The remodeling of maternal uterine spiral arteries (SAs) is an essential process for ensuring low-resistance, high-capacitance blood flow to the growing fetus. Failure of SAs to remodel is causally associated with preeclampsia, a common and life-threatening complication of pregnancy that is harmful to both mother and fetus. Here, using both loss-of-function and gain-of-function genetic mouse models, we show that expression of the pregnancy-related peptide adrenomedullin (AM) by fetal trophoblast cells is necessary and sufficient to promote appropriate recruitment and activation of maternal uterine NK (uNK) cells to the placenta and ultimately facilitate remodeling of maternal SAs. Placentas that lacked either AM or its receptor exhibited reduced fetal vessel branching in the labyrinth, failed SA remodeling and reendothelialization, and markedly reduced numbers of maternal uNK cells. In contrast, overexpression of AM caused a reversal of these phenotypes with a concomitant increase in uNK cell content in vivo. Moreover, AM dose-dependently stimulated the secretion of numerous chemokines, cytokines, and MMPs from uNK cells, which in turn induced VSMC apoptosis. These data identify an essential function for fetal-derived factors in the maternal vascular adaptation to pregnancy and underscore the importance of exploring AM as a biomarker and therapeutic agent for preeclampsia.
Fetal-derived adrenomedullin mediates the innate immune milieu of the placenta

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The remodeling of maternal uterine spiral arteries (SAs) is an essential process for ensuring low-resistance, high-capacitance blood flow to the growing fetus. Failure of SAs to remodel is causally associated with pre-eclampsia, a common and life-threatening complication of pregnancy that is harmful to both mother and fetus. Here, using both loss-of-function and gain-of-function genetic mouse models, we show that expression of the pregnancy-related peptide adrenomedullin (AM) by fetal trophoblast cells is necessary and sufficient to promote appropriate recruitment and activation of maternal uterine NK (uNK) cells to the placenta and ultimately facilitate remodeling of maternal SAs. Placentas that lacked either AM or its receptor exhibited reduced fetal vessel branching in the labyrinth, failed SA remodeling and reendothelialization, and markedly reduced numbers of maternal uNK cells. In contrast, overexpression of AM caused a reversal of these phenotypes with a concomitant increase in uNK cell content in vivo. Moreover, AM dose-dependently stimulated the secretion of numerous chemokines, cytokines, and MMPs from uNK cells, which in turn induced VSMC apoptosis. These data identify an essential function for fetal-derived factors in the maternal vascular adaptation to pregnancy and underscore the importance of exploring AM as a biomarker and therapeutic agent for pre-eclampsia.

Introduction

It is widely accepted that the placenta provides the venue for communication between the mother and fetus and that this communication is essential for their continued health throughout pregnancy. During midgestation, the maternal uterine spiral arteries (SAs) must provide increasing amounts of oxygen and nutrients to the growing fetus, and so they undergo active remodeling in order to transform into low-resistance, high-capacitance blood vessels. SA remodeling is a complex event that involves several distinct processes: (a) endothelial cell vacuolization and relaxation of VSMCs that lead to arterial dilation and growth, (b) induction of local placental factors, including MMPs, chemokines, and cytokines, which in turn cause (c) dissolution of the ECM, (d) dedifferentiation and apoptosis of VSMCs, and (e) replacement of VSMCs by invasive fetal trophoblast cells (1–5). Considering the complexity of this process, it is not surprising that preeclampsia—a life-threatening pregnancy complication caused by poor SA remodeling—is so prevalent and detrimental to both mother and fetus (6, 7). Since the fetus is the primary beneficiary of increased blood flow, it stands to reason that fetal signals may play an active role in enabling SA remodeling. However, very few fetal-derived trophoblast factors involved in SA remodeling have been identified (8–11).

On the other hand, maternal uterine NK (uNK) cells, which do not exhibit the cytotoxic behavior of peripheral NK cells, are known to be required for SA remodeling (4, 11, 12). uNK cells are the most abundant of all decidual leukocytes, accounting for approximately 70% of CD45+ cells, and are histologically identified by large cytoplasmic granules containing perforin and granzymes (1). uNK cells, which are transiently present in the decidua, are frequently aggregated around SAs and play a functional role in remodeling (12–14). Pioneering studies by Croy and colleagues have demonstrated that the SAs of transgenic mice that lack uNK cells show failed SA remodeling and preeclampsia. Several recent studies have begun to reveal important pathophysiological roles for uNK cells in human pregnancy (19–22). However, much remains unknown about the precise molecular interactions between fetal trophoblast cells and uNK cells in the placenta. It is also unclear whether perturbations in this fetal-to-maternal communication can account for, or be diagnostic of, failed SA remodeling and preeclampsia.

Adrenomedullin (Adm; encoding AM) is a hypoxia-induced, angiogenic peptide vasodilator that is elevated approximately 5-fold in the maternal plasma of normal human pregnancies, but often blunted in pregnancies complicated by preeclampsia (23). Moreover, the direct transcriptional induction of Adm gene expression (and that of its receptor components) by estrogen confers high levels of this peptide signaling pathway in female reproductive organs, including the ovaries, uterus, uterine arteries, and placenta (24–27). Yallampalli and colleagues showed that systemic administration of an AM antagonist during late rat gestation caused a myriad of pregnancy abnormalities, including fetal growth restriction, placental and fetal membrane necrosis, and...
fetal edema (28). While the AM antagonist did not change maternal systemic blood pressure, it is likely that many of these fetal-placental sequelae developed as a consequence of acutely reduced uterine perfusion (29).

Using gene-targeted mice, we previously showed that full expression of maternal Adm is required for the early establishment of pregnancy and placentation: modest haploinsufficiency for maternal Adm caused poor uterine receptivity, reduced pinopode formation, and subfertility, even when WT embryos were transferred to the uterus (30). Consequently, pregnant Adm–/– females exhibit abnormal implantation, ectopic placentation, and fetal growth restriction that results in a wave of embryonic lethality around E9.5 that is largely independent of fetal genotype (31). These findings highlight the importance of maternal Adm genetic dosage during the early stages of pregnancy, yet the function of fetal Adm expression during pregnancy remains unclear.

Here, using both loss-of-function and gain-of-function genetic mouse models, we showed that expression of Adm and its G protein–coupled receptor by fetal trophoblast cells was necessary and sufficient to promote appropriate recruitment and activation of maternal uNK cells to the placenta and ultimately facilitate the remodeling of maternal SAs.

Results

Adm-null placentas exhibit parietal trophoblast giant cell death and reduced labyrinth vessel branching. We previously showed that Adm is expressed in trophectoderm cells of the preimplantation blastocyst and that its expression is increased when trophoblast stem cells are differentiated into the giant cell lineage (31). Using in situ hybridization, we found that this robust expression was maintained in vivo within fetal parietal trophoblast giant cells (TGCs); polyploidy derivatives of the primary mural trophectoderm cells that line the implantation site; refs. 32–34) at E9.5 and E13.5 (Figure 1, A–C), consistent with previous studies (35). There was also strong expression of Adm surrounding the invading ectoplacental cone at E9.5 and at the innermost border of the maternal-fetal interface (Figure 1, A and B). Adm expression persisted in TGCs at E13.5, and stromal cells of the maternal decidua expressed moderate levels of Adm (Figure 1C). The chorionic plate, embryo, fetal membranes, labyrinth, and spongiosotrophoblast-containing junctional zone expressed little to no Adm.

Although Adm–/– mice die at midgestation with lymphatic vascular defects (36), we wondered whether genetic dosage of fetal Adm could also contribute to pregnancy outcomes. Histological and morphometric analyses showed that Adm–/– placentas appeared overtly normal, with well-formed layers and labyrinth thickness that were indistinguishable from those of Adm+/+ littermates (Figure 1D and Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI67039DS1). Because a hallmark function of AM peptide is vasodilation, some have speculated that its expression in placenta may be associated with maintaining low-resistance blood flow through the placental vasculature and umbilical cord (37). Contrary to this hypothesis, we found that loss of Adm had no effect on placental impedance, as calculated by umbilical cord blood flow using in vivo Doppler ultrasound (Supplemental Figure 2).

Despite their overtly normal appearance and function, E13.5 Adm–/– placentas had a prominent band of apoptosing cells that spatially localized to the innermost border of the maternal-fetal interface (Figure 1E), where Adm-expressing parietal TGCs are located, which suggests that AM is an essential survival factor for these cells. Consistent with this finding, qRT-PCR analysis of trophoblast marker genes revealed that Adm+/+ placentas had significantly reduced expression of placental lactogen 1—a specific marker of parietal TGCs—while the expression of other TGC lineage markers and spongiosotrophoblast markers did not differ significantly from that of WT placentas (Supplemental Figure 3). Even though Adm is not abundantly expressed in the
labyrinth, we also found a modest, yet significant, reduction in the labyrinth-expressing genes Gcm1, JunB, and Gab1 (Supplemental Figure 3).

Further characterization of the labyrinth layer showed marked defects in the vascular patterning of fetal vessels in Adm–/– placentas. Isolectin B4 staining of the ECM surrounding fetal labyrinth vessels revealed a highly branched network in Adm+/+ placentas, but abnormally large and underbranched fetal vessels in Adm–/– placentas (Figure 2A). Using nucleated and non-nucleated blood cells as markers for fetal and maternal blood sinuses, respectively, we performed quantitative morphometric analysis of the labyrinth vasculature (Figure 2B). Consistent with our finding of normal labyrinth thickness and normal placental impedance, the total area of vascular space in Adm–/– labyrinth did not differ from that in Adm+/+ placentas (Figure 2C). However, we found that individual fetal sinuses of Adm+/+ placentas were significantly larger than those of Adm+/+ placentas (P < 0.01; Figure 2D). This increase in the size of fetal sinuses was accompanied by a concomitant reduction in the size of maternal sinuses in Adm+/+ versus Adm+/+ placentas (P < 0.02; Figure 2D).

To better visualize the architecture and structure of the labyrinth vasculature, we made vascular corrosion casts of the placentas by perfusing pregnant female mice with a liquid polymer. Fetal placental tissue was enzymatically digested away from the hardened maternal casts in order to reveal the interdigitation of maternal and fetal vasculature within the labyrinth placentas of Adm+/+ and Adm–/– littermates. Therefore, in the scanning electron micrographs of Figure 2, E and F, the open spaces represent the footprint of the fetal vasculature as it was interdigitated between the maternal blood sinuses within the central labyrinth layer. Adm–/– placentas had large holes (Figure 2E, arrowheads) and higher-magnification views demonstrated vast spacing between the maternal sinuses (Figure 2F), reflective of the large and underbranched phenotype of Adm–/– fetal vessels. Taken together, these data demonstrated a failure of fetal vessels to branch appropriately in Adm–/– placentas.

Because shallow invasion of chorionic villi can lead to blunted fetal vascular branching, we used alkaline phosphatase staining to evaluate the extent of chorionic villus formation and found no significant difference between WT and Adm+/+ placentas (integrated density, Adm+/+, 11.8 ± 2.0 AU; Adm–/–, 11.3 ± 2.0 AU; Figure 2G). This finding demonstrated that the fetal vascular branching defects of Adm–/– placentas were inherent to the fetal vasculature, rather than secondary to defects in chorionic villus invasion or formation. Collectively, the data presented in Figures 1 and 2
showed that compared with those of their WT littermates, Adm–/– placenta exhibited increased apoptosis of parietal trophoblast cells accompanied by reduced fetal vessel branching.

**Failed SA remodeling and reduced uNK cells in Adm-null placentas.**

Because several clinical studies have shown that maternal levels of AM are reduced in preeclampsia (23), we evaluated another characteristic pathology of preeclampsia in Adm–/– placentas: remodeling of maternal SAs. As expected, SAs of WT placentas progressively lost staining for SMA as they approached the maternal-fetal interface (Figure 3A). In contrast, SAs from Adm+/+ placentas retained a thick smooth muscle layer throughout the decidua that was nearly twice the thickness measured in WT placentas (P < 0.01; Figure 3, A and B). The shedding of VSMCs surrounding SAs is associated with localized degradation of the ECM, a process in which the cathepsin family of proteases has been shown to play a predominant role (10, 38). As expected, SAs from Adm+/+ placentas were tightly surrounded by a prominent band of cathepsin staining, whereas cathepsin staining was weak and diffuse throughout the decidua of Adm–/– placentas and did not surround the maternal SAs (Figure 3C), consistent with a failure of these SAs to shed their smooth muscle coverage. The enlargement of maternal SAs at midgestation is also associated with proliferation of vascular endothelial cells, a process termed reendothelialization. Using BrdU incorporation assays, we readily observed this robust endothelial cell proliferation in all SAs of Adm+/+ placentas, but this was rarely evident in the SAs of Adm–/– placentas (Figure 3D).

Numerous studies have pointed to maternal uNK cells as essential mediators of SA remodeling, and the precise factors contributing to their recruitment, activation, and function remains an area of intense study. Therefore, we stained and counted the number of uNK cells using 2 different uNK markers that identify mature (perforin-containing) and/or activated (Ly49G2+) uNK cells. Consistent with the phenotype of failed SA remodeling, we found that decidua of Adm–/– placentas had half the number of perforin-containing uNK cells observed in Adm+/+ littermates (P < 0.05; Figure 3, E and F). Similarly, levels of Ly49G2+ uNK cells were also reduced by approximately half in Adm–/– placentas, although the difference did not reach statistical significance.

Taken together, our findings demonstrated that lack of fetal AM caused numerous placental pathologies, including reduced fetal vessel branching, failed SA remodeling, and reduced number of uNK cells. We also noted that the incidence and degree of the Adm–/– placental phenotype was strictly and exclusively correlated with the genotype of the individual Adm–/– conceptus, not influenced by the genotype of neighboring conceptuses.

**Adm–/– placental phenotype is independent of maternal genotype.** A confounding factor to the present findings is that the maternal genotype used in the above studies was haploinsufficient for AM; we have
previously shown that during early gestation, Adm+/- females exhibit reduced fertility due to abnormal implantation (30, 31). To eliminate this confounding factor, we performed ovary transplantations in which Adm+/- or Adm-null female mice were surgically sutured in the ovarian bursa of WT 129S6/SvEv recipient mice after removal of their own ovaries. The recipient females were then bred to Adm+/- male mice to generate Adm+/-, Adm+, or Adm-/- offspring (B) Mendelian ratios of offspring were as expected, and Adm-/- mice exhibited characteristic embryonic edema (arrowhead) at E13.5. (C and D) Recovery time to first parturition (C) and average litter size (D). Numbers within bars denote total number of litters analyzed per breeding. (E–I) Adm-/- placentas from Adm-/- ovary→WT recipient females displayed the same pathological phenotypes of Adm-/- placentas from Adm-/- females — including (E) reduced fetal vessel branching in the labyrinth, (F) retention of SMCs (arrowheads) around maternal SAs, (G) reduction in endothelial cell proliferation (arrowheads) in maternal SAs, and (H and I) significantly reduced DBA+ uNK cell numbers — compared with Adm+/- littermates. *P < 0.05. For ovary transplant studies, n = 6–10 placentas per genotype. Scale bars: 50 μM.
The cross of Adm<sup>−/−</sup> males to Adm<sup>+/−</sup> ovary→WT recipient females resulted in the expected Mendelian ratio of fetal genotypes and the previously characterized phenotype of extreme embryonic edema and lethality for Adm<sup>−/−</sup> fetuses (Figure 4B and refs. 36, 39). Consistently, the placental phenotypes of underbranched fetal labyrinth vessels, poorly remodeled SAs, and significantly reduced uNK cell numbers were completely recapitulated in Adm<sup>−/−</sup> placentas born to Adm<sup>+/−</sup> ovary→WT recipient females compared with Adm<sup>+/+</sup> littermate placentas and Adm<sup>+/−</sup> placentas derived from Adm<sup>+/+</sup> ovary→WT recipient females (Figure 4, E-I). Thus, loss of Adm from fetal tissues directly contributed to the observed placental pathologies, even in the context of genotypically normal maternal Adm.

Recapitulation of phenotypes in placentas genetically lacking the AM receptor. AM peptide can bind to and signal through the G protein-coupled receptor calcitonin receptor–like receptor (Calcrl; encoding CLR). We have previously shown that mice with genetic deletion of Calcrl phenotypically recapitulate the embryonic lethality and lymphatic vascular defects observed in Adm<sup>−/−</sup> mice (36, 40), thereby providing genetic and in vivo evidence to substantiate CLR as a bona fide AM receptor.
Figure 6

Genetic overexpression of fetal Adm reverses the placental preeclampsia phenotypes and drives uNK recruitment to the decidua. (A) Targeting vector for generation of Admhi/hi mice consisted of (a) a 6-kb genomic fragment of the Adm gene isolated from a 129S6/SvEv genomic phage library and containing all 4 exons and 5′UTR and 3′UTR of the Adm gene, (b) the bovine growth hormone polyA sequence (bGH 3′UTR), (c) 2 tandem copies of the 1.2-kb 5′ insulator sequence from chicken β-globin gene (2XIns), (d) 1.3 kb of pMC1 promoter–driven neomycin (Neo), (e) 80 bp of AU/U-rich element of the mouse c-fos gene (ARE), and (f) 2 loxP recombination sites. The latter 5 elements were cloned as a cassette, 23 bp downstream of the endogenous Adm stop codon. (B) Southern blot analysis on genomic DNA confirmed correct targeting of the Admhi allele. (C) Adm gene expression in placentas from Adm+/+ and Admhi/hi mice, analyzed by quantitative RT-PCR, showed a significant 3-fold increase in gene expression level. *P < 0.05.

(D–G) Admhi/hi placentas (D) appeared histologically comparable to Adm+/+ placentas and showed (E) highly branched fetal labyrinth vessels, (F) appropriate SA remodeling, and (G) reendothelialization. (H and I) DBA staining revealed that Admhi/hi placentas had a significant 30% increase in uNK cell numbers compared with Adm+/+ littermate placentas. *P < 0.05. Data are mean ± SEM. Scale bars: 1 mm (D); 50 μM (E–H).
Using in situ hybridization, we consistently found high levels of Calcrl expression in WT placentas that spatially and temporally colocalized with that of the peptide ligand Adm. Specifically, Calcrl expression was robustly present in parietal TGCs at both E9.5 and E13.5 (Figure 5, A–C). In contrast to the diffuse expression of Adm surrounding the ectoplacental cone and stromal decidual cells, expression of Calcrl was concentrated to endothelial-like cords of the E9.5 decidua (41) and to maternal endothelial cells lining the maternal SAs of E13.5 placentas (Figure 5C, inset).

Like Adm–/– placentas, the Calcrl –/– placentas appeared overtly normal and contained well-formed placental layers and normal histomorphometric features (Figure 5D). Nevertheless, the Calcrl –/– placentas exhibited the same abnormal phenotypes of Adm–/– placentas, including reduced fetal vessel branching, retention of SMCs surrounding maternal SAs, and significantly reduced uNK cell numbers compared with Calcrl+/+ littermate placentas (Figure 5, E–H). Thus, the recapitulation of the Adm–/– placental phenotype in Calcrl –/– placentas further supports the role of CLR as an AM receptor within this tissue and indicates that lack of AM signaling within the fetal compartment is causally associated with these phenotypes.

Overexpression of fetal Adm reverses the placental phenotype and drives maternal uNK cell recruitment to the decidua. The absence of apoptosis within decidual regions that contain uNK cells (Figure 1E) suggests that the reduced number of uNK cells in Adm–/– placentas may be caused by abnormalities in their recruitment to the tissue, rather than their loss through cell death. To determine whether fetal AM can directly influence the recruitment of uNK cells to the decidua, we generated and characterized a novel gene-targeted mouse model in which AM expression was increased approximately 3-fold (Figure 6, A–C). The gene targeting was designed to increase AM expression approximately 3-fold via stabilization of mRNA levels through genetic modification of the 3′ untranslated region (3′ UTR) of the endogenous Adm gene.

Adm+/hi mice were viable and fertile and were intercrossed in order to compare placental phenotypes between Adm +/+ and Adm hi/hi littermates. The average litter size of Admhi/hi females was 6.04 live births, which was not significantly different than the 6.60 pups per litter observed for WT C57BL/6J mice of the comparable genetic background. Admhi/hi placentas born to Adm +/hi intercrossed mice appeared overtly normal and had appropriately branched labyrinth fetal vessels that were indistinguishable from Adm +/+ littermates (Figure 6, D and E). Moreover, the SAs of Admhi/hi placentas were appropriately remodeled and did not differ from those of Adm+/+ littermates (Figure 6, D and E). Interestingly, Dolichos biflorus agglutinin (DBA) staining and counting of uNK cells revealed that Admhi/hi placentas had a significant 30% increase in the number of uNK cells within the decidua compared with that of Adm+/+ littermate placentas (Figure 6, H and I). These data indicate that 3-fold overexpression of Adm is compatible with normal placental develop-
Development and, moreover, that fetal-derived AM can actively promote the recruitment of uNK cells to the placenta.

AM dose-dependently alters the chemokine, cytokine, and MMP profiles of uNK cells in vivo and in vitro. uNK cells constitute the largest proportion of immune cells in the decidua, and so we expected and found that the dynamic fluctuations in uNK cell recruitment between Adm–/– and Admhi/hi placentas were reflected by concomitant changes in the expression of numerous chemokines and cytokines. Specifically, the expression of Ccl7, Ccl17, Cxcl9, Cxcl10, Xcl1, and TNF were downregulated in Adm–/– placentas and concomitantly upregulated in Admhi/hi placentas compared with their respective WT controls (Figure 7A). To determine whether AM peptide directly causes secretion of chemokines and cytokines from uNK cells, we isolated uNK cells and first determined that these cells expressed high levels of Calcrl and survived up to 48 hours in culture without evidence of apoptosis, as evaluated by TUNEL and trypan blue exclusion (Supplemental Figure 5 and data not shown). Then, using a Luminex-based detection system, we found that treatment of uNK cells with 10 nM AM stimulated marked secretion of a variety of chemokines and cytokines (Figure 7B), several of which have established functions in reproductive immunology (i.e., CXCL10, GM-CSF, and IL-23) (42). Furthermore, we found that AM treatment dose-dependently increased the gene expression and activity of MMP9, but not MMP2, in uNK cells (Figure 7C), consistent with the previously described functions of uNK-derived MMP9 in SA remodeling (43). Collectively, these data suggest that fetal-derived AM potently influences the immune milieu of the placenta in 2 ways: first by recruiting, then activating, uNK cells to secrete chemokines, cytokines, and MMPs, which are important contributors to SA remodeling.

AM-treated uNK cells stimulate VSMC apoptosis. Finally, we sought to determine whether the effects of AM on isolated uNK cells directly relate to the phenotype of reduced SA remodeling in Adm–/– placentas. To test this, we developed an in vitro cell culture system in which primary mouse VSMCs were treated with control media or uNK-conditioned media that had been supplemented or not with 10 nM AM peptide. As predicted, treatment of VSMCs with control media and AM-supplemented control media had no obvious effects on cell growth, morphology, or apoptosis (Figure 8A and B). In contrast, treatment of VSMCs with uNK-conditioned media resulted in dramatic changes in cellular morphology and apoptosis rate. Importantly, these effects were markedly and significantly exacerbated when the uNK-conditioned media was supplemented with AM, with a nearly 4-fold increase in the ratio of proapoptotic Bax to antiapoptotic Bcl2 gene expression (Figure 8C). Thus, we conclude that AM can promote the secretion of a cocktail of chemokines and cytokines from uNK cells that can in turn induce apoptosis of VSMCs.

Discussion

Using both loss-of-function and gain-of-function genetic mouse models, we found that fetal AM, acting through CLR, serves as a trophoblast-derived factor critical for fetal placental vascularization and for the maternal vascular adaptation to pregnancy (Figure 9). There were several pleiotropic consequences of absent AM/CLR signaling within the trophoblast compartment, which could be difficult to causally distinguish from one another. However, the precise replication of the Adm–/– phenotype in Calcrl–/– placentas demonstrated that the predominant factor in initiating the complex phenotype is the absence of ligand signaling and secretion in the trophoblast cell. Ultimately, the mechanisms of AM action appear to be largely mediated through its local effects on the decidual content and activation of maternal uNK cells, which in our experiments expressed either WT or hemizygous levels of CLR. Therefore, the phenotypes associated with Adm–/– and Calcrl–/– placentas, and their reversal by Adm overexpression, demonstrated that the dosage of AM provided by the fetus is a critical communication factor that has a profound effect on the innate immune milieu of the placenta. It is worth noting that nearly all of the characteristic features of Adm–/– and Calcrl–/– placentas were similar to the pathological features that have been characterized in human preeclampsia. Although it is imperative to consider the anatomical and func-
The genetic dosage of fetal AM is directly proportional to the content of maternal uNK cells and their secreted chemokines, cytokines, and MMPs within the placenta. As a consequence, AM expression in fetal trophoblast cells is necessary for appropriate branching of the fetal vasculature and remodeling of maternal SAs. Overexpression of AM within the fetal compartment is sufficient to promote uNK recruitment and activation in the decidua, a process that likely contributes to AM-induced activation of MMP9 and maternal SA remodeling.

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the maternal-fetal interface. Therefore, it becomes of importance to identify genetic, physiological, and/or environmental factors that regulate Adm expression within the fetal trophoblast cell. It has long been appreciated that estrogen and hypoxia can potently upregulate Adm expression in a variety of cell types, including those of reproductive relevance (24, 59). Most interestingly, it has recently become clear that cigarette smoke, likely acting through 2 functional xenobiotic response elements proximal to the Adm start codon (60), can also potently induce Adm expression in trophoblast cells (61). Moreover, exposure of humans and mice to cigarette smoke during pregnancy has recently been linked to a statistically significant increase in uNK cell numbers (62). Thus, although cigarette smoking during pregnancy has a wide range of deleterious effects, the cigarette smoke–induced upregulation of Adm expression in fetal trophoblast cells, and the potential subsequent increase in uNK cells, may provide the mechanistic link for explaining the decades-old clinical observation that moderate smoking during pregnancy is actually protective against pre-eclampsia (63, 64).

In summary, our present findings revealed a critical function for a fetal-derived trophoblast factor in the maternal adaptation to pregnancy. The increased clinical usage of MR-proAM plasma levels as a biomarker for a variety of cardiovascular conditions underscores the importance of exploring AM as a biomarker for preeclampsia. Because CLR is already being exploited as a pharmacological target for the treatment of migraine (65, 66), it is conceivable that direct targeting of the AM/CLR axis may hold tremendous clinical benefit for the amelioration of pregnancy outcomes.

Methods

Mice. Generation of mice with a targeted, homozygous deletion of the Adm and Calrcl genes has been previously described (39, 40). These mice were maintained on an inbred 129S6/SvEv background.

Admhi/c mice were generated by standard gene targeting protocols. The endogenous 3′UTR of the Adm gene was replaced with a cassette (parts of which were provided by M. Kakoki, The University of North Carolina at Chapel Hill) containing the bovine growth hormone UTR (bGH 3′UTR) in E14 ES cells. Southern blot analysis was performed by digesting genomic DNA with BamHI, which revealed an 11.4-kb fragment representing the WT allele and a 3.9-kb fragment corresponding to the Admhi allele. Several chimeric animals were generated, and germline transmission of the targeted allele was established on a mixed genetic background. F1 heterozygotes from male chimera–to–female C57BL/6J breedings were backcrossed to C57BL/6J mice for more than 10 generations. Routine genotyping of the Admhi allele was performed by a 3-primer, PCR-based strategy (primer 1, 5′-AACCTTACACCTTGCTGAGACATC-3′; primer 2, 5′-TTTATTAGGAAAGGACAGTGAGGATG-3′; primer 3, 5′-CCCCATCTCGTGTCACACGCTAC-3′). Primers 1 and 3 amplify a 760-bp WT allele, while primers 2 and 3 amplify a 600-bp targeted allele.

For timed pregnancies, heterozygous male and female Admhi or Admhi/c intercrosses were established, and the day when the vaginal plug was detected was considered E0.5. Fetal genotypes were determined by establishing allele-specific, PCR-based assays (39, 40) using genomic DNA isolated from fetal membranes or tail biopsies.

Ovary transplantation. Ovary transplantation experiments were performed as previously described (67) using donor and recipient animals 8–16 weeks old on the 129S6/SvEv genetic background. Hemi ovaries from Admhi or Admhi/c mice were surgically sutured into the ovarian bursa of recipient WT mice after bilateral removal of the recipient’s own ovaries. 2 weeks after surgery, Admhi or ovary→WT and Admhi/c ovary→WT recipients were used in experiments.

Doppler ultrasound. Intravital Doppler ultrasound was performed on anesthetized female mice at E12.5 using the VisualSonics Vevo 770 system with a 40-MHz probe. Placental impedance was calculated from umbilical cord waveforms by dividing the arterial venous delay by the cycle length.

Placental vascular corrosion casting and scanning electron microscopy. Vascular corrosion casting of E13.5 Admhi and Admhi/c placentas from littersmates of Admhi male and female intercrosses were performed as previously described (68). The castings were processed for scanning electron microscopy as previously described (30) and were viewed and photographed with a Cambridge Stereoscan S200 scanning electron microscope (LEO Electron Microscopy Inc.) using an accelerating voltage of 20 kV.

In situ hybridization. In situ hybridization was performed as previously described (30) using nonradioactive in situ hybridization reagents (1745816, 1585762, 1093274; Roche Diagnostics).

Cytokine expression arrays and cytokine antibody array. RNA was extracted from whole placentas using TRIzol reagent (15596-026; Invitrogen) and purified with Qiagen RNasey Minni (74104; Qiagen). cDNA was made using the RT First Strand Kit (C-03, SAbiosciences), and cytokine expression was determined using RT Profiler PCR Array System for Mouse Chemokines and Receptors (PAMM-022A-2, SAbiosciences). Data were analyzed using the ΔΔCT method. Analysis for the simultaneous detection of cytokines and chemokines was performed with the Cytokine Antibody Array 1 (ARY006; R&D) following the manufacturer’s instructions.

TUNEL and BrdU staining. TUNEL in situ cell death detection kit (1 684 795; Roche Applied Science) was used to identify apoptotic cells. Cell proliferation was marked with a single intraperitoneal injection of 100 mg BrdU per kilogram body weight (B9285; Sigma-Aldrich). BrdU Staining kit (93-3943; Zymed) was used to detect BrdU+ cells, which were quantified using Image J software.

uNK cell isolation and culture of primary aortic VSMCs. uNK cells were isolated as previously described using the Dynal CELLECTION Biotin Binder Kit (115.33D; Dynal) in conjunction with biotinylated DBA lectin (L6533; Sigma-Aldrich) (17). Isolated uNK cells were cultured in complete RPMI media with 10% FCS for 24 hours and then supplemented with 10 nM AM or PBS vehicle. Cells and media were collected for analysis 24–48 hours after isolation. Aortic VSMCs were isolated from C57BL/6J mice and cultured as described previously (69).

Immunohistochemical antibodies. Antibodies were as follows: anti–α-SMA (A2547; Sigma-Aldrich), anti-perforin (804-057-C100; Alexio Biochemicals), ly49G2 (55-5314; BD Biosciences—PharMingen), anti–placental alkaline phosphatase (CBL207; Millipore), cathepsin S M-19 (6505; Santa Cruz), FITC-conjugated isoclin B4 (L2895; Sigma-Aldrich), FITC-conjugated DBA lectin (L9142; Sigma-Aldrich).

Statistics. In all figures, data are presented as mean ± SEM. Statistical analyses were performed with JMP software (SAS). 2-tailed Student’s t test was used to compare 2 groups; 1-way ANOVA was used to compare more than 2 groups. A P value less than 0.05 was considered significant. Exact P values and n for each experiment are provided in the figure legends.

Study approval. All experiments using animals were approved by the Institutional Animal Care and Use Committee of The University of North Carolina at Chapel Hill.

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In summary, our present findings revealed a critical function for a fetal-derived trophoblast factor in the maternal adaptation to pregnancy. The increased clinical usage of MR-proAM plasma levels as a biomarker for cardiovascular conditions underscores the importance of exploring AM as a biomarker for pre-eclampsia. Because CLR is already being exploited as a pharmacological target for the treatment of migraine (65, 66), it is conceivable that direct targeting of the AM/CLR axis may hold tremendous clinical benefit for the amelioration of pregnancy outcomes.
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