivalent mRNA–immunized mice because both routes produced virtually identical results. The trivalent subunit protein and trivalent mRNA mice were both totally protected against clinical disease as measured by death, genital lesions, and signs of generalized illness (Table 1). Differences between the two vaccine formulations emerged when evaluating subclinical infection measured by day 2 and 4 vaginal cultures and HSV-2 DNA in DRG (Table 1). On the basis of clinical and subclinical infections, ID or IM immunization with 10 μg of trivalent mRNA achieved sterilizing immunity in 63 of 64 (98%) mice compared with 23 of 30 (77%) in the trivalent subunit protein group and 15 of 20 (75%) in the gD2 mRNA group, whereas 0 of 25 (0%) in the PolyC RNA group had sterilizing immunity (Table 1). The breakthrough infections in the trivalent subunit protein group included 5 of 10 animals that had HSV-2 DNA in DRG, which represents more DRG infection than we reported in a previous study (22). In the current study, we used a more stringent criterion for a...
negative result based on normalizing the HSV-2 DNA copy number to 10^5 adipsin genes rather than 10^4 previously.

Transmission studies in mice
Transmission studies were performed using day 2 vaginal secretions collected from 10 mice immunized ID with 10 μg of trivalent mRNA-LNP or PolyC RNA-LNP as a control and infected with 5 × 10^5 PFU (275 LD_{50}). The vaginal secretions (20 μl) from each mouse were inoculated intravaginally into a naïve mouse. An additional five naïve mice were inoculated with 5 × 10^5 PFU of HSV-2 as a positive control, and as expected, all five mice became ill and were euthanized. Three of 10 mice inoculated with vaginal secretions from the PolyC RNA group developed disease and were euthanized compared with 0 of 10 from the trivalent mRNA group (fig. S2A). All naïve animals in the HSV-2–inoculated group and 6 of 10 in the PolyC RNA group developed genital disease compared with 0 of 10 in the trivalent mRNA group. Differences were also noted comparing day 2 and 4 vaginal titers of mice inoculated with vaginal secretions from trivalent mRNA or PolyC RNA mice with 6 of 10 mice in the PolyC RNA group positive on both days compared with 0 of 10 in the trivalent mRNA group (fig. S2B). We did not evaluate transmission using vaginal fluids from subunit vaccine–immunized animals; however, 4 of 20 mice in this group had positive day 2 titers compared with 0 of 64 in the modified mRNA group. Therefore, we postulate that more mice in the baculovirus protein group would transmit virus. HSV-2 DNA was detected in the day 2 vaginal fluids of most mice in the trivalent mRNA group, yet no replication-competent virus was detected and no transmission of infection occurred to naïve mice. This result is consistent with the concept that, on day 2, the virus was either coated with neutralizing antibody or degraded. Overall, no naïve mouse inoculated with 20 μl of vaginal secretions from the 10 μg trivalent mRNA group died, developed genital disease, or had positive cultures on day 2 or 4.

Vaccine efficacy in guinea pigs
The guinea pig genital infection model is generally viewed as a more stringent test of vaccine efficacy and more biologically relevant in that guinea pigs develop recurrent genital lesions and recurrent vaginal shedding of HSV-2 DNA, whereas mice do not. Genital HSV-2 shedding indicates reactivation from latency and is a marker of subclinical infection when it occurs in the absence of genital lesions. Nucleoside-modified mRNA-LNP vaccines have not previously been evaluated in guinea pigs; therefore, we decided to use three immunizations, rather than two as in mice. We chose a dose of 20 μg per nucleoside-modified mRNA rather than the 10-μg dose used in mice based on the larger body size of guinea pigs. Female Hartley strain guinea pigs (10 animals per group) were immunized ID three times at 1-month intervals with 20 μg each of gC2, gD2, and gE2 nucleoside-modified mRNA-LNP; IM with 10 μg each of gC2, gD2, and gE2 baculovirus subunit proteins with CpG/alum; or IM with 10 μg of PolyC RNA-LNP as a control or left unimmunized. One month after the final immunization, animals were infected intravaginally with 5 × 10^5 PFU (50 LD_{50}) of HSV-2 MS, which is the dose we used in previous studies with the trivalent baculovirus protein vaccine (16). The group that was not immunized were not infected. Animals were observed for survival, weight loss, and genital lesions. Eight of 10 animals in the PolyC RNA-LNP group became ill and were euthanized by day 20 after infection, whereas no animal in the trivalent mRNA, trivalent protein, or uninfected group died (fig. 3A) or developed weight loss (fig. 3B), hind limb weakness, or urinary retention. Animals in the PolyC RNA-LNP group had extensive genital lesions, whereas no genital lesions appeared in the other groups over 60 days (fig. 3C).

We performed vaginal cultures on days 2 and 4 after infection. Nine of 10 guinea pigs in the PolyC RNA-LNP group had positive titers on day 2. The negative day 2 titer in one animal indicates that the animal did not get infected, which is consistent with the observation that one animal in the PolyC RNA group did not lose weight or develop genital lesions. In the trivalent mRNA and trivalent subunit

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Table 1. Combined ID and IM efficacy studies.

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<th>Outcome</th>
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| Clinical disease:
  Death, genital lesions, or
  generalized illness | 25/25      | 1/10      | 0/44           | 0/20             |
| Subclinical infection          | 25/25      | 4/20      | 0/64           | 4/20             |
| Day 2 vaginal titers           | 25/25      | 3/20      | 0/64           | 4/20             |
| Day 4 vaginal titers           | 25/25      | 3/20      | 0/64           | 4/20             |
| HSV-2 DNA in DRG               | 19/19      | 3/10      | 1/39           | 5/20             |
| Sterile immunity:
  No clinical or subclinical
  infection | 0/25       | 15/20     | 63/64          | 23/30            |

*The primary end point is the comparison between trivalent mRNA-LNP and trivalent subunit protein. The secondary end point is the comparison between trivalent mRNA-LNP and gD2 mRNA-LNP. P values were calculated by two-tailed Fisher’s exact test.
Fig. 3. ID immunization with nucleoside-modified trivalent mRNA-LNP protects against genital HSV-2 in guinea pigs. (A to F) Guinea pigs were immunized three times monthly IM with 10 μg of PolyC RNA-LNP, ID with 20 μg of each trivalent nucleoside-modified mRNA-LNP, and IM with 10 μg of each trivalent subunit protein–CpG/alum or left unimmunized (n = 10 per group). One month after the final immunization, all immunized animals were infected intravaginally with 5 × 10^5 PFU of HSV-2 (50 LD₅₀), whereas unimmunized animals remained uninfected. (A) Survival curves: P < 0.0001, comparing PolyC RNA group with each other. (B) The mean weight loss each day after infection of animals in the trivalent mRNA, trivalent subunit protein, or uninfected group was compared with PolyC RNA controls; P < 0.0001. (C) Cumulative mean genital disease score per guinea pig is shown for each day after infection. No animal developed genital lesions in the trivalent mRNA, trivalent subunit protein, or uninfected group. P < 0.0001, comparing trivalent mRNA, trivalent subunit protein, or uninfected with PolyC RNA. The P values in (B) and (C) were calculated by two-way ANOVA followed by Tukey for significance. (D and E) P values shown in the figures are for the primary end point comparison between trivalent mRNA and trivalent protein. Other P values are noted below. (D) Day 2 and 4 virus titers: P = 0.0013 (day 2) and P = 0.0010 (day 4), comparing PolyC RNA with trivalent mRNA. Dotted line indicates the limit of assay detection at 3.3 PFU/ml. (E) Day 2 and 4 vaginal DNA copy number by qPCR. P values in (D) and (E) were calculated by the two-tailed Mann-Whitney, except that the day 4 comparison between trivalent mRNA and trivalent subunit proteins was performed by the two-tailed Fisher’s exact test. (F) Vaginal shedding of HSV-2 DNA over 21 days from days 28 to 50 after infection. One animal in the trivalent subunit protein group had replication-competent virus recovered on the day HSV-2 DNA shedding was detected (marked in green). P = 0.0052 (calculated by the two-tailed Fisher’s exact test), comparing days of shedding in the trivalent mRNA and trivalent subunit protein groups. (G) HSV-2 DNA copy number per 10^6 copies of GAPDH in spinal cord and DRG at the end of the experiment; n = 2 in the PolyC RNA group and n = 10 in the 3 other groups. (H) Table summarizing key results of guinea pig studies.
protein groups, 5 of 10 animals had positive vaginal titers on day 2. The positive vaginal cultures represent the mean of two determinations in a total of 300 μl of vaginal fluid. Two animals in the trivalent mRNA group and one animal in the trivalent subunit protein group had one virus plaque detected (Fig. 3D). Vaginal titers remained positive on day 4 in 9 of 10 animals in the PolyC RNA group and in 2 of 10 animals in the trivalent subunit protein group. In contrast, all 10 animals in the trivalent mRNA group had negative vaginal cultures on day 4 (Fig. 3D). Therefore, no animal in the trivalent mRNA group had a positive vaginal virus titer beyond day 2. We evaluated day 2 and 4 HSV-2 DNA copy number by qPCR. No significant differences were detected comparing trivalent mRNA with trivalent subunit protein animals, although DNA copy numbers were generally lower in the trivalent mRNA group (Fig. 3E). Some guinea pigs had high copy numbers of HSV-2 DNA despite negative virus cultures, suggesting that the virus was neutralized and/or degraded, as in the murine studies.
The inoculation titer used for the guinea pig experiments was $5 \times 10^7$ PFU, which is 10- to 100-fold higher than the inoculum used in mice. The low positive day 2 titers in the trivalent mRNA and trivalent protein groups may represent residual input virus and/or local replication of virus in genital tissues. An important goal of the vaccine is to prevent the virus from reaching sites where the virus establishes latency in ganglia that serve as a source for reactivation infection. We performed daily vaginal swabs for 21 days starting on day 28 after infection to detect vaginal shedding of HSV-2 DNA as a marker of reactivation infection. Two animals in the PolyC RNA group survived until day 21, but only one was infected on the basis of day 2 titers. We did not detect HSV-2 DNA in vaginal swabs from either the infected or uninfected animal. No animal in the uninfected group shed HSV-2 DNA. In the trivalent subunit protein group, 5 of 10 animals shed HSV-2 DNA on 19 of 210 (9.0%) days. In contrast, 2 of 10 guinea pigs shed HSV-2 DNA in the trivalent nucleoside-modified mRNA-LNP group on a total of 5 of 210 (2.4%) days ($P = 0.0052$; Fig. 3F). Virus cultures were performed to detect replication-competent virus on days that animals had vaginal shedding of HSV-2 DNA. We performed virus cultures only on samples that were positive for HSV-2 DNA because we consistently failed to recover replication-competent virus on days that animals were not shedding HSV-2 DNA (32). In the trivalent mRNA group, 0 of 5 HSV-2 DNA–positive samples yielded replication-competent virus compared with 1 of 19 in the trivalent protein group. Spinal cord and DRG were evaluated for HSV-2 DNA at the end of the experiment. One of two animals in the PolyC RNA group was positive for HSV-2 DNA in DRG and spinal cord (the animal with disease), whereas 1 of 10 in the trivalent mRNA group was positive for HSV-2 DNA in spinal cord but not DRG. The positive animal was the guinea pig that had vaginal shedding of HSV-2 DNA for 4 days from days 47 to 50. In the trivalent subunit protein group, 0 of 10 animals was positive for HSV-2 DNA in spinal cord or DRG, suggesting that detecting HSV-2 DNA in vaginal secretions is a more sensitive assay for subclinical infection than assays performed at the end of the experiment to detect HSV-2 DNA in DRG or spinal cord (Fig. 3G). Key guinea pig results are summarized in Fig. 3H.

**Immunology assays in mice**

In both mice and guinea pigs, vaccine efficacy was better with the trivalent nucleoside-modified mRNA-LNP formulation than with the trivalent subunit protein vaccine. We performed immunology analyses in mice to evaluate the mechanisms of the enhanced protection.

**Serum IgG antibody responses in mice**

ELISA end point IgG titers were evaluated before infection on sera collected 28 days after the first and second (final) mRNA ID immunizations or 28 days after the third (final) trivalent subunit protein IM immunization. ELISA titers increased comparing the first and second immunizations for each nucleoside-modified mRNA (Fig. 4, A to C). The nucleoside-modified mRNA produced significantly higher ELISA end point titers to each immunogen after the second immunization compared with the baculovirus-expressed subunit proteins after the third immunization (Fig. 4, A to C). After the final immunization, the ratio of IgG2a to IgG1 in mice immunized with trivalent mRNA was 0.89, whereas the ratio in animals immunized with trivalent subunit proteins was 1.04, suggesting a balanced T helper 1 (TH1)/TH2 response for both (Fig. S3) (33).

**Vaginal wash IgG antibody responses in mice**

Vaginal mucosa fluids were obtained before infection at 4 weeks after the final immunization. IgG vaginal mucosa antibody responses to gC2 and gE2 were significantly higher in mice immunized with trivalent mRNA-LNP than trivalent subunit proteins (Fig. 4, D and F). Titers were also higher to gD2 in the trivalent mRNA mice, although differences were not significant (Fig. 4E).

**Serum neutralizing antibodies in mice**

HSV-2 neutralizing antibody titers in the presence of complement were evaluated after the second (final) mRNA immunization or third (final) subunit protein immunization. The neutralizing antibody titers in the trivalent mRNA group were higher than those in the trivalent subunit protein group (Fig. 4G). Neutralizing titers were also higher in the trivalent mRNA group than in the gD2 mRNA group, indicating that gC2 and/or gE2 antibodies enhance neutralization (Fig. 4G) (16, 21, 22). HSV-1 is a frequent cause of genital herpes (9); therefore, we evaluated the same sera for neutralizing antibody titers against HSV-1. Each of the vaccine formulations produced high titers of neutralizing antibodies against HSV-1 (Fig. 4H). The trivalent mRNA vaccine produced higher HSV-1 neutralizing antibody titers than trivalent subunit proteins or gD2 mRNA, although the differences did not reach statistical significance (Fig. 4H).

**ADCC responses in mice**

The control PolyC RNA sera failed to stimulate ADCC activity mediated by murine FcyRIII, whereas sera from the trivalent mRNA, the trivalent subunit protein, and the gD2 mRNA mice produced significantly greater ADCC responses (Fig. 4I). ADCC responses did not differ comparing trivalent mRNA and trivalent subunit protein, suggesting that ADCC does not explain the superior performance of the trivalent mRNA vaccine (Fig. 4I).

**Antibodies that block immune evasion domains on gC2 and gE2 in mice**

We reported that immunization with trivalent subunit proteins produced higher titers of antibodies that blocked gC2 binding to C3b than produced by HSV-2 infection (34). Here, we assessed whether trivalent mRNA immunization produces higher titers of blocking antibodies than trivalent subunit proteins. We compared multiple concentrations of IgG purified from sera obtained 4 weeks after the final immunization with trivalent mRNA-LNP or trivalent subunit proteins. The IgG was incubated with 40 ng of gC2 to determine whether the antibodies block gC2 binding to C3b on an ELISA plate (16, 21). IgG from gD2 mRNA–immunized mice was included as a negative control. At IgG of 10 or 20 μg/ml, the trivalent mRNA-LNP significantly outperformed trivalent subunit proteins in blocking gC2 binding to C3b (Fig. 5A). At IgG of 50 μg/ml, both trivalent mRNA and trivalent subunit protein totally blocked gC2 binding to C3b (Fig. 5A). The results indicate that both trivalent mRNA and trivalent subunit proteins produce antibodies that block gC2 binding to C3b but that the trivalent mRNA produces higher titers of blocking antibodies.

HSV-2 gE2 functions as an IgG Fc receptor (22, 35). We evaluated whether gE2 antibody produced by immunization with trivalent mRNA is more effective than trivalent subunit proteins at blocking gE2 protein binding to nonimmune IgG Fc (16, 22). As a control, we assessed whether gE2 antibodies are more effective than gD2 antibodies at blocking gE2 binding to nonimmune IgG. This control...
takes into consideration that gE2 antibodies can bind to gE2 protein by their F(ab′)2 domain and/or Fc domain, whereas antibodies to gD2 protein can only bind by their Fc domain. We used gE2 protein (4 or 2 μg/ml) that was incubated with IgG (62.5 μg/ml) from immunized mice. Comparing trivalent mRNA with trivalent subunit proteins at a gE2 protein concentration of 4 μg/ml, IgG from the trivalent mRNA mice blocked gE2 binding better than trivalent subunit protein, suggesting that the trivalent mRNA produced higher titers of gE2 IgG Fc-blocking antibodies (Fig. 5B). These differences were not apparent at a lower gE2 concentration of 2 μg/ml, although both vaccine formulations blocked gE2 protein binding to IgG Fc significantly better than gD2 antibody (Fig. 5B). These results

Fig. 5. Antibodies that block gC2 and gE2 immune evasion domains and that bind gD2 epitopes that mediate crucial functions. P values are shown in the figures for the primary end point comparison between trivalent mRNA and trivalent protein. Other P values are noted below. (A) Antibodies produced by trivalent mRNA and trivalent subunit protein block C3b binding to gC2. Error bars represent geometric means and 95% confidence intervals. P = 0.0220 (at 10 μg/ml) and P < 0.0001 (at 20 and 50 μg/ml), comparing trivalent mRNA with gD2 mRNA. P < 0.0001 (at 50 μg/ml) and P = not significant (at other IgG concentrations), comparing trivalent subunit protein with gD2 mRNA. (B) Blocking gE2 binding to IgG Fc, P = 0.0079, comparing trivalent mRNA with gD2 mRNA at 4 μg/ml. P = 0.1508 (at 4 μg/ml) and P = 0.0079 (at 2 μg/ml), comparing trivalent subunit protein with gD2 mRNA. (A and B) Results are geometric means with 95% CI. IgG was purified from pooled sera of 10 animals per group. Each entry represents results from a single assay. P values were determined by the two-tailed Mann-Whitney test. (C to H) Sera were evaluated for blocking gD2 binding to prototype mAbs that recognize crucial gD2 epitopes; n = 10 for PolyC RNA, n = 20 for trivalent mRNA, and n = 9 for trivalent subunit protein. Results in (C) to (G) represent means with 95% CI. P values comparing trivalent mRNA with trivalent subunit protein were calculated using the two-tailed Mann-Whitney test.
Antibodies to gD2 epitopes involved in virus entry and cell-to-cell spread in mice

We recently evaluated epitope-specific antibody responses to three linear and four conformational gD2 epitopes that are crucial for gD2 function (23). The gD2 epitopes are involved in virus binding to herpesvirus entry mediator (HVEM) and/or nectin-1 receptors, gD2 interaction with HSV-2 downstream entry molecules gH2 and gL2, or cell-to-cell spread (36–38). We reported that guinea pigs immunized with gD2-CpG/alum produced a range of responses to crucial epitopes. Some animals produced antibodies to few epitopes, whereas others produced antibodies to most or all epitopes. The more crucial epitopes blocked, the better the protection was against intravaginal HSV-2 challenge (23).

Here, we evaluated antibody responses in mice to six gD2 epitopes that are involved in receptor binding, interaction with gH2/gL2, or cell-to-cell spread. The assay places monoclonal antibodies (mAbs) that recognize each of the gD2 crucial epitopes on a biosensor chip. Individual sera obtained after the second immunization with trivalent mRNA or third immunization with trivalent subunit proteins were incubated with gD2 protein and floated over the mAb-coated chip. Sera from mice immunized with PolyC RNA served as a control. Blocking gD2 protein binding to mAb on the biosensor chip indicates that the antibody is present in the mouse serum to the epitope recognized by the mAb. The percent blocking of gD2 binding to prototype mAbs that recognize the six crucial epitopes is shown in Fig. 5 (C to H). The trivalent mRNA-immunized mice produced antibodies that blocked gD2 binding significantly better than trivalent subunit proteins to three of six crucial epitopes. Although significant, the differences were fairly small for two of these three epitopes, 77S and MC5, but more impressive for the MC14 epitope (blocks cell-to-cell spread) (Fig. 5, D, F, and H). These results indicate that most mice immunized with either vaccine formulation produced antibodies to the gD2 epitopes, although trivalent mRNA outperformed trivalent subunit proteins by producing higher titers of blocking antibodies to three of six epitopes.

CD4+ and CD8+ T cell responses in mice

Splenocytes were harvested 2 weeks after the second trivalent mRNA or third trivalent subunit protein immunization. CD4+ and CD8+ T cells were stimulated in vitro with a single gD2 overlapping peptide pool, and gC2 or gE2 overlapping peptides each divided into two pools. The peptide pools span the entire amino acid sequences of the subunit proteins. As a control, T cells were stimulated with dimethyl sulfoxide (DMSO), the vehicle used to dissolve the overlapping peptides. CD4+ T cells in mice immunized with trivalent mRNA significantly increased production of IFNγ alone or both IFNγ and tumor necrosis factor α (TNFα) when stimulated with gD2, gC2, or gE2 peptide pools, compared with DMSO alone (Fig. 6, A and B). Trivalent subunit protein immunization was less effective at stimulating CD4+ T cell cytokine production (Fig. 6, D and E). The trivalent mRNA and subunit protein groups did not produce robust CD8+ T cell responses. In the mRNA mice, only gE2 stimulated a significant increase in CD8+ IFNγ production, whereas in the protein subunit group, gD2 pool 1 stimulated an increase in CD8+ IFNγ production that approached statistical significance (Fig. 6, C and F). We conclude that mice immunized with trivalent mRNA stimulated a more robust CD4+ T cell response than mice immunized with trivalent subunit proteins and a marginally better CD8+ T cell response.

DISCUSSION

The virus titer that results in genital herpes infection in humans is unknown, making it difficult to choose the optimal challenge dose in animals. We selected a high challenge dose by using 275 or 2750 LD50 of HSV-2 in mice and 50 LD50 in guinea pigs. Despite these high doses, no mouse immunized ID or IM with 10 μg of trivalent nucleoside-modified mRNA-LNP and no guinea pig immunized with 20 μg of trivalent nucleoside-modified mRNA-LNP developed genital disease, although it is possible that, at even higher challenge doses, some breakthrough infection may occur. Mice and guinea pigs were also well protected against subclinical infection. Commonly used indicators of subclinical infection in mice include vaginal swab cultures on days 2 and 4 and HSV-2 DNA in DRG, whereas a useful marker for subclinical infection in guinea pigs is vaginal shedding of HSV-2 DNA after resolution of the acute infection (41–43). By these criteria, only 1 of 64 mice and 2 of 10 guinea pigs developed subclinical infection. Overall, guinea pigs shed HSV-2 DNA on 2.4% of days, and no replication-competent virus was recovered in the vaginal secretions, suggesting that the risk of transmission is very low in these animals. Our decision to immunize with three antigens was based on our previous studies that reported that adding gC2 to gD2 subunit protein improved protection in mice and guinea pigs, and although performed in different studies, we noted that adding gE2 to gC2/gD2

CD4+ T cell responses and GC B cell responses. Splenocytes for CD4+ and CD8+ T cell assays were pooled from five animals, and the responses of six replicate wells are shown. (A) CD4+ T cell IFNγ responses in mice immunized with trivalent mRNA or (B) CD4+ polyfunctional IFNγ and TNFα responses in mice immunized with trivalent mRNA. (C) CD8+ T cell IFNγ responses in mice immunized with trivalent mRNA. (D) CD4+ T cell IFNγ responses in mice immunized with trivalent subunit protein or (E) CD4+ polyfunctional IFNγ and TNFα responses in mice immunized with trivalent subunit protein. (F) CD8+ T cell IFNγ responses in mice immunized with trivalent subunit protein. (A to F) Each peptide-stimulated CD4+ or CD8+ T cell assay was compared with DMSO vehicle-treated controls. The CD4+ and CD8+ T cell assays for trivalent mRNA and trivalent subunit protein were run on separate days, which accounts for the variability in the DMSO control results. The percentage of cytokine positive cells is based on the total number of CD4+ T cells counted in the trivalent mRNA and subunit protein assays. The mean numbers of CD4+ T cells counted in the trivalent mRNA experiments and in the trivalent subunit protein were 981 ± 232 and 1118 ± 206, respectively. All values with P < 0.05 are shown in the figures. P values were calculated by the Kruskal-Wallis test with Dunn’s for multiple comparisons. (G and H) Tfh and GC B cell responses that include CpG/alum as a control. Splenocytes were harvested for Tfh and GC B cell assays from individual mice 10 days after a single immunization with CpG/alum alone, trivalent mRNA-LNP, trivalent subunit protein–CpG/alum, or no immunization. P values are shown in the figures for the primary end point comparison between trivalent mRNA and trivalent protein. Other P values are noted below. (G) Tfh cells: P = 0.532, comparing CpG/alum with trivalent subunit protein–CpG/alum. (H) GC B cells: P = 0.90, comparing CpG/alum with trivalent subunit protein. n = 5 animals per group, except n = 4 for CpG/alum. P values were calculated using the two-tailed Mann-Whitney test. Error bars represent geometric means and 95% confidence intervals. (I and J) Tfh and GC B cell responses that include PolyC RNA-LNP as a control. Splenocytes were harvested for Tfh and GC B cell assays from individual mice 10 days after the second immunization with PolyC RNA-LNP, trivalent mRNA-LNP, or no immunization. P values are shown in the figures and were calculated by the two-tailed Mann-Whitney test; n = 5 animals per group.
subunit proteins further improved protection in mice (16, 21, 22). These results are consistent with our current study demonstrating that trivalent mRNA-LNP provides better protection than gD2 mRNA-LNP alone.

Trivalent nucleoside-modified mRNA-LNP immunization outperformed the same immunogens administered as subunit proteins with CpG/alum in mice and guinea pigs. Our choice of 10 μg of each nucleoside-modified mRNA in mice was partially guided by dose-response studies using nucleoside-modified mRNA to express firefly luciferase and vaccine studies of Zika, influenza, and HIV antigens in mice (29, 30, 44). The choice of 10 μg seems appropriate based on the nearly complete protection provided by trivalent mRNA at this concentration and lower efficacy at 2 μg. Immunization with 20 μg of each trivalent nucleoside-modified mRNA in guinea pigs was based on the larger body mass of guinea pigs than mice. Our choice of 5 μg for subunit proteins in mice and 10 μg in guinea pigs was based on our previous results demonstrating excellent protection in mice and guinea pigs against genital lesions at these concentrations (16, 22). A limitation of our study is that we did not evaluate whether higher concentrations of trivalent modified mRNA-LNP or trivalent subunit proteins will further improve vaccine efficacy. We also did not assess the durability of immunity and protection. The potent T FH and GC B cell responses produced by the trivalent mRNA immunogens suggest that durable protection is likely. Impressive durability was demonstrated for HIV, influenza, and Zika when animals were immunized by the same nucleoside-modified mRNA-LNP platform as used here (29, 30). The high antibody titers and potent T FH and GC B cell responses noted with other nucleoside-modified mRNA-LNP immunogens, including influenza, Zika, and HIV, were key reasons that we chose to evaluate nucleoside-modified mRNA as a delivery mechanism for our candidate HSV-2 vaccine (25, 30).

We performed multiple immunology assays to assess mechanisms by which the trivalent nucleoside-modified mRNA outperformed the trivalent protein vaccine. Where differences emerged, mRNA was superior including (i) serum and vaginal IgG ELISA titers; (ii) HSV-2 neutralizing antibody titers; (iii) antibodies that block gC2 and gE2 immune evasion domains; (iv) antibody responses to gD2 epitopes involved in binding to receptors, interaction with gH2/gL2, and cell-to-cell spread; (v) CD4+ T cell responses; and (vi) T FH and GC B cell responses. It seems likely that a combination of these immune responses accounts for the enhanced performance of the trivalent mRNA vaccine. The prolonged expression of antigens using nucleoside-modified mRNA-LNP delivery and the T FH adjuvant activity of LNPs may explain the more potent immune responses (44, 45). Although trivalent nucleoside-modified mRNA outperformed trivalent subunit protein–CpG/alum, it is possible that new adjuvants may improve the efficacy of subunit protein vaccines.

Few preclinical HSV-2 vaccine studies have been published that evaluated the same antigens and adjuvants that were subsequently used in human trials. One exception is the gD2 subunit protein vaccine administered with monophosphoryl lipid A (MPL)/alum that was evaluated in a guinea pig model of genital herpes. The gD2-MPL/alum vaccine prevented genital lesions in 9 of 10 guinea pigs in one study and 10 of 12 in another, but all animals developed subclinical infection as measured by vaginal shedding of HSV-2 DNA (46, 47). In our current study, the trivalent nucleoside-modified mRNA-LNP vaccine prevented genital lesions in 100% of guinea pigs and subclinical genital shedding of HSV-2 DNA in 80%, which surpasses the results obtained using an immunogen that entered phase 3 human trials. We conclude that the trivalent nucleoside-modified mRNA-LNP vaccine is a promising candidate for future human trials.

**MATERIALS AND METHODS**

**Study design**

The primary goal was to assess whether the trivalent nucleoside-modified mRNA-LNP vaccine outperformed the same antigens expressed as baculovirus subunit proteins administered with CpG/alum.

A secondary goal was to evaluate whether adding gC2 and gE2 nucleoside-modified mRNA-LNP to gD2 nucleoside-modified mRNA-LNP improved the protection provided by gD2. Mice and guinea pigs were immunized with nucleoside-modified mRNA-LNP immunogens or baculovirus subunit protein–CpG/alum antigens. Serum and vaginal wash fluids were obtained approximately 1 month after the final immunization, and animals were then infected intravaginally with HSV-2 strain MS. In mice, vaginal swabs were obtained for cultures on days 2 and 4 after infection; animals were weighed daily for 2 weeks and scored for genital disease for 4 weeks; and DRG were obtained either on day 4 after infection or at the end of the experiment. In guinea pigs, vaginal swabs were obtained for cultures on days 2 and 4 after infection; animals were scored for genital disease for 60 days; swabs were obtained for vaginal shedding of HSV-2 DNA for 21 days beginning 28 days after infection; and DRG and spinal cord were harvested for HSV-2 DNA at the end of the experiment. Scoring for genital disease in mice was performed without blinding, whereas scoring in guinea pigs was done blinded.

**Ethics statement**

All animal studies were conducted under protocol 805187 approved by the University of Pennsylvania Institutional Animal Care and Use Committee.

**Production of bac-gC2, bac-gD2, and bac-gE2 subunit proteins; production of mRNA; formulation in LNPs; and Western blots**

The baculovirus proteins were produced as previously described (22, 48, 49). The methods used to synthesize trivalent nucleoside-modified mRNA-LNP and to perform Western blots are described in the legend of fig. S1.

**ELISA, neutralizing antibody, and ADCC titers**

Serum IgG ELISA titers and IgG1/IgG2a subtype titers were evaluated as previously described (16, 50). ELISA vaginal wash fluid titers were assessed by adding 30 μl of Dulbecco’s modified Eagle’s medium (DMEM) with 5% fetal bovine serum to the vaginal cavity and by retrieving the fluid several seconds later. This procedure was repeated once. Mucosal IgG antibody in vaginal fluid was measured at a dilution of 1:50. Neutralizing antibody titers were reported as the serum dilution that reduced the number of virus plaques by 50% (21). ADCC was measured using the murine FcγRIII ADCC Reporter Bioassay (Promega Corporation) that binds murine IgG1 and IgG2a (23, 51).

**Blocking C3b binding to gC2 and IgG Fc binding to gE**

Serum (10 μl) from each mouse in a group was pooled, and the IgG was purified on the NAβ Protein G Spin Column (Pierce Biotechnology). Blocking gC2 binding to C3b was performed by coating ELISA plates with 200 ng of purified C3b protein (Complement Technology), incubating 40 ng of purified gC2 protein with varying concentrations

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of purified IgG, and detecting bound gC2 (34). Blocking gE2 binding to IgG Fc was evaluated using HSV-1/HSV-2 seronegative human IgG (1 mg per well) to coat ELISA wells, by incubating purified IgG (62.5 μg/ml) from immunized mice with bac-gE2 (24-405T) (2 or 4 μg/ml) for 15 min at room temperature, and by detecting bound gE2 (16).

Biosensor-based antibody competition assay
A Cartera Microfluidics continuous flow microspotter surface plasmon resonance imaging system was used to detect epitope-specific antibody responses to crucial gD2 epitopes involved in virus entry and cell-to-cell spread (23, 24).

CD4+ and CD8+ T cell responses and Tfh and GC B cell responses
Spleens were pooled from five animals per group, and 10^6 splenocytes were stimulated at 37°C with gC2, gD2, or gE2 peptide pools consisting of 15 amino acids with 11 overlapping amino acids (JPT Peptide Technologies). After 1 hour, brefeldin A (10 μg/ml) was added for 16 hours at 37°C. Spleen cells were stained with aqua blue (Invitrogen) to distinguish live-dead cells, Pacific blue–conjugated anti-CD8 mouse mAb (BioLegend), and R-phycoerythrin–cyanine 5.5 (PE-Cy5.5) anti-CD4 mouse mAb (BD Pharmingen). Cells were permeabilized with Cytofix/Cytoperm (BD Pharmingen) and stained with Alexa Fluor 700–conjugated anti-IFNγ mouse mAb, PE-Cy7 anti-TNFα mouse mAb, and allophycocyanin–Cy7 anti-CD3 mouse mAb. Spleen cells were fixed with 1% paraformaldehyde and analyzed by fluorescence-activated cell sorter using an 18-color LSR II flow cytometer and FlowJo flow cytometry analytic software (21, 52, 53). Details of the staining and gating strategy for Tfh and GC B cells are presented in the legend of fig. S4.

Immunizations, vaginal infections, and scoring BALB/c mice (Charles River Laboratories)
The gC2, gD2, and gE2 baculovirus subunit proteins were individually incubated at room temperature for 2 hours with 16.7 μg of CpG oligonucleotide 5′-TCCATGAGCTTCCGACGT-3′ (Coley Pharmaceutical) and 25 μg of alum per microgram of protein (Alhydrogel, Accurate Chemical and Scientific Corp.) and combined before immunization in a volume of 50 μl (22). Mock IM immunizations were performed with CpG and alum without proteins. The trivalent (gC2, gD2, and gE2) mRNA-LNP were administered in four sites on the flank using a 29-gauge insulin syringe containing 30 μl for each site, whereas LNP immunizations with LNP-containing immunogens were given in 50 μl into the gastrocnemius muscle. Mice were infected intravaginally 1 month after the final immunization and scored for genital disease (22).

Hartley strain guinea pigs (Charles River Laboratories)
Ten micrograms of gC2, gD2, and gE2 subunit proteins was mixed individually and then combined to form a final concentration of 100 μg of CpG (5′-TGGTTGTTGGTTTGTGTT-3′; Trilink Technologies Inc.) per guinea pig and 150 μg of alum (16). ID immunizations and intravaginal infections 1 month after the final immunization were performed as in mice, except that 40 μl was inoculated ID at each site (16).

Virus cultures from vaginal swabs
Vaginal swabs were placed in 1 ml of DMEM containing 5% fetal bovine serum supplemented with vancomycin (25 μg/ml). For viral titers, 150 and 300 μl were evaluated in mice and guinea pigs, respectively. Serial 10-fold dilutions were added to Vero cells to assess virus titers by plaque assay (21).

HSV-2 DNA isolation and real-time qPCR for HSV-2 DNA copy number in DRG, spinal cord, and vaginal swab samples
Mouse DRG samples were analyzed in duplicate for HSV-2 DNA, as previously described, except that separate reactions were used to amplify HSV-2 DNA and the mouse adipsin gene and DRG HSV-2 DNA copy number was expressed as log10 DNA copies per 10^5 adipsin genes instead of 10^4 (22). HSV-2 DNA copy number in guinea pig DRG and spinal cord samples was calculated as log10 DNA copies per 10^5 gD2 mRNA-LNP. We used the Mann-Whitney test for these two-group comparisons. Some P values are also noted in the figure legends. For some comparisons, the mice in one treatment group all had the same response (e.g., zero as the lower limit of quantification), and the two-tailed Fisher’s exact test was used to compare the values. For comparisons looking across several time points, we used two-way analysis of variance (ANOVA) followed by Tukey adjustment for significance. We used the log-rank test to compare survival distributions between groups. We used Wilcoxon signed-rank test to compare samples from the same animal at different time points. All significance tests were two-sided and performed at P < 0.05. Analyses were done using GraphPad Prism version 7.0 (GraphPad Software Inc.).

SUPPLEMENTARY MATERIALS
immunology.sciencemag.org/cgi/content/full/4/39/eaaw7083/DC1
Fig. S1. Western blot and ELISA demonstrating expression and secretion of gC2, gD2, and gE2 subunit proteins after transfection with nucleoside-modified mRNA-LNP.
Fig. S2. Transmission studies.
Fig. S3. IgG1 and IgG2a titers.
Fig. S4. Flow cytometry gating strategy for the Tfh and GC B cells.
Data file S1. Raw data (Excel).

REFERENCES AND NOTES


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Acknowledgments: We thank B. L. Mui and Y. K. Tam (Acutas Therapeutics, Vancouver, BC, Canada) for their role in formulating the nucleoside-modified mRNA into LNPs. Funding: H.M.F. and S.A. were supported by NIH NIAID grants R01 AI104854 and AI139618. G.H.C. was supported by NIH NIAID grant R01 AI118289. D.W. was supported by NIH NIAID grants R01 AI050484, R01 AI124429, and R01 AI084860; Gates Foundation CAVID OPP1033102; and the New Frontier Sciences division of Takeda Pharmaceuticals. P. Shaw (Department of Biostatistics, Epidemiology and Informatics, Perelman School of Medicine) provided biostatistical support through core services from the Penn Center for AIDS Research (CFAR) NIH grant P30 AI045008. Author contributions: S.A. designed and performed the murine studies and most of the immunology studies, prepared most of the figures, and performed some statistical analysis in collaboration with P. Shaw in the Department of Biostatistics. L.M.H. assisted with the murine studies, performed epitope mapping, and prepared the final version of figures. N.P. prepared the nucleoside-modified mRNA, performed cell transfections, and assisted with murine studies. F.W. performed qPCR for HSV-2 DNA. A.M. and M.P.C. performed the Th1 and GC B cell studies. G.H.C. supervised preparation of subunit proteins and epitope mapping. D.W. supervised the preparation and planning of nucleoside-modified mRNA-LNP studies. H.M.F. oversaw the entire project, performed some of the statistical analyses in collaboration with P. Shaw, and wrote the manuscript. Competing interests: In accordance with the University of Pennsylvania policies and procedures and our ethical obligations as researchers, we report that H.M.F., S.A., and G.H.C. are named on patents that describe the use of multiple subunit glycoprotein antigens for HSV vaccines. H.M.F., S.A., and D.W. have submitted a patent claim that uses nucleoside-modified mRNA as a vaccine for HSV. D.W. is named on patents that describe the use of nucleoside-modified mRNA as a platform to deliver therapeutic proteins. D.W. and N.P. are also named on a patent describing the use of nucleoside-modified mRNA in LNP vaccines as a vaccine platform. We have disclosed those interests fully to the University of Pennsylvania, and we have in place an approved plan for managing any potential conflicts arising from licensing of our patents. Data and materials availability: All data are available within the article and will be made available upon request. Access to subunit antigens must be obtained through a material transfer agreement.

Submitted 17 January 2019
Resubmitted 16 April 2019
Accepted 30 July 2019
Published 20 September 2019
10.1126/sciimmunol.aaw7083

Nucleoside-modified mRNA encoding HSV-2 glycoproteins C, D, and E prevents clinical and subclinical genital herpes

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DOI: 10.1126/sciimmunol.aaw7083

Hinting at a herpes vaccine
Currently, a vaccine for genital herpes does not exist despite the prevalence of this sexually transmitted disease. Previous attempts to make vaccines against herpes simplex virus 2 (HSV-2) included trials with protein subunit vaccine candidates that delayed vaccine onset but were not protective. Awasthi et al. describe a vaccine candidate that is composed of nucleoside-modified mRNA in lipid nanoparticles that encodes the HSV-2 glycoproteins C, D, and E. This trivalent vaccine protected mice and guinea pigs from developing genital lesions and reduced viral shedding. Neutralizing antibody and CD4+ T cell responses were detected in immunized mice. These results suggest that an mRNA-based HSV-2 vaccine may have potential for further preclinical development.