Engraftment of Bone Marrow from Severe Combined Immunodeficient (SCID) Mice Reverses the Reproductive Deficits in Natural Killer Cell-deficient tge26 Mice

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Summary

A large, transient population of natural killer (NK) cells appears in the murine uterine mesometrial triangle during pregnancy. Depletion of uterine (u) NK cells, recently achieved using gene-ablated and transgenic mice, results in pathology. Pregnancies from matings of homozygous NK and T cell–deficient tge26 mice have <1% of normal uNK cell frequency, no development of an implantation site–associated metrial gland, and an edematous decidua with vascular pathology that includes abnormally high vessel wall/lumens ratios. Fetal loss of 64% occurs midgestation and placentae are small. None of these features are seen in pregnant T cell–deficient mice. To confirm the role of the NK cell deficiency in these reproductive deficits, transplantation of tge26 females was undertaken using bone marrow from B and T cell–deficient scid/scid donors. Engrafted pregnant females have restoration of the uNK cell population, induced metrial gland differentiation, reduced anomalies in the decidua and decidual blood vessels, increased placental sizes, and restoration of fetal viability at all gestational days studied (days 10, 12, and 14). Thus, uNK cells appear to have critical functions in pregnancy that promote decidual health, the appropriate vascularization of implantation sites, and placental size.

Large, heavily granulated lymphocytes (LGLs)

Large, heavily granulated lymphocytes (LGLs) are found in the pregnant uteri of many species, including mice, rats, pigs, and humans (1). Due to their dependency on estrogen and progesterone and their absence from the uteri of virgin and postpartum animals, these lymphocytes are expected, but have not yet been proven, to have important, pregnancy-associated functions. By midgestation in rodents (days 10–14 of gestation in mice), up to 35% of the cells in a specialized, pregnancy-induced region within the uterine musculature called the metrial gland are granulated cells commonly called granulated metrial gland cells (2). In pregnant women, uterine LGLs are more widely dispersed throughout decidial tissue and represent 70% of the leukocytes in decidual cell suspensions (3).

Immunophenotyping of the surface and the cytoplasmic granules of rodent and human pregnancy-associated LGLs has led to the hypothesis that uterine LGLs are lymphocytes of the NK cell lineage (4, 5). Studies using immune-deficient mice support this conclusion, since LGLs typical in morphology, location, and frequency have been reported in T cell–deficient mice (6, 7), and NK1.1 T cell–deficient mice (8). However, LGLs and a metrial gland structure were absent from the uteri of pregnant IL-2Rγ null, p56lck null × IL-2Rβ null, and tge26 mice (8, 9). Because an NK cell deficiency accompanies the T cell deficiency in the three latter strains, uterine LGLs are now regarded as members of the NK cell lineage and designated uterine (u)NK cells (8, 9).

Depletion of uNK cells during pregnancy has only been achieved genetically. A transgenic strain, tge26, carrying high copy numbers (30–35) of the full human (hu) CD3ε gene exhibits NK and T cell deficiencies from fetal life (10, 11). In addition to their splenic NK cell deficiency (10), uNK cells are deficient in pregnant tge26 (range from 0 to 3% of the frequency found in immune-competent, random-bred H-2b CD1 mice and inbred H-2k CBA/J mice [8]). Metrial glands also fail to differentiate in tge26 implantation sites (8).

In immune-competent mice, the metrial gland is associated with the maternal portion of the placenta that consists of transformed stromal cells known as the decidua. The fetal portion of the placenta has three layers: giant cell trophoblast, spongiotrophoblast, and labyrinthine trophoblast. Implantation sites in tge26 mice appear normal, both grossly and histologically, until day 9 of gestation (counting from the copulation plug as day 0). By day 10 of gestation,
the decidua lacks normal cellularity and the large ablumenal decidual blood vessels have thickened walls. There are progressive degenerative changes to both the media and endothelium over the next four d of gestation. Vascular anomalies are not found in other organs of pregnant or nonpregnant tge26 mice (8).

Death of >50% of the tge26 fetuses occurs in each litter between days 10 and 14 of gestation, a time correlating with the decidual pathologies. Surviving tge26 fetuses have very small placentae (~50% of normal) and from birth to adulthood tge26 mice are smaller than immune-competent, age-matched controls (36% smaller at 24 h after birth and 27% smaller at 7 wk of age) (8). These data suggested that uNK cells have major functional roles in optimizing pregnancy success.

To understand the role of uNK cells in the reproductive deficits of tge26 females, three further studies were undertaken. First, ectoplacental cones from tge26 mice were examined for expression of the huCD3ε transgene. No expression was detected. Second, morphometry was applied to the maternal decidual arterial sinuses, which are murine equivalents to human spiral arteries (12). Elevated wall to lumen ratios were found, suggestive of hypertension. Third, bone marrow was transplanted from syngeneically mated tge26 females into syngeneically mated CD1 and CBA/J mice (H-2k; The Jackson Laboratory, Bar Harbor, ME), C.B-17 scid/scid (data not shown). Eight median independent tissue sections were analyzed, including the central section and sections on both sides that were at least 36 μm apart. Cross-sectional area measurements of decidual blood vessel wall and vessel lumen and placental cross-sectional areas were measured from a minimum of two implantation sites per pregnancy. The limits used for the definition of placental cross-sectional area were trophectodermal tissue up to and including the trophoblast giant cell layer. The decidua and metrial gland were excluded.

Statistical Analysis. Means, standard errors, standard deviations, P values, and paired t tests were performed using the computer software program Microsoft Excel 5.0 for Windows (Microsoft Corp.).

Results and Discussion

Tge26 Trophoblast Does Not Express the huCD3ε Transgene. One possible explanation for small placentae in tge26 pregnancy is aberrant expression of the huCD3ε transgene in trophoblast cells. We first established that neither CD1 nor tge26 trophoblast expressed immunoreactive murine CD3 (data not shown). Then, day 7 tge26 ectoplacental cone tissue (the fetally derived structure from which all trophoblast arises) was examined for huCD3ε expression by Northern blotting and a more sensitive reverse transcriptase PCR approach. Day 7 was selected because it is the latest developmental time point at which pure trophoblast can be isolated. The positive control was RNA from thymocytes of heterozygous tge600 mice (13), which carry a different copy number of the same transgene. Human CD3ε mRNA was not detected by either method in the trophoblast tissue but was present in the heterozygote thymocyte RNA (data not shown). Thus, it is unlikely that the reduced size of placental trophoblast in tge26 is due to transgene expression within the trophoblastic lineages.

Tge26 Ectodermal Decidua and Quantification of Decidual Vessel Histopathology. The mesometrial decidua in tge26 is very unusual, being relatively acellular and containing vessels with thickened walls. Mason’s trichrome staining did not detect matrix deposition in the cell-deficient regions (data not shown). Since neutral red staining of cryostat sections has also eliminated fat deposition as an explanation for this relative acellularity, we interpret the decidual region as edematous (9).

Normal pregnancy is associated with progressive dilation of uterine blood vessels and their derivative branches. Since this was apparently not occurring in tge26 pregnancies, the anomalous vessels were studied morphometrically. Table 1...
summarizes the ratios of \(tg_{26}\) vessel wall to lumen cross-sectional area measurements and compares them to measurements of the similar vessels in the ablumenal decidua of CBA/J. In immune-competent control animals, a progressive increase in the ratio of the lumen to wall was found. This is consistent with the hypotensive environment promoted in mammalian uterus during mid-gestation (15–17).

At days 10, 12, and 14 of gestation, \(tg_{26}\) vessel walls showed a thickened media. This was measured at 2.7, 2.0, and 4.7 times greater than the media thickness in control, gestational day–matched CBA/J. The decidual vessels in \(tg_{26}\) had relatively less lumen, suggesting vasoconstriction or failure to undergo arterial vessel modification that would facilitate vasodilation. Either mechanism would be expected to promote thickened decidual vessel walls and a more hypertensive environment in the \(tg_{26}\) uterus at midpregnancy.

Masson’s trichrome intensely stained the muscular component of the vessel walls (red). There was little evidence of increased connective tissue reactivity (blue) on days 10, 12, and 14 of gestation. This demonstrates that the media was the thickened component in \(tg_{26}\) decidual vessel walls.

**Table 1.** Summary of Ratios of Mean Decidual Vessel Wall to Lumen Cross-sectional Area Measured at Days 10, 12, and 14 of Gestation in \(tg_{26}\) and CBA/J Mice

| Mouse strain | Day 10 of gestation (mean W/L* ± SEM) | Day 12 of gestation (mean W/L ± SEM) | Day 14 of gestation (mean W/L ± SEM) |
|--------------|--------------------------------------|--------------------------------------|--------------------------------------|
| \(tg_{26}\)  | 2.77 ± 0.55‡                         | 1.32 ± 0.51§                         | 2.48 ± 0.56‡                         |
| CBA/J        | 1.03 ± 0.17                          | 0.67 ± 0.42                          | 0.53 ± 0.10                          |

*\(W/L = \) vessel wall cross-sectional area/lumen cross-sectional area ratio measured at \(×200\).
‡\(P < 0.01\) as compared to CBA/J.
§\(P < 0.1\) as compared to CBA/J.

**Table 2.** Data Summary for SCID Bone Marrow–inoculated \(tg_{26}\) Females at Days 10, 12, and 14 of Gestation

| Day of gestation | No. of mice | Thy-1 tx* | No. of uNK cells (mean ± SEM) | No. of viable/total No. of implants | Percent viability in utero |
|------------------|-------------|-----------|-------------------------------|-------------------------------------|---------------------------|
| N K reconstituted† |             |           |                               |                                     |                           |
| 10               | 1           | no        | 54.2 ± 10.9‖                   | 8/9                                 | 88.8                      |
| 12               | 1           | yes       | 90.6 ± 14.2‖                   | 5/6                                 | 83.3                      |
| 14               | 2           | no        | 108.9 ± 20.3‖                  | 13/14                               | 92.9                      |
| 14               | 3           | yes       | 124.3 ± 26.3‖                  | 24/27                               | 88.8                      |
| N K nonreconstituted† |           |           |                               |                                     |                           |
| 10               | 1           | no        | 4.1 ± 1.9‖                     | 3/4                                 | 75.0                      |
| 12               | 1           | yes       | 5.0 ± 5.8‖                     | 7/10                                | 70.0                      |
| 14               | 1           | no        | 1.45 ± 1.2‖                    | 8/9                                 | 88.8                      |
| \(tg_{26}\)      |             |           |                               |                                     |                           |
| 10               | 6           | no        | 1.3 ± 1.2‖                     | 47/59                               | 79.7                      |
| 12               | 4           | no        | 0.2 ± 0.3‖                     | 25/41                               | 61.0                      |
| 14               | 4           | no        | 1.7 ± 1.2‖                     | 9/25                                | 36.0                      |
| CD1/CBA/J       |             |           |                               |                                     |                           |
| 10               | 11          | no        | 134.8 ± 12.4                   | 110/121                             | 90.9                      |
| 12               | 30          | no        | 161.4 ± 16.8                   | 358/371                             | 96.5                      |
| 14               | 13          | no        | 229.2 ± 12.2                   | 127/138                             | 92.0                      |

*Some SCID bone marrow was pretreated with \(\alpha\)-Thy-1 Ab before inoculation of \(tg_{26}\) females.
†The frequency of uNK cells, as detected by periodic acid-Schiff-reactive granules was counted using a 10 mm² grid, at \(×250\) for all implantation sites, using 10 median sections per implantation site, a minimum of 2 implantation sites per pregnancy.
‡As assessed by splenic effector lytic cell assay against \(51\)Cr-labeled YAC-1 targets.
§\(P < 0.01\) as compared to gestational day–matched pooled CBA/J + CD1 values.
mice (NK \textsuperscript{m} + A) were larger by 27-34\% than gestational age-matched homozygous tge26 mice (B). The NK \textsuperscript{m} + females had well-developed metrial glands (mg) and higher cellularity in the decidua (d), whereas metrial gland development was absent (mg-) and the decidua was edematous in tge26 females. When compared to control immune-competent females, NK \textsuperscript{m} + females had 40-56\% normal frequencies of uNK cells (arrows; C) whereas tge26 females had 0-3\% normal uNK cell frequencies (arrow; D). In the NK \textsuperscript{m} + group of females, uNK cells (arrows) were frequently found surrounding decidual blood vessels and occasionally found within the vessel lumens (E). (F) demonstrates the decidual vessel anomalies found in tge26 mice including thickened vessel walls (asterisks); A and B, bar = 1,000 \( \mu \)m; C and D, bar = 40 \( \mu \)m, E and F, bar = 100 \( \mu \)m; (A, B, and F) stained with hematoxylin and eosin; (C-E) stained with periodic acid-Schiff.

Uterine changes in pregnant reconstituted tge26. uNK cells were present in considerable numbers in the seven NK cell reconstituted females but not in the remaining three females. Implantation sites from two immune-competent strains of mice (CD1 and CBA/J) were used to establish the parameters for normal number, size, and granularity of uNK cells on each of days 10, 12, and 14 of gestation. The frequency of uNK cells was not significantly different between CD1 and CBA/J strains at any gestational day studied. Therefore, the values from these two strains were pooled and are referred to as “control” in Table 2. In non-manipulated tge26, the number of uNK cells were statistically <1\% of control at days 10, 12, and 14 of gestation (reference 8 and Table 2). When compared to control immune-competent females, day 10 NK cell reconstituted females had 40.2\%, day 12 females had 56.1\% and day 14 females had 49.8\% of normal frequencies of uNK cells (P < 0.01; Table 2 and Fig. 1). M any uNK cells in the reconstituted females were associated with the main decidual blood vessels (Fig. 1). In contrast, low numbers of uNK cells (1-3.1\% of immune-competent females, which is also equivalent to the number of uNK cells found in untreated tge26 females) were present in the three bone marrow infused, nonreconstituted tge26 females (Table 2).

Histologically, the placenta from the NK cell reconstituted females appeared larger than those in tge26 females and the implantation sites resembled those in immune-competent females, because they included a well-developed metrial gland (Fig. 1). Reference standards for placental cross-sectional areas were obtained by measurements of CD1 and CBA/J implantation sites. Since these were not statistically different from each other at the gestational days used (10, 12, and 14), the data were pooled to provide a larger immune competent control data set than that used for control reference values in our earlier publication (8). The mean cross-sectional areas of unmanipulated tge26 placentae were 53\% of the mean at day 10 (P < 0.01, n = 6 mice), 50\% of the mean at day 12 (P < 0.01, n = 4 mice) and 47\% of the mean at day 14 (P < 0.01, n = 4 mice) control placentae (Fig. 2).

Placental cross-sectional areas for the NK cell-reconstituted females were significantly larger than cross-sectional areas measured for time-matched homozygous tge26 females or for SCID bone marrow-infused females who failed to show evidence for engraftment (Fig. 2). Placentae from all NK cell reconstituted tge26 mothers (independent

\( \text{mean percentage of specific lysis of 36.2\%} \pm 3.7 \) for unmanipulated tge26 at the same E:T ratio. Three females who had relatively nonlytic spleen cells (mean percentage of specific lysis of 9.1\% \pm 2.0 \) at E:T = 100:1 compared to mean percentage of specific lysis of 4.8\% \pm 3.7 for unmanipulated tge26 at the same E:T) were considered to be non-reconstituted, and were analyzed separately from the first seven females.

References standards for placental cross-sectional areas were obtained by measurements of CD1 and CBA/J implantation sites. Since these were not statistically different from each other at the gestational days used (10, 12, and 14), the data were pooled to provide a larger immune competent control data set than that used for control reference values in our earlier publication (8). The mean cross-sectional areas of unmanipulated tge26 placentae were 53\% of the mean at day 10 (P < 0.01, n = 6 mice), 50\% of the mean at day 12 (P < 0.01, n = 4 mice) and 47\% of the mean at day 14 (P < 0.01, n = 4 mice) control placentae (Fig. 2).

Placental cross-sectional areas for the NK cell-reconstituted females were significantly larger than cross-sectional areas measured for time-matched homozygous tge26 females or for SCID bone marrow-infused females who failed to show evidence for engraftment (Fig. 2). Placentae from all NK cell reconstituted tge26 mothers (independent
of the marrow graft preparation protocol) were 28% (day 10), 27% (day 12), and 34% (day 14) larger than placentae from homozygous tg26 mice (P < 0.01) but were 26% (day 10), 32% (day 12), and 29% (day 14) smaller than placentae from the immune-competent control mice (P < 0.01). Days 10, 12, and 14 placental cross-sectional area measurements from bone marrow-infused, nonreconstituted mice were not statistically different than homoygous tg26 placentaes. Day 10 and day 12 placentaes were measured at ×160 magnification, whereas day 14 placentaes were measured at ×140 magnification.

The vascular anomalies found in unmanipulated pregnant tg26 mice may or may not underlie the limited placental growth seen in these pregnancies. In the NK cell-reconstituted tg26 females, no major anomalies of the decidua or placental vasculature were found (Fig. 1). The lumen size in decidual vessels of the NK cell-reconstituted females was larger than that in nonreconstituted tg26 mice and appeared to increase with gestational length. The wall/lumen ratios were similar to those in the immune-competent controls (Tables 1 and 3), suggesting that engraftment had restored normal dilation of the decidual arterioles. This occurred in spite of the observation that uNK cell numbers were lower than in immune-competent mice. Minor variations from the immune-competent controls were found in implantation sites from only two of the seven NK cell-reconstituted females, the two females with the lowest uNK cell numbers (38.7 and 42.9% of immune-competent controls, respectively), and this suggests a redundancy in the uNK cell numbers regarding this trait. In these two females, the decidual region appeared fluid filled, the width of the myometrial smooth muscle was increased on the mesometrial side with uNK cells scattered within these smooth muscle fibers, and a small developing metrial gland was present. The trophoblast giant cell layer appeared mildly disorganized, having produced isolated islands of trophoblast giant cells, and trophoblast giant cells were found in apposition to decidual vessels. This is the location that many uNK cells occupy in mice. Blood vessels and placentae from bone marrow-infused but NK cell-nonreconstituted tg26 females resembled those from homoygously mated, nonmanipulated tg26 females (Tables 1 and 3).

Viability of the fetuses from the NK cell-reconstituted tg26 females was equivalent to fetal viability in immune-competent mice at all gestational days studied (Table 2). Viability of the fetuses from the single time-matched pregnancies available in bone marrow-infused, nonreconstituted females was below control levels at days 10 and 12 of gestation but, unexpectedly, equivalent to control in the one day 14 female, for which we have no explanation. Thus, reestablishment of 40–56% of uNK cells restored vasodilation in decidual arteries, promoted placental size, and promoted fetal viability.

One interpretation of the tg26 experiments is that uNK cells downregulate vascular smooth muscle cell (VSMC) numbers during pregnancy or promote VSMC relaxation. IFN-γ inhibits proliferation of rat VSMC through the induction of nitric oxide synthase (NOS) activity which in

| Table 3. Summary of Ratios of Mean Decidual Vessel Wall to Lumen Cross-sectional Area Measured at Days 10, 12, and 14 of Gestation |
|-----------------|-----------------|-----------------|-----------------|
| Mice strain     | Day 10 of gestation (mean W/L ± SEM) | Day 12 of gestation (mean W/L ± SEM) | Day 14 of gestation (mean W/L ± SEM) |
| N K +‡          | 1.27 ± 0.32    | 0.44 ± 0.27    | 0.68 ± 0.12    |
| N K −†          | 2.51 ± 0.41    | 1.21 ± 0.39    | 2.13 ± 0.37    |

* W/L = vessel wall cross-sectional area/lumen cross-sectional area ratio measured at ×200 magnification.
† N K − = three N K cell nonreconstituted tg26 females. N K + = seven N K cell reconstituted tg26 females.
‡ N ot statistically different than CBA/J.
† P < 0.01 as compared to CBA/J.
§ P < 0.1 as compared to CBA/J.
turn generates NO (18). NO is a major vasodilator that acts on vascular smooth muscle in many sites, including the uterus (19, 20). Infusion of an inhibitor of NO Ss during pregnancy in rats caused hypertension and fetal growth retardation, suggesting that a reduction in the synthesis of NO may contribute to the pathogenesis of pregnancy-induced hypertension (21). IFN-γ and iNOS are both products of mouse uNK cells (7, 22), as is TNF-α (23), another hypotensive molecule (24, 25). A number of histologists have noted a strong association of rodent uNK cells with blood vessels, and this association was reestablished in bone marrow–reconstituted tge26 females (Fig. 1 F). It has been previously suggested that uNK cells are involved in blood pressure regulation because hypertensive rats in which an artificial deciduomata had been induced experienced a fall of blood pressure at the time uNK cells appeared and an elevation of blood pressure when uNK cells disappeared despite persistence of the deciduomata (26). However, preliminary experiments involving serial blood pressure recordings by tail cuff in four conscious pregnant tge26 females failed to demonstrate statistically significant systemic hypertension over the first 16 d of gestation in comparison to four nonpregnant tge26 females (Luross, J.A., and B.A. Croy, unpublished data).

Three explanations have been considered for incomplete uNK cell reconstitution in tge26 females. First, the number of bone marrow cells or the proportion NK cell progenitors within the bone marrow suspension used for treatment in this study may not have been sufficient for full reconstitution of the uNK cell population. Since bone marrow was pooled from four donors for each recipient, this is not a strong argument. Second, the 3-wk time interval between the time of bone marrow treatment and the time of mating, in addition to the 10–14 d of pregnancy may have been too long, and thus bypassed the optimal interval (3 wk) when splenic NK cell numbers peak in tge26 mice reconstituted with recombination activating gene 2 (RAG-2) null bone marrow (13). Third, the 5-FU pretreatment may not have displaced the recipient marrow cells sufficiently to provide an adequate niche for optimal donor marrow engraftment.

The decidual blood vessel structure and placental morphology were improved or fully restored in the seven reconstituted tge26 mice despite their lower than normal frequency of uNK cells. Therefore, the placental size increments achieved in the N K cell–reconstituted tge26 versus nonmanipulated tge26 may represent the maximal growth promotion by uNK cells with other mechanisms, accounting for the deficit in size between reconstituted tge26 and immune-competent controls. One further requirement to promote full placental growth may be immune-competence in the fetal compartment. We have reported elsewhere that embryo transfers demonstrate that unilateral fetal or maternal NK cell deficiency can reduce placental size (8).

These in vivo studies have better defined the functional role of the immune system and the role of uNK cells in the pregnant uterus in particular. They exclude T cells and NK1 T cells as essential participants in pregnancy success, supporting earlier studies of nu/nu and IL-2 null × β2m null mice (6–8). These studies identify the NK cell lineage as important for normal development of implantation sites and highlight the mesometrial decidua and vasculature as important sites for uNK cell function. These studies do not support the previously postulated function of uNK cells to limiting trophoblast cell growth and invasion. On the contrary, they show that maternal uNK cells promote placental growth and thereby the growth of the developing fetus. It remains unclear whether promotion of trophoblast growth is a direct or indirect effect.

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