Intercellular Transfer of a Soluble Viral Superantigen

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Mouse mammary tumor virus (MMTV) superantigens (vSAgs) can undergo intercellular transfer in vivo and in vitro such that a vSAg can be presented to T cells by major histocompatibility complex (MHC) class II proteins on antigen-presenting cells (APCs) that do not express the superantigen. This process may allow T-cell activation to occur prior to viral infection. Consistent with these findings, vSAg produced by Chinese hamster ovary (CHO) cells was readily transferred to class II IE and IA (H-2k and H-2d) proteins on a B-cell lymphoma or mouse splenocytes. Fixed class II-expressing acceptor cells were used to demonstrate that the vSAg, but not the class II proteins, underwent intercellular transfer, indicating that vSAg binding to class II MHC could occur directly at the cell surface. Intercellular transfer also occurred efficiently to splenocytes from endogenous retrovirus-free mice, indicating that other proviral proteins were not involved. Presentation of vSAg produced by a class II-negative, furin protease-deficient CHO variant (FD11) was unsuccessful, indicating that proteolytic processing was a requisite event and that proteolytic activity could not be provided by an endoprotease on the acceptor APC. Furthermore, vSAg presentation was effected using cell-free supernatant from class II-negative, vSAg-positive cells, indicating that a soluble molecule, most likely produced by proteolytic processing, was sufficient to stimulate T cells. Because the membrane-proximal endoproteolytic cleavage site in the vSAg (residues 68 to 71) was not necessary for intercellular transfer, the data support the notion that the carboxy-terminal endoproteolytic cleavage product is an active vSAg moiety.

Activation of T cells by mouse mammary tumor virus (MMTV) superantigens (vSAgs) is essential for viral transmission (for a review see reference 1). This activation is mediated via interaction of the vSAgs with class II major histocompatibility complex (MHC) proteins on antigen-presenting cells (APCs) and the variable region of the β chain of the T-cell receptor. The vSAgs are produced as glycosylated type II integral membrane proteins that require endoproteolytic maturation to activate T cells (11). In CHO cells, proteolytic processing is effected by furin, a member of a family of endoproteases known as protein convertases (PCs) (17), and results in the generation of one or more proteolytic products. An 18-kDa carboxy-terminal proteolytic cleavage product (p18) has been demonstrated to associate on the cell surface of B cells both with an amino-terminal vSAg proteolytic cleavage product and with the class II MHC protein IAβ (23, 24). Although similar in function to the well-characterized bacterial superantigens, the vSAgs and the bacterial SAgs bear no genetic resemblance, and the structure of the vSAgs and details of their interactions with the class II MHC proteins have not been resolved.

Another feature of vSAgs is their capacity to undergo intercellular transfer. vSAgs will not stimulate T cells in the absence of class II MHC proteins (2), and vSAg intercellular transfer was first observed in mixed bone marrow reconstituted chimeric mice where the donor cells expressed separately a proviral SAg or an appropriate class II MHC protein (14, 19). Intercellular vSAg transfer was evidenced by the ability of the vSAg expressed in the mixed bone marrow chimeric mice to effect intrathymic deletion of reactive T cells. Evidence that vSAg-expressing CD8 T cells, which do not express class II proteins, could induce T-cell deletion in vivo also provided evidence for intercellular transfer (21). Intercellular transfer has also been demonstrated to occur in vitro by coculture of vSAg-reactive T cells with mixtures of independently transfected vSAg and class II MHC-expressing APCs (4). Intercellular transfer of the vSAgs may act during infection by exogenous virus to allow activation of T cells independent of or prior to infection of B cells and/or other APCs.

The vSAgs are integral membrane proteins and therefore unlikely to undergo intercellular transfer in the absence of posttranslational processing that would eliminate membrane tethering. An appealing hypothesis has been that vSAg intercellular transfer is facilitated by the proteolytic generation of a soluble vSAg protein (e.g., p18) (24, 25). Evidence for intercellular transfer of such a moiety would indicate that the regions necessary for interaction of the vSAg with the MHC proteins and the T-cell receptor reside therein. Here, we both confirm and extend a previous study that has demonstrated intercellular transfer of a soluble vSAg in vitro. We provide a formal demonstration that the vSAg, but not class II MHC, undergoes intercellular transfer, and that vSAg binding to class II MHC occurs at the cell surface. It is also shown that intercellular transfer requires furin-dependent proteolytic processing. The data suggest that all regions required for superantigen activity, including the interaction with both the class II MHC proteins and the T-cell receptor, reside on a vSAg carboxy-terminal proteolytic fragment.

MATERIALS AND METHODS

Animals. Inbred mice used as a source of splenocytes were bred at the Wadsworth Center, NYS Department of Health, under institutional guidelines for animal care and use. The MMTV-negative mice (16) were obtained from the laboratory of Philippa Marrack and John Kappler at the Howard Hughes Medical Institute, Denver, Colo.

Cell lines and APCs. All cells were cultured in complete tumor medium as described previously (11). Transfection, drug selection, T-cell stimulation assays, and flow cytometry were also performed as described previously (11). All of the T-cell hybrids and most APCs used in this study have been described previously (11). The CHO/S7 and FD/S7 vSAg7 transfectants were generated by transfection of CHO and FD11 cells, respectively, with the vSAg7 expression

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plasmid pSRoSAg7, as described previously (11). Splenocytes were obtained after passage of tissue through a 100-μm nylon mesh, and erythrocytes were removed using 150 mM ammonium chloride–0.73 mM potassium phosphate. T cells were depleted using the anti-Thy1.2 antibody H35.7 and rabbit complement; the remaining splenocytes were washed with Hanks balanced salt solution (HBSS) and then treated with mitomycin C at a final concentration of 25 μg/ml in HBSS. Cells were fixed by incubation in 0.05% glutaraldehyde in HBSS for 30 s, 1 volume of 0.2 M lysine in HBSS was added immediately, and the cells were washed in HBSS.

**RESULTS**

**Intercellular transfer of vSAg7.** To assess superantigen intercellular transfer, vSAg7 (derived from the MMTV-7 provirus) was expressed in class II-negative CHO (CHO/S7) cells. The vSAg7 transfectants (vSAg donor cells) were incubated with the class II-positive B-cell lymphoma CH12.1 (acceptor APC) and the vSAg7-reactive T-cell hybridomas Oms1.24.6 and Kmls13.11 (responder cells; both Vβ6 [11]). The T cells responded readily to vSAg7 presented in the three-cell culture but did not respond to the vSAg7 transfectants in the absence of the class II-positive acceptor APCs (Fig. 1). Although vSAg7 is expressed at relatively low levels on the CHO transfectants (Fig. 2a), intercellular transfer was nevertheless quite efficient, as IL-2 production engendered in the three-cell culture did not allow presentation by nonfixed CH12 cells (Fig. 4b). Fixation of the vSAg-expressing cells (Fig. 4) in contrast, facilitated intercellular transfer and that class II MHC binding demonstrated that the vSAg, but not the class II protein, underwent intercellular transfer and that class II MHC binding occurred directly at the cell surface of the APC.

To further evaluate the role of class II MHC during intercellular transfer and to test the capacity of normal APCs to accept vSAg7 transfer, the vSAg7 donor cells were incubated with vSAg7-negative, class II-positive spleen cells. Splenocytes from BALB/c (H-2b), B10.A (H-2d), B10.D2 (H-2d), and the MHC recombinant strain B10.A(4R) (which expresses only IAa) all presented the transferred vSAg7 (Fig. 3b). Presentation by IEd was significantly better than that by both IAa [Fig. 3b, compare B10.A with B10.A(4R)] and class II H-2d. These findings recapitulate the hierarchy of class II presentation of endogenously expressed vSAg7 by class II MHC (10, 14) and demonstrate that normal APCs can present the transferred vSAg in vitro.

**vSAg, but not class II IEa, underwent intercellular transfer.** Previous in vivo and in vitro studies did not eliminate the formal possibility that the vSAg responses resulted from transfer of class II MHC proteins from the class II-expressing cells to the vSAg-expressing cells. To address this possibility, experiments were performed using vSAg7- or class II-presenting cells that had been fixed with glutaraldehyde. Significant vSAg intercellular transfer was observed when fixed class II-positive CH12 cells were incubated with nonfixed vSAg7-expressing cells (Fig. 4). In contrast, fixation of the vSAg-expressing cells did not allow presentation by nonfixed CH12 cells (Fig. 4b). Presentation of vSAg7 endogenously produced in CH12 cells was largely unaffected by fixation (Fig. 4c). These experiments demonstrated that the vSAg, but not the class II protein, underwent intercellular transfer and that class II MHC binding occurred directly at the cell surface of the APC.

**Intercellular transfer did not require other MMTV proteins.** Although transfer was readily observed with CH12.1 as the acceptor cell, transfer was not observed when the class II-positive CHO transfectant CHIE was used as an acceptor cell (data not shown). To address whether this may have reflected a requirement for other provirus-encoded MMTV proteins to facilitate intracellular transfer, experiments were performed using splenocytes obtained from a mammary tumor provirus-free mouse strain (16). Intercellular transfer of vSAg7 to the MMTV provirus-free splenocytes was observed (Fig. 5), indicating that additional MMTV proteins were not required for vSAg intercellular transfer. Similar levels of T-cell stimulation were detected using splenocytes obtained from the genetically related strain CBA/J (H-2h), which express endogenous vSAg7, suggesting that presentation of endogenous and transferred vSAg7 by the splenocytes was equally efficient. Further studies suggested that the failure of the CHIE cells to present the transferred vSAg was due to an overall lower efficiency of T-cell activation (data not shown).
were analyzed by flow cytometry for vSAg7 (dashed lines) and/or class II IEk expression. Four
68 to 71, was expressed in class II-negative CHO cells and vSAg7m2 (22), which lacks the PC processing site at positions
was a requisite step in vSAg7 transfer from CHO donor cells. (Fig. 6a and b). Thus, furin-dependent proteolytic processing
ferred vSAg from the furin-deficient class II-negative cells (Fig. 6a). Moreover, treatment of the furin-deficient cells with leupeptin,
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FIG. 2. Surface expression of vSAg7 and class II MHC proteins. Cell lines
expression observed in the CHIE cells (c) was not characteristic of this cell line.

Furin-dependent proteolytic processing was required for vSAg presentation. The PC furin has been demonstrated to
mediated endoproteolytic processing of vSAgs in vitro and in vivo (13, 11), and furin-dependent processing is required for
T-cell activation of vSAg7 in CHO cells (11). To determine if furin was also required for transfer and/or presentation of the
transferred vSAg, vSAg7 was expressed, in the absence of class II, in the furin-deficient CHO cell line FD11 (FD/S7) (Fig. 2)
(6). Activation of T cells upon transfer of vSAg from the furin-deficient cells was approximately 80-fold lower than that
obtained using the furin-positive CHO cells (Fig. 6a). Moreover, treatment of the furin-deficient cells with leupeptin,
which has previously been shown to abrogate the residual presentation of vSAg7 by the furin-deficient class II-positive trans-
fectant FDIE/S7, completely blocked the activity of the transferred vSAg from the furin-deficient class II-negative cells
(Fig. 6a and b). Thus, furin-dependent proteolytic processing was a requisite step in vSAg7 transfer from CHO donor cells.

Proteolytic processing of vSAg7 at positions 168 to 171 was shown to be required for vSAg activity when expressed in CHO
cells (22). In contrast, furin processing at the conserved mem-
brane-proximal cleavage site in vSAg7 (positions 68 to 71) was
found to be inessential for activation of T cells by class II-
positive APCs (22). Because the furin recognition site at positions
68 to 71 is, with one exception, conserved in all known
vSAgs (23), it was considered that proteolytic processing at this
position might be required for intercellular transfer, even
though it was not required for endogenous presentation. To
test this possibility, a previously described vSAg7 variant,
vSAg7m2 (22), which lacks the PC processing site at positions
68 to 71, was expressed in class II-negative CHO cells and examined for its ability to undergo intercellular transfer. Four

FIG. 3. Presentation of transferred vSAg7 by class II MHC. (a) II-2 produc-
tion by T-cell hybridomas was measured after culture with the vSAg7 donor cell
line CHO/S7 and the acceptor APC CH12.1 without antibodies (none) or in the
presence of the following antibodies, as indicated: anti-class II IEk (14-4-4),
anti-class II I4a (17/227 or 10.2-16), both anti-IE and anti-I4a (10.2.16), or
isotype-matched control antibodies (immunoglobulins G2a and G2b). (b) T-cell
hybridomas were incubated with the vSAg7 donor cell CHO/S7 and splenocytes
obtained from the following mouse strains: BALB/c (I4a I5b), B10.A (I4a I5b),
and B10.D2 (I4a I5d) B10.A(4R) (I4a). II-2 production was determined as for
Fig. 1.
the transferred vSAg have not yet been successful. These data nevertheless provide clear evidence that a soluble superantigen was transferred from the vSAg donor cells.

DISCUSSION

Although the vSAgs are synthesized as membrane-bound glycoproteins, this study demonstrates that a functional form of the vSAg can undergo intercellular transfer in vitro and thus confirms and extends the previous in vivo and in vitro studies that demonstrated vSAg intercellular transfer (4, 14, 19). In our studies, intercellular transfer occurred readily from vSAg7-expressing CHO cells to the B-cell lymphoma cells, and to normal spleen cells, and presentation of the transferred vSAg to T cells was inhibited, as expected, by MHC class II antibodies. Although the transferred vSAg was not detectable on the cell surface of the acceptor APCs, presentation to T cells was nevertheless quite efficient, because levels of IL-2 production in some cases approached that obtained when the vSAg was expressed endogenously. These data suggest that relatively few vSAg molecules can stimulate a strong T-cell response, much like that observed for conventional peptide antigens, where as few as 100 peptide molecules are sufficient for T-cell activation (8). The efficiency of intercellular transfer also suggests that vSAg intercellular transfer may be a common or even requisite event during viral infection and concomitant T-cell activation.

In the previous in vitro and in vivo studies, the possibility that intercellular transfer was due to transfer not of the vSAg but of class II proteins was not ruled out. This was addressed in the present studies using cell fixation. Glutaraldehyde-fixed APCs were capable of presenting to T cells vSAg that had been produced by unfixed vSAg7 donor cells. Moreover, vSAg intercellular transfer did not occur when the vSAg-expressing cells were fixed. Because transfer of class II proteins or vSAgs was unlikely to occur from fixed cells, the data indicate that the vSAg protein was the transferred moiety. Moreover, the presentation of vSAg7 by fixed APCs indicated that vSAg association with the class II proteins occurred at the cell surface. Thus, vSAg7-class II binding did not require endocytosis and presentation via the conventional class II antigen presentation pathway (3), and so this association did not require accessory molecules, such as H-2 DM, that are typically required for MHC presentation of conventional peptide ligands (12).

No differences were observed in the hierarchy of class II MHC presentation when vSAg7 was expressed endogenously or upon intercellular transfer. These data suggest that the mode of class II binding of the transferred vSAg is similar or identical to that of endogenously expressed vSAgs, and they demonstrate that vSAgs can bind to stable class II MHC proteins, as has been suggested previously (7).

vSAgs are detected on the cell surface of APCs in a processed form (25), and so it is likely that a proteolytic fragment of the vSAg undergoes intercellular transfer. Furin-deficient CHO cells, which do not express detectable processed vSAg7 (11), were poor vSAg7 donors, and when transfer experiments were performed in the presence of the protease inhibitor leupeptin, the residual transfer and/or presentation of vSAg7 by these cells was completely abolished. These findings suggest that proteolytic processing is required for intercellular transfer. However, one cannot rule out that in the absence of processing the vSAg undergoes intercellular transfer, but the transferred vSAg is not in a form that can be presented to T cells. However, the failure to observe presentation of the unprocessed vSAg indicates that furin or other PCs known to be present on the surface of the acceptor cells (18) were incapable of effecting proteolytic activation. The data are thus consistent with the interpretation that intercellular transfer first requires that the vSAg be proteolytically processed. Furin-dependent processing of vSAg7 in CHO cells has been observed to occur at or near two consensus furin recognition sites (residues 68 to 71 and 168 to 171). Proteolytic processing
The former site was unnecessary, because a mutant vSAg7 that lacked a furin recognition motif at positions 68 to 71 underwent intracellular transfer efficiently. These data, along with the apparent requirement for furin-dependent processing, suggest that the active vSAg is a carboxyl-terminal proteolytic fragment. This interpretation suggests that all of the sites necessary for interaction of the vSAg with both the class II protein and the T-cell receptor are encoded on a carboxy-terminal vSAg proteolytic fragment (residues 171 to 321). It is possible that the vSAg amino-terminal proteolytic processing product may serve to facilitate intracellular transport of the active vSAg or to perform yet uncharacterized roles in viral pathogenesis. Thus, the vSAgs, although produced as integral membrane proteins, may function in their active form in a manner equivalent to the bacterial superantigens, which are produced as small soluble proteins that freely associate with APCs in vivo.

vSAg7 activity could be transferred to class II-positive acceptor cells using cell-free supernatant from vSAg7 donor cells, indicating that cell-to-cell contact was not required for transfer. Although T-cell stimulation was relatively inefficient, these experiments used vSAg obtained after only 2 h of culture and are therefore not directly comparable to those performed under conditions of continuous culture. These findings are consistent, however, with the previous study that demonstrated inefficient vSAg transfer when donor and acceptor cells were separated by a cell-impermeable membrane (4). The inefficiency may be due to the apparent instability of the vSAgs (9). This instability may act to limit the transfer of the vSAg during viral infection to only closely associated APCs.

It is possible that vSAg intercellular transfer is an important facet of MMTV infection. Retroviruses typically require cycling cells for productive infection (15, 26), and so the MMTV may facilitate transfer of the vSAg to noninfected resting B cells. In this model, uninfected B cells that presented transferred vSAgs, upon activation by T cells, would become targets for infection. However, in one study, activation of resting B cells in vivo with lipopolysaccharide did not enhance infection by MMTVs (5), suggesting that resting, not activated, B cells were targets of viral infection. Alternatively, vSAg intercellular transfer may contribute to pathogenesis during viral infection of class II-negative cells such as T lymphocytes (20).
ACKNOWLEDGMENTS

We thank Donal Murphy and William Lee for critical reviews of the manuscript and for the B10 congenic mice, and we thank the Wadsworth Center Immunology Core Facility and the Computational Molecular Biology and Statistics Core Facility. This work was supported by Public Health Service grant CA69710-02.

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