Natural killer cytotoxicity and antibody-dependent cytotoxicity of cells of rat metrial glands

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Cell suspensions of rat metrial gland tissue, containing healthy granulated metrial gland (GMG) cells, were assessed for their ability to lyse natural killer (NK) cell target Yac-1 myeloma cells in a $[^{51}Cr]$-release cytotoxicity assay. Cytotoxicity indices were found to be low even after in vivo stimulation of NK cytotoxic activity by polyinosinic–cytidilic acid and the addition of interferon to test cultures. It is concluded that cell suspensions of rat metrial gland, and rat GMG cells in particular, do not exhibit NK cytotoxicity. However, when a heat inactivated rabbit anti-Yac-1 serum was added to assay cultures, cytotoxicity indices above background levels were obtained, indicating that rat metrial glands contain cells capable of antibody-dependent cytotoxicity.

Introduction

Granulated metrial gland (GMG) cells are pregnancy-specific, bone-marrow derived cells that infiltrate the deciduized endometrium of rodents and have a phenotype suggesting that they are a type of natural killer (NK) cell (Stewart, 1991). Studies of interactions between GMG cells and trophoblast in vivo (Stewart, 1984, 1990) and in vitro (Stewart and Mukhtar, 1988) have indicated that GMG cells kill a small proportion of the trophoblast cells that line the maternal blood spaces of the labyrinthine placenta. However, attempts to show that mouse GMG cells function as NK cells, as defined by their ability to kill specific target cells such as Yac-1 myeloma cells (Trinchieri, 1989), have had little success. One reason for this is the difficulty in obtaining single-cell suspensions of metrial gland cells containing high numbers of viable and normal GMG cells for use in standard chromium-release cytotoxicity assays. It is possible to obtain a relatively pure population of GMG cells by allowing their migration from cultured explants of mouse metrial gland tissue (Mukhtar and Stewart, 1988), but the number of cells obtained is usually insufficient to carry out a full cytotoxicity assay using a range of effector:target ratios and multiple replicates. Low levels of killing have been achieved using migratory GMG cells from cultured mouse deciduall explants but only after the cells were subjected to prolonged treatment with interleukin 2 (Linnemeyer and Pollack, 1991).

In contrast to the mouse, it is possible to prepare single-cell suspensions of rat metrial gland tissue with numerous GMG cells (Bray et al., 1978), although the percentage is still lower than found in vivo (Matthews, 1985), and in large enough quantities to provide sufficient cells for a full cytotoxicity assay. Rat GMG cells have been shown to be cytotoxic to rat and mouse trophoblast cells in vitro (Peel and Adam, 1991) and, in xenogeneic chimaeras, to mouse trophoblast in vivo (Peel and Stewart, 1989), but their ability to kill NK-specific target cells has not been assessed. In the present study, we attempted to determine whether cells from rat metrial glands could kill the NK-target cell Yac-1 using a chromium-release assay. Metrial glands from rats at day 10 of pregnancy which contain GMG cell precursors and immature GMG cells, and metrial glands from day 14 pregnant rats which contain mature GMG cells (Peel, 1989) were used.

An enzyme digestion procedure was used to provide the single cell suspensions of rat metrial gland tissue. In some cases rats were given an injection of polyinosinic–cytidilic acid (poly-IC) to enhance any NK-cell activity (Trinchieri, 1989). Interferon-γ (IFN-γ) was added to some of the assay cultures from the poly-IC-treated rats to continue the stimulation of NK-cell activity initiated in vivo by the poly-IC. Particular attention was given to control material and to the effects of the preparation method for obtaining single cell suspensions of metrial gland tissue on the NK assay. Suspensions of rat metrial gland cells were examined histologically to ensure that the effector cell populations contained viable GMG cells. In addition, to determine whether metrial gland cell suspensions contained cells that could initiate antibody-dependent cytotoxicity, an antisemum to Yac-1 myeloma cells was raised in rabbits and added to some of the cytotoxicity test cultures.

Materials and Methods

Animals

Virgin female Wistar rats, aged 12–15 weeks, were mated overnight with males of the same strain, and the presence of a vaginal plug on the cage floor the following morning was taken as evidence of pregnancy (day 0). Some rats were given a single i.p. injection of 500 μg poly-IC (Sigma) in 0.5 ml phosphate-buffered saline 24 h before they were killed (by cervical dislocation under ether anaesthesia) on day 10 or day 14 of pregnancy.
**Single-cell suspensions of metrial gland tissues**

Metrial glands from normal implantation sites were dissected from the uterine wall and cut into pieces using a surgical blade. Any small for age, or resorbing implantation sites, were discarded. The pieces of metrial gland were digested in 10 ml of Hank’s Balanced Salt Solution (HBSS; Flow Laboratories, Irvine) containing 0.125% collagenase (Type I; Sigma, Poole, UK), 0.1% trypsin (Type II; Sigma), 100 IU penicillin ml⁻¹ and 100 µg streptomycin ml⁻¹ for 1 h at 37°C under constant stirring. After digestion, the cell suspension was filtered through a 100 mesh wire gauze, washed three times by centrifugation at 150 g for 5 min and resuspended in Minimum Essential Medium (Eagle) with Earle’s salts (Flow Laboratories) supplemented with 10% fetal calf serum (Seralab, Sussex), 0.058% L-glutamine, 100 IU penicillin ml⁻¹ and 100 µg streptomycin ml⁻¹ (MEM). Cells were counted and their viability assessed by trypan blue exclusion. Only cell suspensions with a viability greater than 90% were used. The cell concentration was adjusted to 10⁶ viable cells ml⁻¹ and the cells kept on ice until required.

**Chromium release assay**

Yac-1 myeloma cells (2 x 10⁷ viable cells) in 1 ml MEM, from a culture growing in logarithmic phase, were added to 100 µl saline containing 1 mCi Na₂⁵¹CrO₄ (ICN Radiochemicals, High Wycombe) and maintained in culture for 1 h at 37°C. The cells were washed three times by centrifugation (150 g, 5 min) and resuspension in MEM and their concentration adjusted to 5 x 10⁶ viable cells ml⁻¹ for use as target cells.

**Histology of metrial gland cell suspensions**

Excess cells from the metrial gland cell suspensions, when available, were fixed in a mixture of 2% glutaraldehyde and 4% formaldehyde in 0.1 mol phosphate buffer l⁻¹, pH 7.2-7.4 (modified from Karnovsky, 1965). The cells were pelleted, supported in 1% agar, washed in 0.1 mol phosphate buffer l⁻¹, dehydrated in ethanol and embedded in glycol methacrylate (Rudell, 197). Sections (1-2 µm) were cut, reacted with the Periodic acid–Schiffs reagent with or without prior diastase digestion, and counterstained with haematoxylin. Cell counts were made of nucleated cells that were categorized as GMG cells, lymphocytes/lymphocyte-like cells (including GMG precursor cells) or other leucocytes and stromal cells (this group would include endothelial cells or smooth muscle cells). Only cells with an apparently normal nucleus were counted. Only cells with a nucleus and at least one of the typical cytoplasmic

**Antibody-dependent cytotoxicity**

A New Zealand white rabbit was given five s.c. injections, each of approximately 5 x 10⁷ lyed Yac-1 myeloma cells in Freund’s complete adjuvant, over 6 months. Some immune serum was heat inactivated at 60°C for 1 h. Immune activity of the serum was tested in a ⁵¹Cr-release assay in which release was determined from viable Yac-1 cells incubated in the presence of the immune serum at various concentrations and in the presence of heat-inactivated serum with or without normal non-immune rabbit serum. In these test assays, maximal lysis of Yac-1 cells was obtained when cells were incubated in heat-inactivated immune serum at concentrations of 1 in 100, or higher, in the presence of normal non-immune serum. When required for experimental purposes aliquots of Yac-1 cells were added to MEM containing heat-inactivated anti-Yac-1 serum diluted to 1 in 10, giving a final concentration of 1 in 100 in the culture wells for the chromium-release assays. Otherwise, chromium-release assays to determine antibody-dependent cytotoxicity were carried out as described above.

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glycoprotein-containing granules were categorized as GMG cells. At least 200 cells were counted from each cell suspension.

**Results**

**Metrial gland cell cytotoxicity**

Most of the metrial gland cell suspensions obtained from rats at day 10 or day 14 of pregnancy killed fewer than 5% Yac-1 cells (Fig. 1a, b). In only two rats (one at day 10 and one at day 14) were cytotoxicity indices above 10% obtained at an E:T ratio of 100:1 (Fig. 1a, b).

When animals were given an injection of poly-IC 24 h before death to increase NK cytotoxicity by IFN stimulation, cell suspensions of their metrial glands still failed to kill Yac-1 cells (Fig. 1c, d). Even under the additional stimulation of *in vitro* IFN, cytotoxicity levels were unchanged (Fig. 1e, f).

**Spleen cell cytotoxicity**

Spleen cell suspensions assayed in parallel with the metrial gland cell suspensions gave a high cytotoxicity index at an E:T ratio of 200:1 that decreased with each successive reduction in E:T ratio (Fig. 1a, b). At an E:T ratio of 100:1, spleen cell suspensions from control rats at day 10 of pregnancy gave a mean cytotoxicity index of 33% (Fig. 1a). With spleen cells from rats at day 10 of pregnancy, that had been given an injection of poly-IC (Fig. 1c), higher cytotoxicity indices were obtained (56% at an E:T ratio of 100:1). When assay cultures from poly-IC-stimulated rats at day 10 of pregnancy were supplemented with IFN there was no change in cytotoxicity indices (Fig. 1e). Similar responses were obtained from spleen cell suspensions prepared from rats at day 14 of pregnancy (Fig. 1b, d, f).

**Effects of cell preparation methods**

Metrial gland cell suspensions for use in test assays were prepared only using collagenase–trypsin digestion at 37°C. No differences were found between the cytotoxicity indices of spleen cell aliquots that had been exposed to the various cell preparation procedures used (4°C, 37°C and collagenase–trypsin at 37°C) indicating that collagenase–trypsin digestion did not adversely affect cytotoxic activity.

**Determination of antibody-dependent cytotoxicity**

Cytotoxicity indices were higher when aliquots of Yac-1 cells were added to medium containing heat-inactivated rabbit anti-Yac-1 serum than in parallel cultures in which the anti-Yac-1 serum was absent from the culture medium and replaced with medium free of rabbit serum or containing non-immune rabbit serum (Fig. 2). Even when the normal metrial gland cell suspensions failed to kill Yac-1 cells (the majority of experiments) a positive cytotoxicity index was obtained in the presence of anti-Yac-1 serum (Fig. 2). In control cultures where Yac-1 myeloma cells were incubated in the absence of metrial gland effector cells there were no differences between the chromium-release counts when anti-Yac-1 serum was added or omitted from the culture medium. The addition of anti-Yac-1 serum to wells with spleen cell effectors did not affect the level of cytotoxicity.

**Histological assessment of metrial gland cell suspensions**

All samples of metrial gland cell suspensions prepared for examination contained GMG cells that appeared healthy and contained the typical glycoprotein-containing cytoplasmic granules (Fig. 3). Lymphocytes and lymphocyte-like precursor cells were also found as well as other leucocytes and stromal cells (Table 1).
Fig. 2. Cytotoxicity indices of metrial gland cell suspensions against Yac-1 cells for individual rats at day 14 of pregnancy. Three replicates were performed at each effector:target ratio shown. (□) Cells tested in MEM containing normal rabbit serum; (■) cells tested in MEM alone; (○) cells tested in MEM containing heat inactivated rabbit anti-Yac-1 serum.

Fig. 3. Granulated metrial gland cells (arrows) and stromal cells (S) in a rat metrial gland cell suspension fixed after aliquots of cells had been taken for incorporation into chromium-release assay test cultures. Diastase, periodic acid–Schiff and haematoxylin. Bar = 10 μm.

Discussion

Suspensions of rat metrial gland cells containing apparently healthy GMG cells generally failed to kill Yac-1 myeloma cells. Spleen cell suspensions, including some that had been subjected to the same enzyme digestion regimen as the metrial gland cells, gave consistently high levels of cytotoxicity at high E:T ratios and showed a regular decline in cytotoxicity with reducing E:T ratios. These results show that metrial glands do not contain cells with natural killer cell activity as defined by the capacity to kill Yac-1 myeloma cells. Numerous studies have shown that cells in rodent (Clark et al., 1990) and human decidua (Lala et al., 1990) can produce factors that can inhibit NK activity. The metrial gland may be considered as an extension of the decidua basalis (Stewart, 1993) and may have similar functional activities, including the release of suppressor factors that inhibit NK activity. It is possible, therefore, that NK activity in crude cell suspensions of metrial glands could be similarly affected. However, metrial gland cell suspensions from rats treated with poly-IC did not show any increased cytotoxicity to Yac-1 cells, even when the assay was performed in the presence of interferon, suggesting that suppression of any cytotoxic potential of metrial gland cells, including GMG cells, did not occur.

Immunohistochemical studies have shown that rat and mouse GMG cells express the asialo-GM1 antigen (Mukhtar et al., 1989; Redline and Lu, 1989; Peel and Adam, 1991) which characterizes NK cells. More extensive immunohistochemical studies on mouse GMG cells have shown that they express other surface antigens that are also characteristic of the NK cell phenotype such as Fcy receptors (Daki et al., 1989), Thy-1 (Mukhtar et al., 1989; Redline and Lu, 1989) and perforin (Parr et al., 1987, 1990a). Rat GMG cells, however, do not express Thy-1 (Mitchell and Peel, 1984; Peel and Adam, 1991) nor Fcy receptors (Bray et al., 1978); no studies to detect perforin in rat GMG cells have been reported. Cells analogous to rodent GMG cells in the human endometrium, endometrial stromal granulocytes, express the NK cell antigen CD56 but not other NK antigens such as CD16 (Bulmer et al., 1991). Human endometrial stromal granulocytes are reported to be cytotoxic to the target cell K562 used to define human NK cells but at a lower level of cytotoxicity than peripheral blood mononuclear-cell isolates (Ferry et al., 1990). Indeed, it has been proposed that the NK-cytotoxicity activity of decidual-cell isolates, enriched with endometrial stromal granulocytes, may be due to a minor CD16+ population of CD56+ cells (Christmas et al., 1990) which could be blood contaminants rather than tissue endometrial stromal granulocytes (Stewart, 1991). It is well established that large numbers of viable mouse GMG cells are difficult to isolate. Parr et al. (1990b), using single-cell suspensions obtained by mechanical and enzymatic digestion, found no evidence of natural cytotoxicity, although the GMG cells used...
in their study were lacking the normal cytoplasmic granules found in GMG cells. Croy et al. (1991) and Linnemeyer and Pollack (1991) used GMG cells isolated from cultured metrial gland or decidual explants but these cells failed to kill Yac-1 cells unless they were treated with interleukin 2. An alternative hypothesis to GMG cells being a type of NK cell is that they differentiate from NK cells (Parr et al. 1991) have recently suggested that mouse GMG cells differentiate from LGL-1 positive NK cells but expression of the LGL-1 antigen is lost with differentiation), and are, therefore, members of the NK-cell lineage without necessarily retaining all the functions of NK cells. Head (1990) has reported that cells in the developing rat metrial gland at day 10 of pregnancy express the NK activation marker 3.2.3, although this antigen was not expressed by mature GMG cells. Our studies included the assessment of NK cytotoxic activity of cell suspensions prepared from metrial glands obtained from rats killed at day 10 of pregnancy. At this time GMG cells, as well as many lymphocyte-like cells and potential precursor GMG cells, are present but evidence of NK cytotoxicity was not found even after interferon treatment. We have, therefore, found no evidence that rat metrial glands contain cells capable of NK cytotoxicity.

It is possible that the immunohistochemical studies that identified NK cell associated antigens on GMG cells may have been misleading and that these antigens are expressed by a wider range of cell populations, not all of which are cytotoxic for the classic targets. There is evidence that rat GMG cells can kill rat trophoblast cells in vitro (Peel and Adam, 1991) and mouse GMG cells can kill mouse trophoblast cells in vitro (Stewart and Mukhtar, 1988) and in vivo (Stewart, 1984, 1990). In contrast, normal mouse trophoblast cells in vitro are not susceptible to lysis by cytotoxic T cells or splenic NK cells (Drake and Head, 1989a, 1989b). A more broad-based approach to studying the cytotoxic potential of GMG cells is required if the mechanism by which the killing process takes place is to be determined and the function of GMG cells in pregnancy success or failure established. Our finding that single-cell suspensions of rat metrial gland tissue express antibody-dependent cytotoxicity provides the basis for a renewed approach. It will be of interest to determine whether it is the GMG cells, or other cells, such as macrophages, that form a minority population in the rat metrial gland (Peel, 1989), in the cell suspensions that are responsible for the antibody-dependent cytotoxicity.

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**Table 1. Percentage (mean ± SEM) of granulated metrial gland (GMG) cells and other cell types in cell suspensions of rat metrial gland**

<table>
<thead>
<tr>
<th>Day of pregnancy</th>
<th>Number of rats</th>
<th>GMG cells</th>
<th>Lymphocyte (-like) cells</th>
<th>Other leucocytes</th>
<th>Stromal cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>4</td>
<td>4.7 ± 1.2</td>
<td>8.4 ± 1.2</td>
<td>0.6 ± 0.4</td>
<td>86.3 ± 1.4</td>
</tr>
<tr>
<td>14</td>
<td>6</td>
<td>7.4 ± 1.3</td>
<td>5.9 ± 1.5</td>
<td>2.3 ± 0.9</td>
<td>84.0 ± 1.4</td>
</tr>
</tbody>
</table>

Data include counts from cell suspensions prepared from rats receiving or not receiving an injection of polyinosinic–cytidilic acid. Cell counts were made from plastic sections.

References


Christmas SE, Bulmer JN, Meager A and Johnson PM (1990) Phenotypic and functional analysis of human CD3- decidual leucocyte clones Immunology 71 182–189


Drake BL and Head JR (1989a) Murine trophoblast can be killed by allospecific cytotoxic T lymphocytes generated in CIBCO Opti-MEM medium Journal of Reproductive Immunology 15 71–77

Drake BL and Head JR (1989b) Murine trophoblast can be killed by lymphokine-activated killer cells Journal of Immunology 143 9–14

Ferry BL, Starkey PM, Sargent JL, Watt GMO, Jackson M and Redman CWG (1990) Cell populations in the human early pregnancy decidua: natural killer activity and response to interleukin-2 of CD56-positive large granular lymphocytes Immunology 70 440–452

Head JR (1990) Distribution of natural killer (NK) cells in the pregnant rat uterus American Journal of Reproductive Immunology 22 74


Lala PK, Srodras JM, Graham CH, Lynjak JJ and Parhar RS (1990) Activation of maternal killer cells in the pregnant uterus with chronic indomethacin therapy, IL-2 therapy, or a combination therapy is associated with embryonic demise Cellular Immunology 127 368–381

Linnemeyer PA and Pollack SB (1991) Murine granulated metrial gland cells at uterine implantation sites are natural killer lineage cells Journal of Immunology 147 2530–2535


Mitchell BS and Peel S (1984) Identification of cells bearing leucocyte surface antigens in metrial gland tissue from rats of different gestational ages, strains or parities Immunology 53 63–68

Mukhtar DDDY and Stewart JI (1988) Migration of granulated metrial gland cells from cultured explants of mouse metrial gland tissue Cell and Tissue Research 253 413–417


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Parr EL, Young LHY, Parr MB and Young JD-E (1990a) Granulated metrial gland cells of pregnant mouse uterus are natural killer-like cells that contain perforin and serine esterases Journal of Immunology 145 2365–2372


Peel S and Stewart I (1989) Rat granulated metrial gland cells differentiate in pregnant chimeric mice and may be cytotoxic for mouse trophoblast Cell Differentiation and Development 28 55–64


Trinchieri G (1989) Biology of natural killer cells Advances in Immunology 47 187–376