Preexisting antibodies can protect against congenital cytomegalovirus infection in monkeys

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Human cytomegalovirus (HCMV) is the most common congenital infection and a known cause of microcephaly, sensorineural hearing loss, and cognitive impairment among newborns worldwide. Natural maternal HCMV immunity reduces the incidence of congenital infection, but does not prevent the disease altogether. We employed a nonhuman primate model of congenital CMV infection to investigate the ability of preexisting antibodies to protect against placental CMV transmission in the setting of primary maternal infection and subsequent viremia, which is required for placental virus exposure. Pregnant, CD4+ T cell–depleted, rhesus CMV–seronegative (RhCMV-seronegative) rhesus monkeys were treated with either standardly produced hyperimmune globulin (HIG) from RhCMV-seropositive macaques or dose-optimized, potently RhCMV-neutralizing HIG prior to intravenous challenge with an RhCMV mixture. HIG passive infusion provided complete protection against fetal loss in both groups. The dose-optimized, RhCMV-neutralizing HIG additionally inhibited placental transmission of RhCMV and reduced viral replication and diversity. Our findings suggest that the presence of durable and potently neutralizing antibodies at the time of primary infection can prevent transmission of systemically replicating maternal RhCMV to the developing fetus, and therefore should be a primary target of vaccines to eliminate this neonatal infection.

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Preexisting antibodies can protect against congenital cytomegalovirus infection in monkeys

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Introduction

The current epidemic of Zika virus in the Americas has raised significant awareness of the societal and pediatric health impact of a congenitally transmitted neuropathogen. Yet nearly 1 million infants worldwide (1 out of every 150 live births) are born each year with congenital human cytomegalovirus (HCMV) infection (1, 2). Congenital HCMV, like congenital Zika virus, can cause microcephaly, hearing/vision loss, and abnormalities in nervous system development (1, 3). And though HCMV accounts for more congenital disease than all 29 newborn conditions currently screened for in the United States combined (4), public knowledge and effective interventions are severely lacking (5, 6). In recognition of this burden of disease, a congenital HCMV vaccine has remained a Tier 1 priority of the National Academy of Medicine for the past 17 years (7).

Drawing upon standard vaccine strategies, the HCMV field has previously attempted to induce potent antiviral immunity by various approaches including attenuation of live viruses (8–10), formulation of HCMV glycoprotein subunit vaccines (11–13), use of viral vectors for epitope expression (14–16), and delivery of single and bivalent DNA plasmid vaccines (17, 18). However, our understanding of anti-HCMV humoral immunity is complicated by the number of distinct glycoprotein complexes that contain neutralizing epitopes, most notably glycoprotein B (gB; fusion protein) and the pentameric complex gH/gL/UL128-131A (PC; necessary for entry into epithelial/endothelial cells) (19). While all vaccine platforms tested thus far ultimately failed...
to reach target endpoints in human clinical trials, the HCMV gB subunit vaccine demonstrated moderate (~50%) efficacy at preventing primary HCMV infection, which is promising for future vaccine-development efforts (12, 13). Additional retrospective human studies have reported that neutralizing antibodies targeting HCMV surface glycoproteins are correlated with reduced incidence of congenital virus transmission after primary maternal HCMV infection (20–22). However, a recent clinical trial failed to demonstrate a significant reduction in rates of congenital HCMV infection following passive infusion of hyperimmune globulin (HIG) to pregnant women with primary HCMV infection. Results from these past studies emphasize possible challenges for the development of an efficacious antibody-based vaccine (23).

One major barrier the HCMV vaccine field has faced is the lack of a highly translatable animal model of congenital virus transmission. For the past 50 years, vaccine efficacy studies that evaluate protection against congenital HCMV transmission have been reliant upon either (a) small-animal models using species-specific viruses with limited HCMV sequence homology or (b) costly and arduous human clinical trials. We recently reported on the first nonhuman primate (NHP) model of congenital CMV transmission following primary infection of rhesus monkey dams with rhesus CMV (RhCMV) (24). There are numerous logistical advantages to an NHP model including enhanced immunologic tools (e.g., monoclonal antibodies) and extensive knowledge of immune cell subsets (e.g., cellular markers, B/T cell receptor repertoire databases). Furthermore, we hypothesize that this NHP model may more accurately reflect HCMV pathogenesis than small-animal models owing to increased similarity of host and virus. The rhesus monkey placenta is anatomically, physiologically, and immunologically quite similar to that in humans, which likely enhances the translational applications of the model (25). Additionally, RhCMV is genetically more homologous to HCMV than guinea pig or murine CMV (24, 26, 27) and employs highly conserved immune-evasion pathways including viral IL-10 (28) and MHC downregulation (29).

Our previous study established that RhCMV could cross the placenta and cause congenital infection following intravenous (i.v.) inoculation of either immune-competent or CD4+ T cell–depleted seronegative dams during the second trimester of pregnancy. The i.v. route of RhCMV inoculation was selected for this experimental congenital infection model over mucosal routes of exposure, as it induced reproducible high levels of viremia and virus shedding in the urine and saliva of infected macaques, and hence best mirrored the prerequisite for systemic CMV replication prior to placental virus transmission in congenital CMV infection. In this study, we observed that CD4+ T cell–depleted dams frequently aborted their fetus following virus inoculation, exhibited higher plasma and amniotic fluid viral loads, and had delayed production of autologous neutralizing antibodies.

These results suggested that maternal humoral immunity may impact systemic and intrauterine RhCMV replication, thereby influencing the severity of congenital infection. As such, we sought to use this NHP model to investigate whether preexisting antibody alone, in the absence of virus-specific T cell responses, could protect against placental RhCMV transmission or reduce congenital RhCMV infection severity. The passive infusion of antibodies in this study was not intended to model a treatment strategy for the prevention of either maternal CMV acquisition or fetal virus transmission, but rather as a means to understand the protection conferred by preexisting maternal antibody. These insights will inform rational development of a vaccine to elicit the most critical immune responses required to eliminate congenital HCMV disease.

**Results**

**HIG production and study design.** Two separate preparations of HIG were purified from plasma of RhCMV-seropositive rhesus monkey donors. The first preparation (hereafter referred to as standard) was obtained from donors with high levels of total RhCMV-specific IgG, while the second preparation (hereafter referred to as high-potency) was purified from plasma of donors exhibiting robust epithelial cell–neutralizing IgG antibody titers (ID50 > 1:1,000). The high-potency preparation had approximately 4-fold more potent epithelial cell neutralization (standard = 22.6 μg/ml vs. high-potency = 5.9 μg/ml) and slightly increased overall glycoprotein binding (RhCMV gB/RhCMV PC [RhB/RhPC]: standard = 1.7/3.9 μg/ml vs. high-potency = 1.0/3.1 μg/ml) versus the standard preparation. All RhCMV-seronegative dams in our current and previous study were infused with a CD4+ T cell–depleting antibody at week 7 of gestation, as previously described (24). T cell phenotyping (Supplemental Figure 1 and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.94002DS1) revealed a decline in CD4+ T cell–depleting antibody at week 7 of gestation, as previously described (24). T cell phenotyping (Supplemental Figure 1 and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.94002DS1) revealed a decline in CD4+ T cells with changes in the CD8+ T cell population in response to primary RhCMV infection (Supplemental Figure 2). The dams were subsequently divided into 3 groups: control,
standard, and high-potency. Each animal, including those in the control group \((n = 6, \text{ including 4 historical controls; see ref. 24})\) were i.v. inoculated with a mixture of fibroblast-tropic 180.92 (30) and epithelial cell–tropic UCD52/UCD59 (31) RhCMV viruses at week 8 of gestation (Figure 1A). The standard group \((n = 3)\) was administered a single dose (100 mg/kg, which mimics a previously published human trial protocol; see ref. 23) of the standard hyperimmune globulin (HIG) preparation 1 hour prior to RhCMV infection (Figure 1B), while the high-potency group \((n = 3)\) was given a dose-optimized regimen of the high-potency HIG preparation (Supplemental Figure 3) 1 hour prior to (150 mg/kg) and 3 days following (100 mg/kg) RhCMV inoculation (Figure 1C and Supplemental Figure 4).

**Glycoprotein targets of RhCMV-neutralizing antibodies.** To determine the RhCMV antibody specificity of the administered HIG, both standard (Figure 2A) and high-potency HIG (Figure 2B) were depleted for antibodies specific for RhgB, RhPC, as well as RhgB/RhPC combined. Sufficient and selective depletion of glycoprotein-specific antibodies was confirmed by ELISA against the depleted epitope. We confirmed a greater than 75% decrease in EC\(_{50}\) magnitude against depleted epitope and less than 20% change in EC\(_{50}\) against nondepleted epitope (Table 1). For the standard HIG preparation, adsorption with RhgB resulted in a 2.2-fold reduction in the neutralization potency of plasma (mock-depleted IC\(_{50}\) = 32.3 vs. RhgB-depleted IC\(_{50}\) = 71.8), whereas adsorption with RhPC resulted in a slightly higher 2.9-fold reduction in neutralization potency (RhPC-depleted IC\(_{50}\) = 94.6) (Figure 2A and Table 1). This same trend was observed for antigen depletion of the high-potency HIG, with a 2.9-fold reduction following RhgB depletion and a 3.3-fold reduction following RhPC depletion (Figure 2B and Table 1). Additionally, depletion of antibodies specific for both antigens reduced the neutralization potency of sera beyond that of depletion with either RhgB or RhPC alone (Figure 2, A and B, and Table 1). Altogether, these findings suggest that antibodies targeting both RhgB and RhPC contribute to overall RhCMV neutralization activity in roughly equivalent proportions. Furthermore, both RhgB- and RhPC-specific antibodies were found to contribute to the RhCMV-neutralization titer in individual RhCMV-seropositive rhesus monkeys (Supplemental Figure 5).
Antibody kinetics following HIG infusion. Total RhCMV-binding IgM and IgG titers (Figure 3, A and B), RhgB- and RhPC-specific IgG titers (Figure 3, C and D), and fibroblast/epithelial cell neutralizing antibody titers (Figure 3, E and F), were measured following HIG infusion. Based on the known kinetics of antibody responses to primary viral infections, we attributed early detection of RhCMV-specific activity (within the first 10 days) to reflect passively infused antibodies in the HIG preparations. Peak RhCMV-binding IgG as well as RhgB- and RhPC-specific antibody titers in the high-potency HIG group surpassed those of the standard HIG group (Figure 3, B–D), while peak neutralization titers were similar between the 2 HIG groups (Figure 3, E and F). However, levels of IgG binding and antibody neutralization were sustained near peak levels for 10 days in the dose-optimized, high-potency HIG group compared with only 3 days in the standard HIG group (Figure 3, B–E). It is noteworthy that average binding-IgG and epithelial cell–neutralization titers in the high-potency HIG group surpassed those of chronically infected rhesus monkeys (Figure 3, B and E, and Supplemental Figure 5) (24).

The standard HIG group had the most robust natural host immune response, with a sustained elevation of RhCMV-specific IgM (Figure 3A), as well as an exponential rise in binding-IgG (Figure 3B) and neutralization titers (Figure 3, D and E) that outpaced each of the other groups.

RhCMV viral load, placental transmission, and shedding. Following RhCMV inoculation, both control and standard HIG group animals showed a rapid onset of viremia that peaked at 2 weeks after infection (Figure 4A). In contrast, peak viremia in the high-potency HIG group was delayed until 4 weeks after infection (Figure 4A), with a median value nearly 2 logs lower than the control group (Figure 4E; P = 0.047, corrected Wilcoxon exact test). RhCMV DNA copy number in amniotic fluid was used as a marker for placental virus transmission because it is the gold-standard clinical test for congenital HCMV infection (32). All 6 control animals had detectable RhCMV DNA in amniotic fluid between 1 and 3 weeks following infection (Figure 4B), indicating 100% placental transmission. Additionally, 2 of 3 animals in the standard HIG group had detectable congenital RhCMV infection (67% transmission) (Figure 4B). However, none of the 3 animals in the high-potency HIG group had detectable RhCMV DNA in the amniotic fluid, indicating complete inhibition of congenital RhCMV transmission (Figure 4B), and indicating a potential dose effect of HIG-mediated protection. It is possible that systemic viral load influenced placental transmission, as animals that transmitted the virus had the highest peak plasma viral loads (Figure 4E) and peak plasma viral load correlated with initial amniotic fluid viral load (Figure 4F; r = 0.812, P = 0.001, Spearman correlation). Finally, while there was no difference in the peak magnitude of viral shedding in maternal urine

| Table 1. Antigen-binding and neutralization activity of RhgB- and RhPC-depleted plasma |
|-------------------|-------------------|-------------------|-------------------|-------------------|
|                    | Standard HIG          |                    | High-potency HIG       |                    |
|                    | RhgB EC<sub>50</sub> (μg/ml)<sup>a</sup> | RhPC EC<sub>50</sub> (μg/ml)<sup>a</sup> | IC<sub>50</sub> (μg/ml) | RhgB EC<sub>50</sub> (μg/ml)<sup>a</sup> | RhPC EC<sub>50</sub> (μg/ml)<sup>a</sup> | IC<sub>50</sub> (μg/ml) |
| Undepleted         | 1.7                 | 3.9                 | 22.6               | 1.0               | 3.1                 | 5.9                 |
| Mock-depleted      | 5.9                 | 15.7                | 32.3               | 3.1               | 9.0                 | 13.1                |
| RhgB-depleted      | >33.3<sup>a</sup>    | 17.7                | 71.8               | 10.9              | 7.1                 | 38.3                |
| RhPC-depleted      | 5.26                | >33.3<sup>a</sup>   | 94.6               | 3.8               | >33.3<sup>a</sup>   | 42.6                |
| Both depleted      | >33.3<sup>a</sup>    | >33.3<sup>a</sup>   | >100<sup>c</sup>   | 15.3              | >33.3<sup>a</sup>   | 82.5                |

<sup>a</sup>EC<sub>50</sub> values measured in μg/ml of HIG equivalent. <sup>b</sup>Starting ELISA concentration of 33.3 μg/ml. <sup>c</sup>Starting neutralization concentration of 100 μg/ml. HIG, hyperimmune globulin; RhgB, rhesus CMV glycoprotein B; RhPC, rhesus CMV pentameric complex.
between treatment groups, there was delayed onset of shedding in both urine (Figure 4G) and saliva (Figure 4H) following high-potency HIG infusion (both $P = 0.047$, corrected Wilcoxon exact test). Altogether, these findings suggest that the presence of neutralizing antibodies can reduce systemic replication and block dissemination of the virus to other anatomic compartments.

Short NGS amplicon population profiling (SNAPP) of plasma virus. Using a validated (Supplemental Figure 6) next-generation sequencing (NGS) technique, short 400-bp regions within the RhCMV gB (Figure 5) and gL (Supplemental Figure 7) genetic loci were amplified and sequenced at great read depth. We found that, as previously reported (24), the viral populations in plasma were composed primarily of strain UCD52 at each sampled time point and in all treatment groups. The reason for the consistent predominance of UCD52 over other viral strains 180.92/UCD59 is unclear, though may indicate enhanced in vivo fitness of this isolate. Additionally, viral diversity was measured by calculating mean nucleotide diversity ($\pi$) across each sample’s haplotype sequences (Supplemental Table 2). Overall, the diversity of the dominant plasma UCD52 viral subpopulation was restricted in the high-potency HIG–treated animals compared with the control group at both the gB (Figure 5C) and gL (Supplemental Figure 7) loci (both $P = 0.036$, Wilcoxon exact test). Nucleotide diversity was not significantly different between the control and standard HIG groups.

Fetal outcome and RhCMV congenital infection. While 5 of 6 control dams aborted their fetus at 3 weeks after RhCMV infection, administration of either HIG regimen was sufficient to protect fetuses from spontaneous abortion ($P = 0.015$, exact log-rank test based on Heinze macro; Figure 6A) (33). Furthermore, we observed a reduced rate of congenital RhCMV transmission among HIG-treated animals (2 of 6) compared with control animals (6 of 6) ($P = 0.049$, exact log-rank test based on Heinze macro; Figure 6B) (33). Intriguingly, in the standard HIG group only 1 of 3 fetuses remained uninfected, whereas all 3 fetuses in the high-potency HIG group were protected from infection (Figure 6C), suggesting that preexisting, potently CMV-neutralizing HIG can reduce the incidence of congenital infection. Of note, fetal growth curves (Supplemental Figure 8) and systemic cytokine profiles (Supplemental Figure 9) were similar between treatment

(Figure 4C) or saliva (Figure 4D) between treatment groups, there was delayed onset of shedding in both urine (Figure 4G) and saliva (Figure 4H) following high-potency HIG infusion (both $P = 0.047$, corrected Wilcoxon exact test). Altogether, these findings suggest that the presence of neutralizing antibodies can reduce systemic replication and block dissemination of the virus to other anatomic compartments.

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groups, indicating no obvious off-target effects of the infused HIG. Reassuringly, all dams with detectable RhCMV DNA in amniotic fluid also had detectable virus in placenta, amniotic membrane, and umbilical cord (Figure 6C). However, there was significant heterogeneity in viral burden and distribution among fetal tissue types; some fetuses had high copy number virus in nearly every tissue tested (174-97, IM67), while others only had virus detectable in the cochlea (GI73, HD79) (Figure 6C). This heterogeneity is consistent with previous reports for HCMV suggesting that the virus can replicate in the amniotic membrane without observable fetal infection (34). It is unclear from the data whether cochlear infection might have occurred via direct inoculation of the ear canal or if fetal immune control could have led to isolated virus detectable in the cochlea. However, the persistence of virus in the cochlea further validates this model as consistent with congenital HCMV infection in human fetuses. Congenital infection was additionally confirmed in placentas by histology and immunohistochemical staining for the RhCMV IE-1 protein (Figure 6D and Supplemental Table 3). Of note, the placental tissue PCR and histology/IHC data do not agree completely (Supplemental Table 3), suggesting that PCR may be a more sensitive measure after resolution of the acute infection.

**Placental transcriptome.** Congenital RhCMV infection appears to modify the placental transcriptome more radically than HIG infusion or fetal abortion, as more than 300 genes were differentially expressed between RhCMV-transmitting and nontransmitting dams, but only 78 when comparing HIG infusion/fetal outcome.
This finding suggests that HIG predominantly mediated protection through its impact on maternal systemic virus replication rather than through modification of the placental transcriptome. A large proportion of these differentially expressed genes were immune related. Additional cellular functions that appeared altered by RhCMV infection included metabolism, oxidoreductase activity, and cellular growth/proliferation (Supplemental Table 4). Gene interaction analysis of differentially expressed genes between RhCMV-transmitting and nontransmitting dams (Figure 7B) revealed that the placental transcriptome was heavily biased towards upregulation of genes in transmission. Furthermore, certain gene nodes had numerous connections to other differentially regulated genes (e.g., VCAM1, EGFR, GZMB, LYN, FYN, and GAPDH), suggesting that these keystone genes are central to the cellular pathways induced by RhCMV infection. Focusing on a subset of genes involved in innate versus adaptive immunity (Figure 7C), we note that there is a potential bias for activation of innate immune pathways by placental RhCMV infection. Finally, cell-type enrichment analysis based upon previously described genes involved in cellular activation/processes (35) demonstrated no preference at the transcriptome level for myeloid versus lymphoid lineages (Figure 7D). However, NK cell–specific genes had the most robust differential expression in RhCMV infection (Figure 7D),
and we identified 21 unique genes known to be associated with NK cell function and/or chemotaxis (Figure 7E and Supplemental Table 5) including killer immunoglobulin-like receptors (KIR2DL4) and killer lectin-like receptors (KLRD1, KLRC1, KLRC3, and KLRB1). Furthermore, nonclassical MHC molecules HLA-E and HLA-G (rhesus analog inferred by IPA database) were upregulated.

**Discussion**

The role of preexisting antibodies in prevention of congenital HCMV infection remains controversial within the HCMV vaccine field, yet our findings indicate that a dose-optimized preinfusion with potently neutralizing HIG can block placental RhCMV transmission in a rhesus monkey model of congenital CMV. Our study illustrates that factors such as maternal antibody titer and neutralization potency at the time of peak CMV viremia are important considerations for antibody-mediated protection against congenital CMV infection. In this study, given the experimental setup, we have not definitively proven that
it is neutralization titer that is protective against congenital RhCMV transmission. Other possibilities include the total preexisting binding-antibody level, non-neutralizing antibody functions such as antibody-dependent cellular cytotoxicity (ADCC) activity, a reduction in maternal viral load, or reduction in the number of infected monocytes that might traffic to the placenta. Nevertheless, these data substantiate observational human cohort studies suggesting that maternal HCMV-specific antibodies decrease risk of placental virus transmission (20–22, 36). Furthermore, these data affirm findings from the guinea pig model of congenital infection that demonstrate that preexisting antibodies (from either glycoprotein immunization or passive HIG/monoclonal antibody infusion) can reduce the incidence and/or severity of congenital guinea pig CMV infection (37–40).

The therapeutic efficacy of HIG in preventing placental transmission in mothers with primary HCMV infection is a topic of ongoing investigation. Several nonrandomized small-scale clinical trials (41, 42) and case-controlled studies (43, 44) concluded that postinfection HIG infusion significantly reduced rates of placental HCMV transmission, reduced placental pathology, and improved infant outcomes. Animal studies in the guinea pig model of congenital infection further corroborated these findings (39, 45). However, a large-scale, randomized, placebo-controlled trial did not find significant reduction in rates of congenital infection in mothers with primary HCMV infection that were subsequently treated with HIG, and revealed no difference in virus-specific antibody titers, effector T cell count, or the level of plasma viremia.
between the HIG-treated and placebo groups (23). Several confounding factors may explain the lack of efficacy of HIG treatment in this large-scale clinical trial. First, it was unknown in this study whether congenital transmission had already occurred prior to initiating therapeutic HIG treatment. Second, there is no concrete evidence for the therapeutic efficacy of the clinically utilized product (Cytotect), and therefore it is unknown whether HIG infusion achieved a titer that was both sustained and/or sufficient for effective anti-HCMV immunity.

This study was not intended to model a therapeutic treatment regimen but rather to directly address the clinical question, can preexisting antibodies alone prevent congenital CMV infection? The advantages of our NHP model are that the timing of infection is predefined and serum antibody titer can be titrated. By utilizing passive infusion experiments with different dosing regimens and potency of RhCMV-specific antibodies prior to RhCMV inoculation, we were able to model the requirements for preexisting antibodies to confer protection against congenital RhCMV infection. Moreover, conducting the experiments in a CD4+ T cell–depletion model also allowed us to address the protective role of antibodies to the exclusion of RhCMV-specific cellular immune responses. Our results suggest that a single 100 mg/kg dose of standard neutralizing potency HIG (donor plasma screened by elevated ELISA OD, as is done for the clinical product Cytotect; see ref. 46) delivered immediately prior to RhCMV infection can reduce the severity of congenital infection (0 of 3 monkeys aborted) but is insufficient to prevent congenital CMV transmission (2 of 3 monkeys transmitted).

To our knowledge, the epitope specificity of RhCMV-neutralizing antibodies has not been previously reported. It has been well established for HCMV that, while gB is an important target of neutralizing antibodies, the majority of neutralization activity in seropositive individuals is directed against the gH/gL/UL128-131A pentameric complex (47, 48). Furthermore, the pentameric complex has also been shown to be a target of neutralizing antibodies for guinea pig CMV (49). In this study, we determined that antibodies targeting both RhgB and RhPC contribute to the total neutralization activity of RhCMV-seropositive monkey sera in approximately equivalent proportions (Figure 2). It is certainly reassuring for the translational applicability of this animal model system that RhPC-directed antibodies are neutralizing; however, it is unclear why the relative proportions of PC-directed neutralization activity would differ between HCMV (~85%) (47) and RhCMV (~50%). We hypothesize that this could explain why the magnitude of RhCMV-neutralizing antibody titers are roughly equivalent in magnitude when measured in both fibroblasts and epithelial cells (Figure 3), whereas HCMV-neutralizing antibody titers are nearly an order of magnitude lower in fibroblasts than epithelial cells (50). Subsequent studies are needed to improve our understanding of this potential difference in viral biology and/or antiviral humoral immune factors between RhCMV and HCMV.

It has been hypothesized that passively infused antibodies can reduce systemic maternal viral load and thereby reduce the likelihood of placental transmission (20, 41), though this correlation has not been observed in some human primary infection cohorts (51). In this study, dams with reduced peak plasma viral load in the presence of antibodies did not transmit the infection to their fetus in utero (Figure 4E). This trend was accompanied by a significant reduction in nucleotide diversity of the plasma virus population in the high-potency HIG group, assessed at both the gB and gL loci (Figure 5 and Supplemental Figure 7). Thus, we hypothesize that in the setting of antibody-mediated immune pressure, the number of unique viral variants is restricted following primary infection and systemic replication is inhibited, resulting in decreased viral transmission to the fetus.

The placental transcriptome was examined to investigate whether HIG-mediated protection against congenital infection occurs via gene regulation at the level of the placenta/decidua. Intriguingly, RhCMV infection impacted the transcriptional profile of placental decidua cells to a much greater degree than HIG treatment or fetal abortion, suggesting that HIG predominantly mediated protection through its impact on maternal systemic virus replication. The preferential activation of innate over adaptive immune processes in RhCMV-infected placenta (Figure 7D) is perhaps due to the tolerogenic environment at the maternofetal interface (52, 53). Furthermore, the apparent upregulation of NK cell–specific genes in RhCMV-infected placenta (Figure 7D and E) may be attributable to either NK cells being the major leukocyte in the decidua during the early stages of pregnancy (54) or to the propensity for NK cells to control CMV infection in vivo (55, 56). However, this analysis has several caveats including (a) placental samples were from full-thickness decidua/placenta tissue and therefore lack anatomic specificity and (b) all tissue samples were isolated from CD4+ T cell–depleted animals. As such, we cannot be certain whether the trends observed are broadly applicable outside of this model system.
Our findings are based on a relatively small cohort of animals ($n = 12$) due to the severely limited availability of breeding, seronegative female monkeys. One consequence of our limited animal population is the need to amplify the biologic effects of placental RhCMV infection through both CD4$^+$ T cell depletion and i.v. viral inoculation. Though both these artificial methods distort the reality of natural CMV infection and bias our results in favor of placental transmission, we propose that these interventions strengthen the major finding in this study that preexisting antibody alone (in the absence of cell-mediated immunity) can prevent placental transmission in the setting of acute maternal viremia. One additional limitation to our study design is that, although HIG is known to nonspecifically modulate immune responses to prevent inflammatory disease (44, 57, 58), we had to limit the number of individual control groups in this study and did not have a cohort of animals infused with non–RhCMV-reactive HIG. Because of this deficiency, we cannot definitively rule out nonspecific immunomodulatory effects of the infused HIG, though all infusion groups had similar cytokine expression profiles following infection (Supplemental Figure 9). Furthermore, since protection against congenital transmission may have been achieved as a result of a dose effect in the high-potency HIG group versus the standard group, antibody-mediated protection is likely due to anti-viral function rather than immunomodulatory properties of the infused HIG.

Despite these limitations, this is to our knowledge the first primate study to demonstrate that preexisting maternal antibody alone has the potential to block and ameliorate placental CMV transmission. Thus, a maternal vaccine that elicits durable, potently neutralizing humoral immunity could be an ideal intervention to eliminate pediatric disabilities associated with congenital HCMV infection. Based on our findings we hypothesize that the prevention of maternal HCMV acquisition (i.e., sterilizing immunity), which has been the target of several phase II vaccine trials (12, 13), may not be the most appropriate measurable outcome for a congenital HCMV vaccine that ultimately seeks a reduction in the frequency of congenital HCMV transmission. Since we observed that antibody-mediated protection against congenital HCMV was associated with a decrease in maternal systemic viral load and constrained viral diversity, these surrogate markers could potentially serve as clinical study endpoints. Subsequent studies should investigate the epitope specificity and properties of protective immunoglobulins for future rational design of a maternal congenital HCMV vaccine.

Methods
Animals. Indian-origin rhesus macaques were housed at the Tulane Primate National Research Center and maintained in accordance with institutional and federal guidelines for the care and use of laboratory animals (59). All females were from the expanded specific pathogen–free (eSPF) colony and confirmed to be RhCMV seronegative by whole-virion ELISA screening for RhCMV-specific IgM and IgG plasma antibodies. RhCMV-seronegative males and females were then cohoused and females were screened every 3 weeks for pregnancy via abdominal ultrasound. All animals in this study were between 4 and 9 years of age at the time of pregnancy/enrollment (JC65 = 4.7, GI73 = 8.8, IM67 = 5.7, JM52 = 4.6, HR73 = 7.6, HD82 = 8.7, HD79 = 8.7, GM04 = 8.8, and HD79 = 8.7 years old). Gestational dating was performed by sonography, and was based on gestational sac size (average of 3 dimensions) and crown-rump length of the fetus. At week 7 of gestation, 12 RhCMV-seronegative dams were administered an i.v. 50 mg/kg dose of recombinant rhesus CD4$^+$ T cell–depleting antibody (CD4R1 clone; NIH Nonhuman Primate Reagent Resource).

One week following CD4$^+$ T cell depletion, 3 seronegative dams were administered a single dose (100 mg/kg) of HIG produced from the plasma of RhCMV-seropositive monkeys screened by RhCMV whole-virion IgG-binding ELISA (IC$_{50}$ = 22.64 μg/ml) 1 hour prior to i.v. challenge with a mixture of RhCMV strains 180.92 (originally isolated from the lung of an immune-compromised rhesus macaque) (30) and UCD52/59 (originally isolated from urine of seropositive monkeys) (31). Each monkey was administered $2 \times 10^8$ 50% tissue culture infective dose (TCID$_{50}$) RhCMV 180.92 in one arm and $1 \times 10^8$ TCID$_{50}$ each of RhCMV UCD52 and UCD59 in the other arm, diluted in serum-free RPMI. The remaining 3 dams received a dose-optimized (2 dose) regimen of a potently neutralizing HIG product purified from RhCMV-seropositive monkeys with a plasma neutralization titer in epithelial cells greater than 1,000 (IC$_{50}$ = 5.88 μg/ml) at 1 hour prior to i.v. RhCMV inoculation (150 mg/kg) and 3 days after infection (100 mg/kg), respectively. This dose-optimized regimen was based on pharmacokinetic modeling of the neutralization activity from the standard group (Supplemental Figure 4). Six dams were noninfused control animals (including 4 historical controls) (24).
Blood draws were performed on the standard potency–infused animals on the following days after infection/infusion: 0, 2, 4, 7, 10, 14, 21, 28, 35, and 42. The high-potency–infused animals were sampled at days 0, 1, 2, 3, 4, 7, 10, 14, 21, 28, 35, 42, 49, 56, 63, 70, 77, and 84. Amniotic fluid (by amniocentesis), urine (by clean pan collection), and saliva (by oral saline wash), were also collected on all animals on a weekly to biweekly intervals until fetal harvest. In the case of fetal abortion, placental and fetal products were obtained from the cage and fixed in formalin for H&E staining and immunohistochemistry or frozen. Fetal growth was monitored by measuring biparietal diameter and femur length during weekly to biweekly fetal sonography over the course of gestation. Sonography was also used to screen for signs of congenital RhCMV-associated sequelae. The fetuses of dams in the standard HIG group were harvested at 6 weeks following infection, and those of the high-potency HIG group at 12 weeks following infection. The standard and high-potency HIG groups were followed for different periods of time because we did not observe any fetal abortion in the first group (standard HIG), and therefore wanted to ensure in the subsequent year that HIG treatment did not simply delay fetal abortion.

**IgG purification for HIG preparations.** Seropositive rhesus monkey plasma donors were screened by either high-OD binding against UCD52 virions (top 50% seropositive OD; standard HIG) or by epithelial cell–neutralization titer (ID$_{50} >$ 1,000; high-potency HIG). Donor plasma was mixed with an equal volume of Protein G IgG Binding Buffer (Pierce) and spun at 5,000 g for 25 minutes at room temperature. Pellets were discarded and supernatants were filter sterilized through a 0.22-μm filter and used for purification. The sera mixture was passed through a gravity column packed with Protein G Plus Agarose beads (Pierce). Beads were washed with 5 column volumes (CVs) of IgG Binding Buffer and eluted with 2 CVs of IgG Elution Buffer. Eluted IgG was immediately neutralized with 1 M Tris buffer, pH 8. Beads were washed with 5 CVs of IgG Binding Buffer. Serum was passed through Protein G Plus Agarose beads 10 times and IgG was eluted each time. All IgG elutions were concentrated and buffer exchanged into 1× PBS using Millipore Amicon 30K filters (Fisher Scientific). Final preparations were tested by ELISA for the presence of bacterial endotoxin. See quality control sheet for high-potency HIG preparation in Supplemental Figure 3.

**Pharmacokinetic modeling for dose-optimized HIG infusion.** Plasma concentration–time data of IgG in rhesus monkeys were adjusted using the baseline value of IgG for each individual animal and analyzed using WinNonlin (version 6.3.0; Certara). One- and 2-compartment PK models with first-order elimination were evaluated to characterize the observed data. PK parameters including clearance (CL), volume of distribution (V), and half-life associated with the elimination phase ($T_{1/2}$) were estimated for each animal. Simulations for optimizing dosing regimen in the second PK study were performed to ensure IgG concentrations above 1–1.2 mg/ml (~ID$_{50} >$ 1,000) for 2 weeks after initiating treatment.

**Cell culture and virus growth.** Telomerized rhesus fibroblasts (TeloRFs) were maintained in Dulbecco’s modified Eagle medium (DMEM) containing 10% FCS, 2 mM L-glutamine, 50 U/ml penicillin, 50 μg/ml each streptomycin and gentamicin, and 100 μg/ml geneticin G418 (Invitrogen). Immortalized TeloRFs were used for in vitro assays and viral growth since they (a) robustly support viral replication indefinitely (i.e., without issues of cellular senescence) and (b) facilitate standardization/consistency between assays (60). Monkey kidney epithelial (MKE) cells were cultured in DMEM-F12 supplemented with 10% FCS, 2 mM L-glutamine, 1 mM sodium pyruvate, 50 U/ml penicillin, 50 μg/ml each streptomycin and gentamicin, and 5 ml of Epithelial Cell Growth Supplement (ScienCell). Cell lines were tested for mycoplasma contamination every 6 months.

Epithelial cell–tropic UCD52 and UCD59 were propagated on MKE cells for 3–4 passages, and then passaged once on TeloRFs in order to increase viral titer. This single round of fibroblast amplification was intended to minimize any alteration of viral tropism or glycoprotein surface expression. Virus infections were performed with similar media but with 5% FCS for 4 hours at 37°C and 5% CO$_2$. Virus was harvested when cells showed 90% cytopathic effects (CPEs) by cell scraping. For harvest, infected cells were pelleted by low-speed centrifugation, and the supernatant was placed on ice. Cell pellets were then resuspended in infection media and subjected to 3 freeze/thaw cycles. Following centrifugation, supernatants were combined, passed through a 0.45-μm filter, overlaid onto a 20% sucrose cushion, and ultracentrifuged at 70,000 g for 2 hours at 4°C using an SW28 Beckman Coulter rotor. Virus pellets were resuspended in DMEM containing 10% FCS and titered in TeloRFs using the TCID$_{50}$ method of Reed and Muench.

**Characterization of T cells by flow cytometry.** Maternal peripheral CD4$^+$ and CD8$^+$ T cells were phenotyped by mixing 100 μl of EDTA-anticoagulated blood with a pool of fluorescently labeled monoclonal antibodies including CCR7, CD95, CD28, CD4, CD20, CD3, CD8, and CD45 (Supplemental Table 1).
After a 30-minute incubation at 4°C, cells were washed with PBS supplemented with 2% FCS and pelleted at 100 g for 5 minutes. Red blood cells were then lysed by adding 1× BD lysis buffer for 40 minutes at room temperature. Intact cells were centrifuged at 150 g for 5 minutes, washed once with PBS containing 2% FCS, and resuspended in 100 μl of PBS with 2% FCS. Cells were fixed by adding 15 μl of 2% formaldehyde and processed by flow cytometry. Gating of CD4+ and CD8+ populations/subpopulations was completed in FlowJo as is outlined in Supplemental Figure 1.

**Antibody kinetics and glycoprotein binding specificity.** Virus-specific IgM and IgG antibody kinetics were measured in maternal plasma by whole-virion ELISAs, and glycoprotein specificity of HIG and maternal plasma samples was measured by ELISA against soluble RhCMV gB protein and RhPC. Plates were coated with antigen diluted in PBS with Mg2+ and Ca2+ (PBS+) overnight at 4°C — either 5,120 PFU/ml of filtered RhCMV UCD52 virus for whole-virion ELISA or 0.13 μg/ml soluble glycoprotein. Following incubation, plates were blocked for 2 hours with blocking solution (PBS+, 4% whey protein, 15% goat serum, 0.5% Tween-20), and then 3-fold serial dilutions of plasma (1:30 to 1:5,314,410) were added to the wells in duplicate for 1.5 hours. Plates were then washed twice and incubated for 1 hour with anti-monkey IgM or anti-monkey IgG HRP-conjugated antibodies (Rockland) at a 1:10,000 dilution. After 4 washes, SureBlue Reserve TMB Microwell Peroxidase Substrate (KPL) was added to the wells for 10 minutes, and the reaction was stopped by addition of 1% HCl solution. Plates were read at 450 nm. The lower threshold for antibody reactivity was considered to be 3 SDs above the average OD measured for the preinfection, RhCMV-seronegative samples at the starting plasma dilution (1:30). The end-point dilution reported is the highest dilution of plasma that exceeds this positivity threshold.

**Neutralization assays.** TeloRFs and MKE cells were seeded into 96-well plates and incubated for 2 days at 37°C and 5% CO2 to achieve 100% confluence. After 2 days, serial dilutions (1:10 to 1:30,000) of heat-inactivated rhesus plasma were incubated with RhCMV 180.92 or RhCMV UCD52 (MOI = 1) in 50 μl for 45 minutes at 37°C. The virus/plasma dilutions were then added in duplicate to wells containing TeloRFs or MKE cells, respectively, and incubated at 37°C for 2 hours. After washing, cells were incubated at 37°C for an additional 16 hours. Infected cells were then fixed for 20 minutes at −20°C with 1:1 methanol/acetone, rehydrated in PBS+ 3 times 5 minutes, and processed for immunofluorescence with 1 μg/ml mouse anti–RhCMV IE-1 monoclonal antibody (provided by Daniel Cawley, Oregon Health and Science University, Portland, Oregon, USA) followed by a 1:5,314,410 dilution of goat anti–mouse IgG–Alexa Fluor 488 antibody (Millipore). Nuclei were stained with DAPI for 5 minutes (Pierce). Infection was quantified in flow cytometry. Gating of CD4+ and CD8+ populations/subpopulations was completed using the method of Reed and Muench.

**Soluble protein bead coupling and antibody depletion.** Cyanogen bromide–activated (CNBr-activated) Sepharose beads (GE Healthcare) were rehydrated in 1 mM HCl, and then suspended in coupling buffer (0.1 M NaHCO3 + 0.5 M NaCl, pH 8.3) at room temperature. Every 100 μl of bead slurry was combined with 150 g soluble RhgB or RhPC ligands. Coupling was allowed to proceed for 1 hour at room temperature on an inversion rotator. Excess soluble protein was washed off with 5 column volumes of coupling buffer. CNBr unbound active groups were blocked by incubation in 0.1 M Tris HCl, pH 8.0 for 2 hours at room temperature on an inversion rotator. Protein-bound beads were washed with 3 cycles of alternating pH: first 0.1 M acetic acid, pH 4.0, followed by 0.1 M Tris-HCl, pH 8.0. Beads were suspended in PBS and stored at 4°C. For depletion of antibodies for a given specificity, 100 μl of protein-coupled bead slurry was loaded into a spin microelution column (Pierce, TFS). Plasma (50 μl) was filtered (0.22 μm) and then loaded into the column. Plasma was centrifuged through the column 5 times without elution between steps. Bound IgG was eluted using 0.2 M glycine elution buffer, pH 2.5. The column was recalibrated by washing with 3 cycles of solutions of alternating pH (as described above). Adequate and specific depletion of plasma was confirmed by ELISA against the depleted protein (>75% change in EC50 against depleted epitope; <20% change in EC50 against nondepleted epitope).

**Tissue processing and staining.** Standard immunoperoxidase staining for RhCMV was performed on formalin-fixed, paraffin-embedded sections of multiple tissues. Tissue sections were deparaffinized in xylene, rehydrated in graded alcohol, and subsequently blocked with 0.3% hydrogen peroxide in PBS for
μdUTP, 0.01 U/μl with a linear dynamic range from 100 to 10^6 copies in the presence of genomic DNA.

RhCMV DNA was detected and as copies per microgram input DNA for tissues/saliva/urine. RhCMV DNA was quantified using extension at 60°C for 1 minute. Data are expressed as copies per milliliter for plasma/amniotic fluid.

PCR conditions consisted of an initial 2-minute cycle at 50°C followed by 10 cycles at 95°C, and then 45 cycles of denaturation at 95°C for 15 seconds and combined annealing/extension at 60°C for 1 minute. Data are expressed as copies per milliliter for plasma/amniotic fluid and as copies per microgram input DNA for tissues/saliva/urine. RhCMV DNA was quantified using a standard consisting of a plasmid containing the entire RhCMV IE1 gene (61). Between 5 and 10 ng of DNA was added as template to 50 μl of 1× PCR mixture containing 300 nM of each primer, 100 nM probe, 2 mM MgCl_2, 200 μM each of dATP, dCTP, and dGTP, 400 μM dUTP, 0.01 U/μl AmperaseUNG, 0.025 U/μl Taq polymerase, and reaction buffer that includes passive reference dye ROX. PCR conditions consisted of an initial 2-minute cycle at 50°C followed by 10 minutes at 95°C, and then 45 cycles of denaturation at 95°C for 15 seconds and combined annealing/extension at 60°C for 1 minute. Data are expressed as copies per milliliter for plasma/amniotic fluid and as copies per microgram input DNA for tissues/saliva/urine. RhCMV DNA was detected with a linear dynamic range from 10^2 to 10^6 copies in the presence of genomic DNA.

**SNAPP**. Viral DNA was isolated from plasma using the High Pure Viral Nucleic Acid kit (Roche Life Science). Hypervariable regions 400 bp in length within gB and gL were amplified in duplicate by a single round of PCR using 5′-AAGTGCTCGAAGGCTTCTC-3′ and 5′-CTGGACATTGACGTGCTGTTTGG-3′ for the gB region and 5′-GGGCGGCACACATTATCCTAC-3′ and 5′-GGTGAGTGCTGCTGTTTGG-3′ for the gL region. Overhang regions were conjugated to primer locus–specific sequences for subsequent Illumina index primer addition and sequencing: forward primer overhang = 5′-TCGTCGCGACGCTAGTGATATAAGAGACAG-[locus]-3′ and reverse primer overhang = 5′-GTCTCTGAGGCGAGCTGAGATATAAGAGACAG-[locus]-3′. Approximately 10 ng DNA was added as template to 50 μl of 1× PCR mixture containing 100 nM of each primer, 2 mM MgCl_2, 200 μM each of dNTP mix (Qiagen), and 0.025 U/μl Phusion Taq polymerase. PCR conditions consisted of an initial 2-minute denaturation at 98°C, followed by the minimum number of PCR cycles to achieve adequate amplification (98°C for 10 seconds, 65°C for 30 seconds, and 72°C for 30 seconds), and then a final 72°C extension for 10 minutes. Products were purified using Agencourt AMPure XP beads (Beckman Coulter), and then Illumina Nextera XT index primers were added by 15 cycles of amplification. The resultant PCR product was gel purified using ZR-96 Zymoclean Gel DNA Recovery Kit (Zymogen). The molar amount of each sample was normalized by real-time PCR using the KAPA library amplification kit (KAPA Biosystems). The library of individual amplicons was pooled together, diluted to an end concentration of 10 pM, combined with 20% V3 PhiX (Illumina), and then sequenced on Illumina MiSeq using a 600-cycle V3 cartridge (Illumina). For each primer set we confirmed that there was no significant primer bias through mixing viral DNA in known ratios and applying the SNAPP technique (Supplemental Figure 6).

**SNAPP amplicon reconstruction and nucleotide diversity.** Sequences of the targeted regions on the gB and gL genes were reconstructed by merging the paired-reads sets using the PEAR software under default parameters (62). The fused reads were then filtered using the extractor tool from the SeekDeep pipeline (http://baileylab.umassmed.edu/seekdeep), filtering sequences according to their length, overall quality scores, and presence of the primer sequences. To homogenize the number of sequences per sample due to variable bias through mixing viral DNA in known ratios and applying the SNAPP technique (Supplemental Figure 6).
coverage, a random set of at most 10,000 sequences was retrieved from each sample to estimate strain frequency and nucleotide diversity. The number of sequences analyzed per sample varied from 2,000–10,000, depending on the sample coverage. Next, haplotype reconstruction was performed using the quiver cluster tool from SeekDeep, which accounts for possible sequencing errors by collapsing fragments with mismatches at low-quality positions. For each given sample, the haplotypes had to be present in 2 sample replicates to be taken into account. Each haplotype was assigned to 1 of the 3 inoculated viral strains by first calculating the nucleotide distance (nucleotide substitutions) between the haplotype and the reference viral strain, and then assigning the haplotype to the strain with the shortest nucleotide distance. Nucleotide diversity was computed as the average distance between each possible pair of sequences belonging to the same reference strain (63). Historical control animal 274-98 (24) was omitted from this analysis (Figure 5 and Supplemental Figure 7) because it was only inoculated with a single RhCMV strain (180.92).

Placental RNA isolation and microarray. Tissue sections weighing approximately 30 mg were obtained from flash frozen, full-thickness rhesus placenta/decidua, lysed using a TissueLyser LT (Qiagen), and then total RNA extracted using an RNeasy Mini Kit (Qiagen). RNA was assessed for quality with Agilent 2100 Bioanalyzer G2939A (Agilent Technologies) and Nanodrop 8000 spectrophotometer (Thermo Scientific/Nanodrop) and confirmed to have A_{260}/A_{280} greater than 1.8, A_{260}/A_{230} greater than 1.0, and RIN greater than 7.0. Note that the analysis was not completed for animal HR73 (high-potency HIG infused) because high-quality RNA could not be obtained from the tissue sample. Hybridization targets were prepared with MessageAmp Premier RNA Amplification Kit (Applied Biosystems/Ambion) from total RNA, hybridized to Affymetrix GeneChip Rhesus Macaque Genome arrays in an Affymetrix GeneChip hybridization oven 645, washed in an Affymetrix GeneChip Fluidics Station 450, and scanned with an Affymetrix GeneChip Scanner 7G according to standard Affymetrix GeneChip Hybridization, Wash, and Stain protocols. Affymetrix microarray data were initially processed, underwent QC, and were normalized with the robust multi-array average method using the affy (64) Bioconductor (65) package from the R statistical programming environment (R studio version 0.99.902). Differential expression was carried out using a moderated t-statistic from the limma (66) package. In an attempt to minimize the effect of gestational age on our transcriptome analysis, we corrected for this variable by using a linear model that included time as a covariate (gene expression = condition + time). The FDR was used to control for multiple hypothesis testing. Gene set enrichment analysis (67) was performed to identify differentially regulated pathways and gene ontology terms for each of the comparisons performed. Gene sets specific for innate/adaptive immunity, myeloid/lymphoid lineages, and immune cells were obtained from previous reports (35) and supplemented from the literature such that each comparison group had more than 25 characteristic genes (gene lists in Supplemental Dataset 1). Gene networks were constructed using NatureAnalyst (68), with first-order gene interactions inferred from the InnateDB interactome (Figure 7B) (69). Genes related to NK cell function (Supplemental Table 5) were determined using Qiagen’s Integrated Pathway Analysis (IPA) (NK cell activation and movement gene lists) and supplemented with those from the KEGG database (NK cell mediated cytotoxicity gene list) (70).

Statistics. Comparisons of peak viremia and shedding (Figure 4, E, G, and H) in monkeys from each group were performed using the exact Wilcoxon test. The correlation between peak plasma viral load and initial amniotic fluid viral load (Figure 4F) was calculated with the Spearman correlation. The Benjamini-Hochberg FDR P value correction was used to correct for multiple comparisons (71). For sequence diversity analysis (Figure 5 and Supplemental Figure 7), UCD52 diversity was compared between treatment groups because it was the dominant strain replicating in plasma. First, median UCD52 nucleotide diversity was calculated for each animal. Median values were compared first by Kruskal-Wallis test (P = 0.036), followed by post-hoc Mann-Whitney U test. Analysis of survival curves between standard HIG group and high-potency HIG–infusion group was completed with the exact log-rank test based on the Heinzle macro (Figure 6, A and B) (33). Statistical analysis was performed with SAS v9.4 (SAS Institute). A P value less than 0.05 (2-tailed) was considered significant for all analyses.

Study approval. The animal protocol was approved by the Tulane University and the Duke University Medical Center IACUCs.

Data and materials availability. SNAPP sequence data used in this manuscript are available in the NCBI Sequence Read Archive (SRA) (PRJNA386504, https://www.ncbi.nlm.nih.gov/bioproject/PRJNA386504). Placental transcriptome microarray data are available in the NCBI’s Gene Expression Omnibus (GEO GSE87395; http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE87395).
Author contributions
AK and SRP designed the research. CSN, DT, KMB, MG, RB, XA, and LS performed the research. CSN, DVC, RB, XA, HI, HW, and MC analyzed data. AD, FC, FW, DJD, NV, MW, PAB, MCW, and KK contributed analytic tools/expertise/reagents. CSN, AK, and SRP wrote the manuscript.

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