INTRODUCTION
Vaccination significantly reduces the health burden of many infectious disease, especially in high-risk populations. Pregnant women, unborn fetuses, and neonates represent three populations of high-risk individuals that can all be simultaneously protected from vaccine-preventable infectious disease with strategic maternal immunization protocols. Infectious microbes that pose significant health risks during pregnancy can be divided into three broad categories, based on the pathogenesis and disease outcome (Fig. 1), with some microbes falling within more than one category. First are maternal infections, which are defined by heightened disease severity in pregnant females, but with rare or inconsequential transmission and disease in the fetus. Second are fetal or congenital infections, which are characterized by mild or no disease in pregnant females, but occasional vertical transmission and severe congenital disease in the fetus. Third are neonatal and infant infections, which are not considered to pose significant risk to pregnant women or unborn fetuses, but can cause severe, and sometimes fatal disease in neonates and infants that lack protective maternal immunity following birth.

The vaccination strategies employed differ for microbes within each of these categories and vary based on the at-risk individual (i.e., mother, fetus, and/or neonate/infant), the timing of the greatest risk of infection (i.e., early pregnancy, late pregnancy, or post-natal), and on the duration of protective immunity following vaccination. In this review, we discuss evidence to suggest that immunization strategies for pregnant women should be tailored to optimize protection for the mother, fetus, neonate, infant, or all individuals. We review vaccine-preventable infections during pregnancy and the current vaccination strategies employed to reduce the burden of infectious diseases, including influenza. Further, we examine novel vaccine platforms and consider how their application may provide safe alternatives for enhancing protection of pregnant women. Finally, we discuss vaccine development and prevention strategies for combating emerging infectious diseases, including Zika, that pose a threat to pregnant women and their fetuses.

VACCINATION AGAINST MATERNAL INFECTIONS
Owing to physiologic and immunologic changes that support pregnancy and tolerance of a semi-allogenic fetus,1 pregnant women demonstrate increased susceptibility to certain infectious agents including hepatitis E, varicella zoster, and influenza viruses. Infection with these viruses during pregnancy results in severe maternal disease, increased maternal mortality and associated pregnancy complications, which are observed most frequently during the third trimester and peripartum period. For example, the case fatality rate among pregnant women infected with hepatitis E virus is estimated to be 5–25%,2,3 compared with 1–3% in the general population.4 Approximately 28% of cases of varicella pneumonia in adults reported from 1965–1989 were from pregnant women5; and pregnant women infected with the pandemic 2009 H1N1 influenza A virus (IAV) were reportedly 4 times more likely to be hospitalized or die than the general population.6 Overall, vertical transmission of these viruses is relatively uncommon, but adverse pregnancy outcomes, including spontaneous abortion and pre-term birth can still occur as an indirect consequence of maternal inflammation.7,8 Reports during the 2009 H1N1 pandemic in Australia and New Zealand indicated that among pregnant women who were hospitalized with suspected H1N1 IAV infection, 50% of their infants required intensive care, and 10% were either stillborn or died shortly after birth; only 2 infants, however, had detectable 2009 H1N1 infection.9

The primary goal of vaccination strategies for protecting against maternal infections is the generation of protective maternal immunity either prior to or during early pregnancy. Optimally, vaccination should prevent or reduce disease by inducing sterilizing immunity (i.e., immunity that completely prevents infection). Despite reported reductions in antiviral proteins during pregnancy and tolerance of a semi-allogenic fetus,1 pregnant women demonstrate increased susceptibility to certain infectious agents including hepatitis E, varicella zoster, and influenza viruses. Infection with these viruses during pregnancy results in severe maternal disease, increased maternal mortality and associated pregnancy complications, which are observed most frequently during the third trimester and peripartum period. For example, the case fatality rate among pregnant women infected with hepatitis E virus is estimated to be 5–25%,2,3 compared with 1–3% in the general population.4 Approximately 28% of cases of varicella pneumonia in adults reported from 1965–1989 were from pregnant women5; and pregnant women infected with the pandemic 2009 H1N1 influenza A virus (IAV) were reportedly 4 times more likely to be hospitalized or die than the general population.6 Overall, vertical transmission of these viruses is relatively uncommon, but adverse pregnancy outcomes, including spontaneous abortion and pre-term birth can still occur as an indirect consequence of maternal inflammation.7,8 Reports during the 2009 H1N1 pandemic in Australia and New Zealand indicated that among pregnant women who were hospitalized with suspected H1N1 IAV infection, 50% of their infants required intensive care, and 10% were either stillborn or died shortly after birth; only 2 infants, however, had detectable 2009 H1N1 infection.9

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pregnancy, studies comparing vaccine responses between pregnant and nonpregnant women find no difference in either the magnitude or duration of antibody responses against influenza A viruses. In fact, surveillance data from Taiwan reveal that influenza vaccination during pregnancy results in higher levels of seroprotection than does vaccination prior to conception, with no effect of gestational age on vaccine-induced antibody responses. As of 2004, the Centers for Disease Control (CDC) Advisory Committee on Immunization Practices (ACIP) categorizes pregnant women as a target population for receiving the inactivated influenza vaccine and recommends that pregnant women be immunized during any trimester. Several studies evaluating adverse vaccine reactions in pregnant women have concluded that there is no link between pregnancy complications or adverse fetal outcomes among women who are vaccinated during pregnancy. Although the live attenuated intranasal influenza vaccine is not recommended for pregnant women, accidental administration during pregnancy was not associated with an increased risk of adverse reactions. Despite the plethora of data that support the benefits and safety of influenza vaccination during pregnancy, coverage remains low, with a less than 50% maternal vaccination rate during the 2010–2011 influenza season in the United States. Misconceptions about the safety and benefits of influenza vaccination represent the largest barriers to vaccine acceptance among pregnant women.

Varicella zoster virus (VZV) is another vaccine-preventable infection associated with increased severity during pregnancy. VZV is an alpha herpes virus and the causative agent of varicella or chickenpox. In temperate climates, seroprevalence among individuals over 30 years of age is estimated to be 95%, with almost 90% of infections occurring prior to 15 years of age. The first modified-live vaccine against varicella zoster virus was licensed in the United States in 1995, and is now recommended for children under 12 months of age. Primary VZV infection during pregnancy is therefore uncommon, as most women of childbearing age have been either infected or immunized. In women who have not been previously exposed, however, primary VZV infection between weeks 8 through 26 of gestation is associated with a 2% risk of congenital transmission and disease in the offspring. Because all licensed VZV vaccines contain live-attenuated virus, their use during pregnancy is contraindicated. Instead, the CDC recommends that nonpregnant women of childbearing age be vaccinated against VZV at least one month prior to conception. As a herpes virus, infection with VZV is life-long, and reactivation occurs in approximately 10–30% of individuals, which results in a painful skin condition known as shingles or herpes zoster.

Acute viral hepatitis caused by hepatitis E virus (HEV) is an emerging infectious disease that causes severe disease in pregnant women, with a fatality rate of up to 30% in endemic regions. In addition to heightened maternal disease severity, HEV infection during pregnancy is associated with increased rates of premature birth and prenatal mortality. Although vertical HEV transmission rates are high, with estimates between 23–50%, the relative contributions of fetal HEV infection to adverse perinatal outcomes is unclear. A recombinant HEV subunit vaccine has been developed and proven safe and effective following completion of Phase II and III clinical trials, but commercial use is currently limited to China. Furthermore, the vaccine is not approved for use in pregnant women despite being 100% efficacious in participants receiving all three doses. Additional HEV vaccine candidates are being tested in preclinical pregnant animal models, and one recombinant HEV vaccine has been shown to be safe and highly immunogenic in pregnant mice. Additional studies in a susceptible animal model are needed to confirm efficacy following virus challenge.

**Vaccination Against Congenital Infections**

Developing fetuses are extremely vulnerable to both infectious and noninfectious insults. Certain infectious agents that are often clinically silent in healthy adults can cause severe birth defects if
they breach the placental barrier during critical developmental periods during pregnancy. An increasing number of pathogens are being recognized for causing congenital disease, and what was originally designated as the TORCH complex (Toxoplasma gondii, “other,” Rubella virus, Cytomegalovirus, and Herpes Simplex virus) is now expanded to include other infectious agents including Zika virus. Development of congenital disease can depend on the timing of infection during gestation, the infectious burden, and the pathogenesis in the fetus. The congenital syndrome for each pathogen is characterized by a variety of different developmental abnormalities, and commonly impact hearing, vision, and central nervous system function. For many congenital infections, the timing of infection during gestation determines the relative risk to the fetus and dictates the spectrum of disease that results. For example, while infection with rubella virus during the first 9 weeks of gestation is associated with an 85% risk of congenital rubella syndrome (CRS), this is reduced to a 52% risk between 9–12 weeks, and minimal risk for infections occurring after 16 weeks of gestation. In contrast, the risk of congenital toxoplasmosis has been demonstrated to be highest during third trimester pregnancy, which is hypothesized to be due to differential expression of placental toll-like receptors, including TLR6, within first compared with third trimester trophoblast cells. Similarly, maternal infection with Listeria monocytogenes is typically associated with adverse pregnancy outcomes during the third trimester, though infection during the first trimester in nonhuman primates also leads to rapid fetal demise.

The primary goal of vaccination strategies for protecting against fetal infections is generation of protective maternal immunity prior to pregnancy. Because congenital infections can occur in the absence of maternal symptoms, vaccines against congenital agents should ideally provide complete sterilizing immunity. Rubella is included in a live-attenuated combination vaccine for measles, mumps, and rubella (MMR), which confers lifelong protective immunity. Because of the long duration of protective immunity following rubella vaccination, target populations include children and adolescent girls. However, incomplete vaccination coverage can lead to paradoxical increases in CRS due to an increase in the average age of infection, and so it is also recommended that unvaccinated women of childbearing age be counseled to receive the rubella vaccine at least one month prior to conception. The implementation of large-scale rubella vaccination programs has resulted in sufficient population-level immunity, significant reductions in CRS, and elimination of rubella virus from several developed countries, including the United States.

Following successful implementation of MMR vaccination programs, cytomegalovirus (CMV) has emerged as the most common congenital viral infection in the developed world. The incidence of congenital CMV varies based on geographic region and socioeconomic status, but overall birth prevalence is estimated to be 0.64%, which is similar to the incidence of Down syndrome and fetal alcohol syndrome. In contrast to rubella virus, however, there is currently no licensed vaccine available for CMV, and with seroprevalence approaching 100% in some developing countries, vaccine development has been identified as a priority public healthcare goal. While CMV infection of healthy adults is usually asymptomatic, adaptive immune responses are insufficient to clear the infection, which results in lifelong latent infection of myeloid precursor cells. Although latent or reactivated CMV is less likely to cause congenital infection than a primary CMV infection during pregnancy, neonatal infections is generation of robust maternal antibody prior to conception. Vaccination during pregnancy as opposed to before conception is likely to result in the greatest benefit to the neonate. Further, the efficiency of placental antibody transfer in females increases throughout gestation, with less than 8% maternal IgG transferred to the fetus through the placenta prior to birth, which contrasts with most veterinary species, in which maternal antibody is transferred via colostrum immediately following birth. Regardless of species, vaccination during pregnancy increases circulating maternal antibodies and enhances transfer to the fetus/neonate.

VACCINATION AGAINST NEONATAL AND INFANT INFECTIONS

Owing to the limited exposure to foreign antigen and blunted innate immune responses in utero, the neonatal immune system is immature at birth, making neonates (i.e., less than one month of age) particularly susceptible infections. Infectious diseases are responsible for over 60% of child mortality, and over 40% of these deaths occur within one month of age. During the neonatal period of immune system maturation, protection against pathogens relies primarily on passive immunity from maternal-derived IgG antibodies. In humans, most maternal antibodies are transferred into the fetal circulation through the placenta prior to birth, which contrasts with most veterinary species, in which maternal antibody is transferred via colostrum immediately following birth.

The goal of vaccination strategies for protecting against neonatal infections is generation of robust maternal antibody responses during pregnancy to enhance placental transfer. Further, because neonatal protection is exclusively conferred by maternal-derived antibody, vaccines aimed at protecting infants should prioritize induction of humoral over cellular immune responses, with the induction of IgG1 being most important because this IgG isotype is associated with the highest placental transport efficiency in females. Moreover, the kinetics of maternal vaccine-induced antibody response, the efficiency of placental antibody transfer, and the half-life of the antibody in the neonate should inform the optimal timing of vaccination during pregnancy. Because the peak antibody response is typically observed 1–3 weeks following immunization, vaccination during pregnancy as opposed to before conception is likely to result in the greatest benefit to the neonate. Further, the efficiency of placental antibody transfer in females increases throughout gestation, with less than 8% maternal IgG transferred to the fetus.
in the first 16 weeks of gestation.\textsuperscript{65} Significantly more transferred during the second and third trimesters, and at delivery fetal IgG often exceeds maternal levels.\textsuperscript{56} Vaccination of females during the second and third trimesters of pregnancy is most likely to generate the greatest level of protection in the neonate, but the precise timing for maximum protection is debated. Controversy over the timing of pertussis vaccination in pregnancy has been reviewed elsewhere,\textsuperscript{67} with some reports claiming peak cord blood antibody concentrations following vaccination in the second trimester, and others reporting peak antibody concentrations following vaccination in the third trimester. Antibody avidity also influences the efficiency of placental transfer, with higher avidity antibodies crossing the placenta with greater efficiency than low avidity antibodies.\textsuperscript{68} More consideration should be given to the development of high avidity antibodies in the timing of vaccination during pregnancy, as protective immunity in the infant depends on both the concentration and avidity of the maternal-derived antibody.

The half-life of maternal antibodies in infants also must be considered in vaccine development and administration. Maternal-derived IgG1 is reported to have a half-life of approximately 48 days in serum,\textsuperscript{69} and depending on serum antibody titers present at birth, this translates into protective immunity for approximately the first 3–9 months of life for most infant pathogens.\textsuperscript{70} The half-life of the antibody also dictates the vaccination schedules for infants, as the presence of maternal-derived antibody interferes with vaccine efficacy, and it is not until maternal-derived antibody has waned below a certain threshold that an infant can mount its own active vaccine response. The goal of the infant vaccine series is to time vaccination to coincide with the time that maternal-derived antibody drops below the threshold at which it can neutralize the vaccine antigen. Because the precise timing of these events is unpredictable, infant vaccination schedules are designed so that vaccines are administered in a series that spans the duration of this window, and minimize susceptibility to natural infection. In the United States, infant vaccines are recommended at 2, 4 and 6 months of age.\textsuperscript{41}

*Bordetella pertussis* is a vaccine-preventable respiratory pathogen of significant public health importance, and it is a major cause of mortality in infants lacking protective maternal immunity. Vaccination of women during pregnancy, however, significantly enhances the transfer of maternal antibody to the fetus,\textsuperscript{71,72} and these newborns are 11 times more likely to have protective antibody titers at birth compared with those born from women who were not vaccinated during pregnancy.\textsuperscript{71} Inactivated pertussis antigen is combined with tetanus and diphtheria toxoids in a single vaccine (Tdap), which the CDC ACIP recommends for all pregnant women, regardless of previous vaccine history.\textsuperscript{73} In contrast to vaccine formulations that contain killed whole *B. pertussis* organisms, the Tdap vaccine contains only select antigens and confers relatively weak and only transient protective immunity that declines after 1 year.\textsuperscript{74} Vaccination of women either prior to conception or during early pregnancy does not provide adequate neonatal protection against pertussis.\textsuperscript{73} Consequently, the CDC considers the third trimester to be the optimal time to administer the Tdap vaccine to pregnant women.\textsuperscript{75} Adverse events reported following Tdap vaccination are generally mild, and there are no reported risks of adverse pregnancy outcomes related to Tdap vaccination during pregnancy.\textsuperscript{76} Despite consistent evidence that supports the benefit and safety of Tdap vaccination during pregnancy, coverage remains low, with an estimated 42% of pregnant women receiving the Tdap vaccine in the United States in 2013.\textsuperscript{77}

Receipt of the Tdap vaccine during pregnancy also confers protection against neonatal tetanus, which is associated with case fatality approaching 100% in the absence of medical care.\textsuperscript{78} Disease is caused by the toxin produced by *Clostridium tetani*, and infection occurs most commonly due to contamination of the umbilical stump following delivery. Consequently, the incidence of disease is much greater in developing countries, where maternal vaccination is scarce and perinatal hygiene practices are poor.\textsuperscript{79} In 1989, the World Health Assembly called for the elimination of neonatal tetanus, which has inspired an initiative to improve vaccination coverage and birth hygiene in 59 countries with high disease prevalence. As part of this initiative, immunization standards have been expanded and recommend that pregnant women with unknown or inadequate vaccination history receive two doses of the toxoid-containing vaccine, administered one month apart.\textsuperscript{80} Maternal anti-tetanus antibodies are passively transferred to the fetus, and it is estimated that maternal immunization reduces neonatal tetanus mortality by 94%.\textsuperscript{81}

Respiratory syncytial virus (RSV) is the most common respiratory viral pathogen of newborns and infants, and accounts for 50–90% of acute bronchiolitis and 5–20% of pneumonia cases in hospitalized children less than 2 years of age.\textsuperscript{82} RSV is also reported to cause severe disease and hospitalization in pregnant women when infection occurs during the third trimester,\textsuperscript{83,84} and therefore dualy qualifies as a maternal infection as well. A licensed vaccine against RSV is currently unavailable, but several vaccine candidates have shown promise in various animal models.\textsuperscript{85–88} Given the importance of this pathogen during early life, vaccine development strategies have focused on maternal immunization, with three maternal vaccines currently in clinical trials.\textsuperscript{89} Maternal vaccination against RSV has direct and indirect benefits to the neonate; neonates are directly protected through passive transfer of maternal antibody through the placenta, and they are indirectly protected because a vaccinated mother is less likely to transmit the infection to her infant.\textsuperscript{90}

**CONTRIBUTIONS OF PREGNANT ANIMAL MODELS**

Vaccination of pregnant women is controversial, and immunization with live (i.e., replication-competent) viral or bacterial vaccines is generally contraindicated due to the theoretical risk of congenital infection and teratogenic effects from the vaccine strains. However, in a report of over 2000 pregnant women who were unknowingly immunized with live attenuated rubella vaccine, there were no cases of vaccine-associated congenital rubella infection,\textsuperscript{91} and live virus strains of influenza or yellow fever viruses administered to pregnant women also have no link with pregnancy complications.\textsuperscript{21,92} Vaccination with inactivated vaccines such as influenza and Tdap during pregnancy have low uptake, with concerns of safety among both patients and their healthcare providers being a primary barrier. The safety of vaccine adjuvants is debated, and although neither the Tdap nor seasonal influenza vaccine recommended during pregnancy contain adjuvants, retrospective studies evaluating safety of the adjuvanted pandemic H1N1 influenza vaccine in pregnant women found no relationship with adverse pregnancy outcomes.\textsuperscript{93} The conservative approach to vaccination protocols for pregnant women stems from the lack of controlled safety and efficacy studies for this population. For ethical reasons, pregnant women are exempted from almost all clinical and vaccine trials, and health care providers are less likely to endorse prophylactic treatments for which safety and efficacy profiles have not been adequately characterized.

Whereas study in pregnant women is not possible, pre-clinical testing in animal models may provide a useful alternative, and vaccine preclinical trials in pregnant animal models may provide information to inform healthcare policies for pregnant women. Although there are some differences in the length of gestation, placental structure, and fetal development between humans and animal models, many structural and functional parallels exist,\textsuperscript{94–96} which serve as tractable platforms for evaluating the safety and efficacy of various therapies during pregnancy.
Similar to humans, pregnant mice, rats, and rabbits have a hemochorial placenta, and their relatively short gestation and large litters are advantageous for performing high throughput screening of candidate therapeutics for safety and efficacy. Preclinical behavioral testing of rodent offspring has proven to be a promising avenue for identifying and predicting adverse effects associated with prenatal drug exposure in children.97 Both rodent and rabbit models have been instrumental in testing teratogenic effects of artemisinin-based combination therapies for treating malaria in pregnant women. These studies concluded that drug-related teratogenic effects are limited to the first trimester, which supports the World Health Organization (WHO) recommendation that artemisinin may be administered only during the second or third trimester in pregnant women.98,99 One limitation of mouse and rat models, however, is their inability to recapitulate certain elements of human congenital disease. For instance, because murine CMV is not transmitted vertically as it is in humans, other animal models, including guinea pigs and nonhuman primates, are required for studying this aspect of disease pathogenesis. Studies in pregnant nonhuman primates have been instrumental for the identification of CD4 + T cell responses as critical for early control of CMV infection and transmission during pregnancy,100 and studies in guinea pigs have demonstrated that a single-cycle infectious CMV vaccine induces immune responses similar to natural infection and protects against congenital infection.101 Guinea pigs are also a useful model of chlamydial genital infection in humans. Experimental venereal infection with Chlamydia pneumoniae mimics disease associated with C. trachomatis in humans, including both sexual and perinatal transmission. Guinea pigs have therefore served as a useful model for testing candidate vaccines and treatments.102

Rabbits continue to serve as an important model of venereal infection with Treponema pallidum, the causative agent of syphilis, which is associated with congenital disease in humans. While natural infection in rabbits is associated with the species-specific T. paraluiscaneculi, rabbits can be experimentally inoculated with human T. pallidum, and have been instrumental in testing the efficacy of candidate vaccines.103 Many mammalian species, including rodents,104 ruminants,105 and nonhuman primates,40,106 are susceptible to infection with Listeria monocytogenes and demonstrate similar fetal complications when infection occurs during pregnancy. Studies in various animal models have uniquely contributed to our understanding of placental listeriosis and serve as a platform for evaluating prevention strategies.107,108 Finally, mice,85 cotton rats,86 guinea pigs,88 and sheep87 are all susceptible to infection with RSV, and vaccination of pregnant animals has facilitated the development and testing of maternal immunization strategies for protecting against neonatal RSV. Based on preliminary studies in guinea pigs,88 an experimental RSV recombinant F nanoparticle vaccine is now being evaluated in third-trimester pregnant women (Clinicaltrials.gov, NCT02247726).

Beyond the direct modeling of human congenital infection in animals, information can also be gained from the study of related pathogens in pregnant women, for which similar studies cannot be ethically performed.

VACCINE STRATEGIES FOR EMERGING INFECTIOUS DISEASES

Zika virus (ZIKV) is a unique Flavivirus that causes mild or subclinical disease in pregnant women,111,112 but can have devastating effects for the fetus and neonate. Infection during pregnancy is linked with spontaneous abortion and a variety of birth defects, including microcephaly and impaired neurocognitive function.113 Since its initial discovery in African macaques in 1947, ZIKV has expanded its geographical range and evolved into separate virus lineages, with environmental pressures resulting in the emergence of virulent substrains, raising concerns about vaccine escape mutants once a vaccine is approved.114 The combination of its unique pathogenesis, diverse modes of transmission, and rapid global spread has increased efforts toward development and licensing of a ZIKV vaccine.

A basic understanding of ZIKV biology and pathogenesis is essential for development of an effective vaccine against ZIKV. Although we can extrapolate some biological information from related Flaviruses, such as yellow fever, West Nile, and dengue viruses, ZIKV has unique characteristics following in vivo infection, which pose significant challenges to vaccine design. Unlike other Flaviruses, ZIKV has a tropism for reproductive tissues, including the testes, semen, and sperm in males,115,116 and the placenta in pregnant females,117−119 which is hypothesized to contribute to sexual and vertical modes of transmission, respectively. In addition to unique tissue tropisms, ZIKV persists in reproductive tissue following clearance of systemic viremia. In males, ZIKV RNA can be detected for months following recovery from symptomatic infection,120,121 and virus persistence in the placenta of pregnant women is hypothesized to contribute to prolonged viremia in this population.122 Evidence of virus persistence suggests that ZIKV may have evolved mechanisms for evasion of host immune responses when infection occurs in certain immune-privileged tissues. Scenarios involving persistent ZIKV infections should be considered in developing and testing candidate vaccines. Considering the potential for viral persistence in the semen, vaccinating men may serve as an additional strategy to reduce transmission to the fetus, as pregnant women may be infected by their sex partners.

Characterization of the immune response to ZIKV infection is essential for determining correlates of protection for vaccine efficacy. Following infection of both humans and nonhuman animals, ZIKV induces neutralizing antibodies against the ZIKV E protein, which prevent fetal infection and demise when administered to pregnant mice.123 Further, 26 MHC Class I epitopes have been identified that conferred protection against ZIKV challenge in immunocompetent mice.124 To date, 45 candidate ZIKV vaccines have been developed, and as of February 2017, nine have entered Phase I clinical trials.125 Vaccine candidates have been developed using diverse platforms, including DNA, mRNA, and purified inactivated and live-attenuated virus, many of which have been tested in non-pregnant mouse and nonhuman primate models for their ability to generate immune responses that mimic responses to natural infection and protected against ZIKV challenge. A candidate DNA plasmid vaccine induced robust cellular immunity and neutralizing antibody responses in both nonhuman primates and immunocompetent mice, and conferred complete protection against lethal ZIKV challenge in type I interferon receptor deficient (IFNAR −/−) mice.126 LNP-mRNA vaccines induced similar protective immunity, which was characterized by high neutralizing antibody titers and sterilizing immunity against ZIKV challenge in non-pregnant mice and nonhuman primates.127,128 Whether these candidate vaccines induce protective immunity in pregnant females that is sufficient to prevent fetal and neonatal infections129 requires further evaluation. Also, whether pre-existing immunity to other Flaviruses that co-circulate with ZIKV, including dengue virus and West Nile virus, affects the efficacy of ZIKV vaccines in pregnant
| Pathogen                      | Category          | Maternal susceptibility and/or disease severity | Vertical transmission and/or disease | Congenital Disease                                                                 | Neonatal susceptibility and/or disease severity | Prevention                                                                 | References |
|-------------------------------|-------------------|------------------------------------------------|-------------------------------------|-------------------------------------------------------------------------------------|-----------------------------------------------|----------------------------------------------------------------------------|------------|
| *Bordetella pertussis* bacteria | =                 | Postpartum                                     | Not documented                      |                                                                                    | ↑                                             | Maternal vaccination (Tdap) during second or third trimester pregnancy    | (73) (143) |
| *Chlamydia trachomatis* bacteria | =                 | Peripartum                                     | conjunctivitis                      |                                                                                    | ↑                                             | antibiotics                                                              | (144)      |
| *Clostridium tetani* bacteria | =                 | Peripartum                                     | Not documented                      |                                                                                    | ↑                                             | Maternal vaccination (Tdap) during second or third trimester pregnancy    | (81) (78) |
| **Group B Streptococcus** bacteria | =                 | Peripartum > Transplacental                    | Preterm birth, vision and hearing loss, mental retardation, cerebral palsy       |                                                                                    | ↑                                             | Intrapartum antibiotic prophylaxis                                        | (145) (146) |
| *Listeria monocytogenes* bacteria | ↑                 | Transplacental                                 | Miscarrage, perterm birth, sepsis, meningitis                                   |                                                                                    | ↑                                             | avoid high risk foods                                                     | (147) (39) |
| *Treponema pallidum* bacteria | =                 | Transplacental                                 | Miscarrage, preterm birth, stillbirth, bone deformities, anemia, hepatosplenomegaly, blindness, deafness, meningitis, skin rash | ↑                                                                                     | Intrapartum antibiotic prophylaxis            |                                                                             | (148) (149) |
| *Plasmodium spp.* protozoa ↑ | Transplacental     | Variable                                       |                                                                                    |                                                                                    | ↑                                             | artemisinin-based combination therapies                                   | (150) (151) |
| *Toxoplasma gondii* protozoa = | Transplacental     | stillbirth, chorioretinitis, deafness, microcephaly, encephalitis, developmental delay | ↑                                                                                     | Intrauterine blood transfusion                                                         |                                                                             | (153) (154) |
| *Cytomegalovirus* virus =     | Transplacental     | Sensorineural hearing loss, mental retardation, cerebral palsy, seizures, chorioretinitis | ↑                                                                                     | Vaccination (MLV) prior to pregnancy                                                   |                                                                             | (155) (156) |
| *Ebola virus* virus ↑         | Transplacental likely | Miscarrage or neonatal death most common        | ↑                                                                                     | Vaccination “at-risk” pregnant women                                                  |                                                                             | (158) (159) |
| *Hepatitis B virus* virus =   | Transplacental or Peripartum | chronic hepatitis                              | ↑                                                                                     | Vaccination (licensed in China only - not licensed during pregnancy)                  |                                                                             | (58) (60) |
| *Hepatitis E virus* virus ↑   | Transplacental or Peripartum | influence of transplacental transmission on congenital disease unknown | ↑                                                                                     | HAART, cesarean delivery, formula feeding                                              |                                                                             | (32) (3) (4) |
| *Herpes simplex virus* virus ↑ | Peripartum > Transplacental | spontaneous abortion, preterm birth            | ↑                                                                                     | No vaccine, antivirals not effective                                                  |                                                                             | (161) (162) |
| *Human immunodeficiency virus* virus ↑ | Transplacental or Peripartum | Not documented                                | ↑                                                                                     | HAART, cesarean delivery, formula feeding                                              |                                                                             | (163) (164) |
| *Influenza virus* virus ↑     | Transplacental     | Not documented                                 | ↑                                                                                     | Maternal vaccination (TIV) during any trimester                                       |                                                                             | (1) (9) (8) (13) |
| *Parvovirus B19* virus =      | Transplacental     | spontaneous abortion, hydrops fetalis, congenital anomalies, aplastic anemia (rare) | ↑                                                                                     | No vaccine; Intrauterine blood transfusion                                             |                                                                             | (165) |
| *Respiratory syncytial virus* virus ↑ | Postpartum | Not documented                                | ↑                                                                                     | No vaccine; immunoprophylaxis for at-risk neonates                                     |                                                                             | (84) (85) |
| *Rubella virus* virus =       | Transplacental     | Sensorineural hearing loss, ocular abnormalities, heart disease, mental retardation | ↑                                                                                     | Vaccination (MLV) prior to pregnancy                                                   |                                                                             | (166) (43) |
APPLICATION OF NOVEL VACCINE PLATFORMS

Conventional vaccines are formulated from either live, attenuated pathogen strains or from inactivated pathogens, but there are notable disadvantages to each of these platforms. Live, attenuated vaccines are replication-competent with the potential of becoming virulent and causing adverse effects in individuals with weakened immune systems. Due to the unknown risk to the developing fetus, live virus vaccines are not recommended for use in pregnant women. Inactivated vaccines, on the other hand, are not associated with a risk of reacquisition of virulence, but they tend to induce a weaker host immune response. Efforts to balance safety with immunogenicity have led to the development of several novel vaccine technologies, including replication-deficient nanoparticle-based vaccines and self-assembling recombinant virus-like particles (VLPs), replication-competent recombinant viral vectors, and single-cycle infectious viruses that can infect, but not replicate in host cells.

Nanoparticle delivery platforms, including liposomes and synthetic polymers, can be engineered to enhance selective tissue homing for high potency, targeted delivery of antigen in its native conformation. Recombinant VLPs combine highly immunogenic surface viral proteins with encapsulated adjuvants that are devoid of infectious nucleic acid, but induce strong cellular and humoral immune responses. Both nanoparticle and VLP vaccines are devoid of genetic material and are therefore replication incompetent, enhancing safety in vulnerable individuals. Although current production yields and costs associated with these technologies may prohibit large-scale use, these platforms may be well-suited for targeted use in high-risk populations, including pregnant women. Since the first nanoparticle-based vaccine was licensed for hepatitis B virus in 1981, the technology has been applied to develop licensed vaccines against human papilloma virus, hepatitis E virus, and malaria. Demonstration of safety and efficacy during pregnancy has not yet been documented.

Recombinant viruses can be engineered to combine the antigenic genes of one virus with the structural genes of another. This targeted manipulation of the virus genome is used to remove virulence genes to enhance safety and alter envelope proteins to change cell tropism. Recombinant viruses can be engineered to retain the ability to infect and replicate in the host, while preserving infectious potential and enhancing the generation of innate and adaptive immune responses that mimic natural infection. Recombinant virus vaccines, however, warrant careful consideration of the safety of the vector itself, especially in pregnant women. Once the safety and efficacy profiles of a viral vector platform have been established, engineering new antigenic targets into the viral genome are relatively simple, and do not require extensive re-validation, as safety and efficacy is most influenced by the vector virus. This can significantly reduce the time to develop and manufacture vaccines against new viral pathogens. Viruses from many different families can be used as vectors provided they can infect the host and elicit a productive immune response without causing disease. Of note, poxviruses are practical vectors due to ease of growth and manipulation in vitro, wide host range, and robust induction of protective immune responses. Although vaccinia virus vectors are contraindicated during pregnancy due to risk of disseminated disease, recent testing of a raccoonpoxvirus-vectorized rabies vaccine in pregnant mice proved safe and effective.

Other novel vaccine platforms are based on genetic engineering and creation of targeted loss-of-function mutations in the viral genome. Reverse genetics technology has contributed the identification of the sequences within the viral genomes that females should be considered in both preclinical animal models and human clinical trials.
are essential for infection and replication in vivo. It is now possible to engineer recombinant virus vaccines with targeted mutations in a cost- and time-efficient manner. In contrast to inactivated vaccines, reverse genetics allows for controlled manipulation of the viral genome that targets a specific process in the virus life cycle. This enables production of a vaccine strain with maximum efficacy and safety, and represents another promising alternative to conventional attenuated or killed virus vaccines. Among the safest recombinant vaccine approaches are the single-cycle infectious (sclAV) viruses, which have been developed from influenza A virus (IAV) backbones by deleting or truncating viral proteins necessary for completion of the virus life cycle within the host. Such genetic modifications render the virus replication-incompetent, but capable of infecting and inducing an immune response in the host. The infectious capacity of sclAV vaccines results in strong cellular and humoral immune responses, without the risks and adverse effects associated with live-attenuated vaccine strains. Studies in nonpregnant mice have demonstrated that a single dose of sclAV vaccine confers protection against heterosubtypic lethal challenge without any adverse effects. Similar safety and efficacy profiles of sclAV have been replicated in ferret and pig models, but studies in pregnant animals have not been performed. Compared with the risks associated with live virus vaccination and concerns of efficacy with inactivated vaccines, novel vaccine platforms, such as nanoparticle-based technologies, VLPs and replication-deficient viruses have proven benefits. Additional safety and efficacy studies in pregnancy models are warranted to validate and expand the use of these vaccine platforms for pregnant women. (Table 1)

CONCLUSIONS
Strategic immunization of women, either prior to or during pregnancy, can eliminate or substantially reduce the risk of maternal, fetal, and neonatal infection and disease. The effectiveness of an immunization protocol depends on both the efficacy of the vaccine in inducing protective immune responses and on the timing of vaccine delivery during pregnancy to synchronize the peak vaccine response with the period of greatest susceptibility in the host. Optimization of vaccination protocols to achieve this goal requires an understanding of the mechanisms of infection and pathogenesis of disease during pregnancy. Successful implementation of vaccine protocols for pregnant women requires consideration of additional challenges, such as the frequency of unplanned pregnancies and access to prenatal health care. Human surveillance data provide correlative clues of the character of specific infections, but mechanistic understanding requires additional study in comparative animal model systems.

ACKNOWLEDGEMENTS
The authors thank the Johns Hopkins Fisher Center Discovery Program for funding.

AUTHOR CONTRIBUTIONS
All authors (M.S.V. and S.L.K.) were involved in the conception and drafting of this review, and approved the manuscript before it was submitted by the corresponding author.

ADDITIONAL INFORMATION
Competing interests: The authors declare no competing financial interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

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