Activation of NK cells by HIV-specific ADCC antibodies
Role for granulocytes in expressing HIV-1 peptide epitopes

Vijaya Madhavi,† Marjon Navis,† Amy W. Chung,† Gamze Isitman,† Leia H. Wren,† Robert De Rose,† Stephen J. Kent,* and Ivan Stratov†

1Department of Microbiology and Immunology; University of Melbourne; Melbourne, VIC Australia

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Introduction

Developing an HIV vaccine is a global priority. Several lines of evidence suggest antibodies that trigger NK cell mediated killing of virus-exposed cells termed antibody-dependent cellular cytotoxicity (ADCC) could contribute to the prevention or control of HIV infection. Several human cohort studies suggest ADCC antibody responses correlate with slower progression to HIV.1-4 Passive antibody transfer studies in macaques demonstrate a role for ADCC antibodies in controlling SHIV infection.5 Macaque-SIV vaccine studies have suggested a role for ADCC antibodies in protective immunity.6-8 The Thai RV144 human HIV vaccine efficacy trial, which induced high levels of HIV-specific ADCC antibodies, showed partial protection from infection that has been linked to non-neutralizing antibodies.9-11 There is considerable interest in understanding how HIV-specific ADCC could be utilized in an HIV vaccine strategy.9

Most commonly studied in vitro ADCC assays measure the ability of these antibodies to mediate killing of immortalized cell lines expressing HIV proteins.1,7,12 These assays have been important in defining the utility of ADCC antibodies. Our group has described a whole blood based ADCC assay that measures activation of NK cells (e.g., expression of IFNγ or the de-granulation marker CD107) in response to ADCC antibodies in HIV-infected blood and overlapping 15-mer HIV peptides.13,14 Serum transfer experiments showed the activity was mediated by IgG immunoglobulin within the HIV+ serum. Linear HIV ADCC epitopes could be mapped using individual peptides from within the overlapping peptide pool. Using this assay we recently reported the emergence of viral escape variants following ADCC selection pressure15 and that ADCC responses to particular epitopes are associated with slow HIV progression.16 Furthermore, other groups have also reported HIV-specific NK cell activation in reaction to HIV-peptide stimulation.17,18

The mechanism of activation of NK cells by exogenous HIV peptide ADCC epitopes is investigated in this manuscript. In order for ADCC activity to occur, three key components are generally required, namely: (1) target cells that express the HIV antigen, (2) antibodies that bind the viral antigen and (3) effector cells expressing Fcγ receptors, such as NK cells, which bind the Ag-Ab complex. Activated NK cells will secrete a number of
CD4 T cell line with HIV Envelope protein and showing that CD4 cells are target for ADCC related killing. We compared HIV Envelope gp140 protein pulsed CD4 T cells in the RFADCC with Envelope peptide stimulated whole blood in the NK activation ADCC assay. A comparable number of individuals responding to the protein also responded against the peptide antigen.13 Furthermore, comparison of Envelope gp140 protein and Envelope peptides in our NK activation ADCC assay indicated similar percentages of CD107 and IFNγ expression.13 Within the whole blood assay we have described, we envisage that one or more primary blood cells express the peptide epitopes and may serve as a target for NK cell-mediated ADCC in the presence of HIV+ plasma. This study sought to further understand the mechanisms behind HIV-1 peptide specific NK cell activation within the whole blood ADCC assay. We hypothesized that we could identify this population using fluorescently labeled peptide epitopes and this population would specifically undergo apoptosis and reduction in cell numbers in association with NK cell-mediated ADCC in the presence of HIV+ plasma.

Results

ADCC peptide epitope binding is not HLA restricted. We have previously demonstrated that HIV-1 peptide antigens within whole blood samples can induce CD56+ NK cells to liberate cytokines via an ADCC mechanism.14 We have also observed CD107 de-granulation13 however we have not confirmed whether cytotoxicity of target cells is induced. Our previous work has shown that peptide-antibody complexes added to whole blood do not activate substantial numbers of NK cells.13 We hypothesized that a cell population within whole blood express these HIV-1 peptides to ADCC antibodies in HIV+ plasma, which in turn bind and activate NK cells to express IFNγ (Fig. 1A), resulting in killing of the peptide-expressing cell. We have previously shown that cytokine producing cells were not CD3+ T cells, that optimal responses were generated with a peptide length of 14 amino acids or longer and that responses were contingent on the presence of HIV positive IgG and not contingent on the presence of autologous cells.14 Although it seems unlikely that HLA molecules would present peptides for ADCC recognition, we have not previously excluded this possibility.
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termed Vpu19; sequence EMGHHAPWDVDDL14). This Vpu subtype B linear ADCC epitope in the HIV-1 Vpu protein ADCC responses were HLA-restricted. We utilized an HIV-1 and 9-control subjects (C1 and C9 are shown in Fig. 1B CD56+ NK-cells from HIV+ subject cells (S1; top right plot) lymphocytes (expression after a 5-h incubation by gating on CD3-CD56- negative control subjects (C1-C9). We evaluated NK cell IFNγ expression after a 5-h incubation by gating on CD3-CD56+ lymphocytes (Fig. 1B). The Vpu19 peptide elicited activation of CD56+ NK-cells from HIV+ subject cells (S1; top right plot) and 9-control subjects (C1 and C9 are shown in Fig. 1B, lower plots) in the presence of HIV+ plasma. HLA-A, B, C, DRB1, DQA1, DQB1 typing of S1 and C1-C9 showed that although some alleles were shared by several donors (e.g., A*02:01 by 4 donors), there were no common class I or class II alleles identified among all 9 donors despite the activation of NK cells by ADCC antibodies in all 9 donors (Table 1). These results indicate that presentation of peptides to ADCC antibodies is not dependent on HLA restriction.

Peptides bind to granulocytes inducing killing during the ADCC assay. Identifying cells within whole blood that bind ADCC peptide epitopes and potentially act as peptide-expressing cells was accomplished by using flow cytometry and tracking Vpu19 peptide conjugated with FITC fluorochrome (Vpu-FITC; GL Biochem, Shanghai, China), SIV Gag peptide KC10 (sequence KKFGAEVVPC) was conjugated to AlexaFluor 488 fluorochrome (GL Biochem) and used as a control. The fluorescently labeled Vpu peptide (Vpu19-FITC) was still able to elicit antibody-mediated NK cell-IFNγ expression (Fig. 2A, upper plots). Vpu-FITC bound almost exclusively to cells with high forward and side scatter properties suggesting they were granulocytes (Fig. 2A lower plots). The peptide-expressing cells with high scatter properties were confirmed to be granulocytes by surface marker staining with the granulocyte marker CD66c (98% of cells). The peptide expressing cells were largely negative for CD3, CD8, CD14, CD20 and CD56 expression (< 6% of these subsets) in the assay.

We hypothesized that in the presence of HIV+ plasma, cells bound with Vpu ADCC peptide would be killed over time compared with cells incubated with a control peptide. Similarly, a major outcome of NK cell-mediated ADCC activation is the release of granzymes and perforin from the effector cells that lyse antigen expressing target cells, leading to a chain of intracellular events culminating in apoptosis of the target cell. Targets of cell lysis within the whole blood ADCC assay should be identifiable by the presence of apoptosis markers such as Annexin V expression. Membrane phosphatidylserine is translocated from the inner to the outer leaflet of the plasma membrane early during apoptosis.20 Annexin V binds with high affinity to surface phosphatidylserine and is a useful flow cytometry reagent to identify cells undergoing early stages of apoptosis.21

After a 5-h incubation in the presence of HIV+ plasma, CD66c+ cells bound with the Vpu-FITC peptide decreased by 10% (Fig. 2B, lower right plot) while CD66c+ cells did not decrease over time when incubated with a control peptide (Fig. 2B, upper right). Reduction in CD66c+ granulocytes over time (Fig. 2C, upper right bar graph) was associated with increasing expression of the apoptosis marker Annexin V (Fig. 2C, lower right bar graph). As a comparison, there was no reduction in CD56 cells numbers over time (Fig. 2C, left upper bar graph) or Annexin V expression by CD3+ cells (Fig. 2C, lower left bar graph). Further, Annexin V expression by CD2+ cells did not increase during the incubation time course while Annexin V expression by CD20+ and CD14+ cells did increase by approximately 0.5% and 1% of total number, respectively, during the incubation but not relative to control samples, suggesting some cell apoptosis during the incubation unrelated to the ADCC peptide. These results suggest that cells expressing the ADCC peptide epitope were targets for NK-mediated killing in the presence of ADCC Abs from HIV+ plasma during the assay. Replicate experiments using 5 different healthy donor NK cells were performed and we found a consistent and significant loss of peptide-expressing granulocyte numbers over time when incubated in the presence of HIV+ plasma that elicits peptide-specific ADCC (Fig. 2D).

It is important to note that granulocyte loss does not occur in the absence of other leukocytes. Granulocytes numbers decrease when the ADCC assay is performed in whole blood containing NK cells (Fig. 2E, second bar) but not in the presence of Percoll purified granulocytes alone (Fig. 2E, fourth

Table 1. NK cell activation ADCC response to HIV is not HLA-restricted

<table>
<thead>
<tr>
<th>NK cell donor</th>
<th>HLA A</th>
<th>HLA B</th>
<th>HLA C</th>
<th>HLA DRB1</th>
<th>HLA DQA1</th>
<th>HLA DQB1</th>
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<td>27:05;27:05</td>
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<td>1,3</td>
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<td>03:04;15:04</td>
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<td>04:03;07:02</td>
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<td>18:01;38:01</td>
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*% NK cells expressing IFNγ.

Granulocytes express ADCC epitopes and are targets for killing. (A) Fluorescently labeled Vpu peptide bind to granulocytes. The ADCC epitope Vpu peptide 19 (sequence EMGHAPWDVDDL) was conjugated with FITC fluorochrome. The fluorescent Vpu peptide induced robust ADCC activity in the presence of HIV+ plasma (top right plot). Based on FSC and SSC criteria, cells expressing fluorescent peptide localized primarily to granulocytes (bottom right plot). (B) Loss of CD66c+ granulocytes (bottom right plot) occurs in the presence of HIV+ plasma during the 5 h incubation following stimulation by Vpu peptide compared with a control peptide (sequence KKFGAVVPC) that has no known ADCC inducing ability (top right plot). (C) Granulocytes are the major cells that undergo apoptosis in the NK cell activation ADCC assay in the presence of HIV+ plasma. Decrease in CD66c+ granulocytes (top right graph) but not CD56+ NK cells (top left graph) occurred over time. Annexin V staining indicated that CD66c+ granulocytes were undergoing apoptosis (bottom right graph) compared with CD3+ T cell populations over time. Cell loss and Annexin expression in the presence of Vpu peptide (black bars) was compared with a control peptide (white bars). (D) Across 5 donors, granulocyte numbers reproducibly fell in the presence of HIV+ plasma following Vpu peptide stimulation (circles) compared with control peptide stimulation (triangles; mean and SD shown). (E) Granulocyte loss does not occur in the absence of other leukocytes. Granulocytes numbers decrease in the presence of HIV+ plasma when the ADCC assay is performed in whole blood containing NK cells (2nd bar) but not in the presence of purified granulocytes alone (4th bar). Loss of granulocytes in the presence of Vpu peptide (black bars) was compared with incubation in the presence of a control peptide (white bars).

Loss of granulocytes in the presence of Vpu peptide and HIV+ plasma was compared with incubation in the presence of a control peptide, suggesting the presence of both the ADCC peptide antigen and the peptide-expressing cell (i.e., granulocytes in this case) is necessary for ADCC to occur in the presence of HIV+ plasma.
Higher ADCC activity is observed in the presence of granulocytes. The uptake of peptides by granulocytes, the expression of the apoptosis marker Annexin V by granulocytes and the loss of granulocytes suggested that granulocytes were important in mediating peptide-specific ADCC responses. To further assess the importance of granulocytes within the peptide-specific ADCC assay, we studied the effects of granulocyte-depletion on the ADCC response. We evaluated ADCC-mediated NK cell activation in the presence of either whole blood (WB), PBMC alone (isolated over a Ficoll gradient; P-G) or PBMC preparations replenished with autologous granulocytes isolated over a Percoll gradient (P+G). We attained over 95% depletion of granulocytes, using the Ficoll gradient (Fig. 3A, middle plot), and found that the ADCC-mediated NK cell activation was significantly reduced in the absence of granulocytes (Fig. 3B, middle plot) compared with WB (Fig. 3B, left plot). The ADCC-mediated NK cell activation was significantly restored when autologous granulocytes were added back into the incubation (P+G, Fig. 3B right plot). This experiment was repeated with 5 different HIV-negative donors, and a consistent and significant restoration of ADCC activity was detected when granulocytes were replenished in the cultures (Fig. 3C). The individual donors results are graphed in Figure 3D.

HIV-1 gp140 Envelope protein binds to CD4 T cells which are targeted by ADCC. Although our results confirm that cells that express ADCC peptide antigens are targets for killing and this is associated with expression of IFNγ from CD56+ NK cells, the role of granulocytes in expressing ADCC epitopes was unexpected. Given that HIV principally infects via binding to the CD4 receptor, we therefore investigated whether similar results could be obtained using whole gp140 envelope protein. Akin to our approach with FITC-labeled peptides, we incubated whole blood from an HIV positive individual with biotinylated-gp140 and then tracked gp140 binding and cell culture kinetics using PE-conjugated streptavidin.

We first confirmed that biotinylated-gp140 effectively induced ADCC-mediated NK cell IFNγ expression in the presence of HIV+ plasma, similar to our previous work with unlabeled gp140 (Fig. 4A). Cells from an HIV negative subject were incubated with gp140 protein in the presence of HIV negative plasma or plasma from a person with ADCC responses to gp140. We found that the number of lymphocytes

**Figure 3.** Higher ADCC Activity is observed in the presence of granulocytes. Autologous granulocytes were added back to PBMC and assessed for their ability to restore ADCC activity. Whole blood (WB), PBMCs (isolated by ficoll gradient, P-G) and PBMCs with autologous granulocytes (isolated by percoll gradient) added back (P+G) were analyzed by flow cytometry. (A) Granulocyte population for the 3 assay conditions circled. (B) Depletion of granulocytes markedly reduces IFNγ expression by CD56 NK cells in the presence of HIV+ plasma after stimulation with Vpu peptide (top middle plot) compared with whole blood (top left plot); this response is significantly restored by adding back granulocytes (top right plot); a control peptide showed no responses (bottom plots). (C) Across 6 donors, there is a significant decrease in IFNγ expression by NK-cells in the presence of HIV+ plasma during the ADCC assay when granulocytes were depleted (p = 0.005); this response was nearly completely restored when granulocytes were added back to the incubation (p = 0.002; Mann-Whitney) (D) Shows the same data from C normalized to NK cell activation in the whole blood culture.
co-expressing CD4 and gp140 decreased sequentially over an 8-h incubation in the presence of HIV,3 the ability to target virus-infected cells, no HLA restriction and the ability to be transferred between subjects.14 We have previously demonstrated that CD56+ NK cells liberate cytokines in the presence of HIV peptide antigens, mediated by IgG, indicating an ADCC mechanism.14 We have shown that such responses are generated not only against Envelope but also against internal HIV proteins (including Vpu, Tat and Pol23), providing a potential additional benefit to ADCC-based vaccine strategies. However, the mechanism by which internal HIV proteins elicit such antibody responses was unknown. Possibilities currently under exploration include (1) parts of the virus being presented on the surface of HIV-infected cells (akin to HLA presentation of T cell epitopes) or (2) viral debris from lysed cells presented by antigen presenting cells.

Our results show that HIV-specific ADCC responses are not based on the HLA phenotype. NK cells from all 9 HIV negative donors expressed IFNγ when incubated in the presence of HIV+ plasma. It is of interest to note that the responses were not of a uniform magnitude, suggesting there may be other phenotypic features within individuals that influence these responses such as KIR phenotype25 or FcR polymorphism.24

We previously showed that HIV-antibody specific IFNγ expression by NK cells correlates with NK cell degranulation as assessed by the marker CD107a and granzyme B loss,13 however we have not previously identified the cell targets being killed by ADCC in this whole blood assay. We now show that cells bound with HIV ADCC peptide antigens expressed Annexin V and decrease in number during the assay incubation period indicating that actual killing of peptide-expressing cells is occurring. Further, we found that HIV ADCC peptides associate with CD66c+ granulocytes and that when granulocytes are depleted from the culture there is reduction antibody-mediated NK-cell IFNγ expression. This NK cell effector function is then significantly restored when granulocytes are added back to the assay culture. It is unclear as to how granulocytes are participating in the assay. HLA presentation by granulocytes leading to activation of immune responses has been described.25 However, our data indicate that HLA restriction is not important for ADCC-mediated NK cell activity; a fact confirmed by the observation that not all HLA bearing cells take up the fluorochrome-labeled antigen. Preliminary in vitro studies with lactacystin (proteasome inhibitor) and cytochalasin D (phagocytosis inhibitor) suggest that these inhibitors do not block ADCC peptide presentation. Possible mechanisms for ADCC peptide presentation

Discussion

Since the partial success of the RV144 vaccine trial, the role of non-neutralizing antibodies (such as antibodies that mediate ADCC) in protection from HIV has gained prominence.9 ADCC antibodies have numerous potential advantages: presence at mucosal entry sites of HIV,3 the ability to target virus-infected cells, no HLA restriction and the ability to be transferred between subjects.14 We have previously demonstrated that CD56+ NK cells liberate cytokines in the presence of HIV peptide antigens, mediated by IgG, indicating an ADCC mechanism.14 We have shown that such responses are generated not only against Envelope but also against internal HIV proteins (including Vpu, Tat and Pol23), providing a potential additional benefit to ADCC-based vaccine strategies. However, the mechanism by which internal HIV proteins elicit such antibody responses was unknown. Possibilities currently under exploration include (1) parts of the virus being presented on the surface of HIV-infected cells (akin to HLA presentation of T cell epitopes) or (2) viral debris from lysed cells presented by antigen presenting cells.

Figure 4. Cells co-expressing CD4 and HIV-1 gp140 Envelope protein reduce over time during the NK cell activation ADCC assay. (A) HIV-1 gp140 envelope protein antigen stimulation of the whole blood samples in the NK cell activation ADCC assay in the presence of HIV+ plasma induces NK cell IFNγ expression. (B) Cells co-expressing CD4 and gp140 reduced in number over time (% baseline) in the presence of HIV positive serum (right graph) but not in the presence of HIV negative serum (left graph). (C) The reduction in the number of cells co-expressing CD4 and gp140 was associated with an increase in Annexin V expression over time in the same population when incubated in the presence of HIV+ plasma (white bars) compared with HIV- plasma (black bars).
by granulocytes include FcR mediated uptake or other protein binding cell surface molecules such as integrins, a function which has mainly been studied in the context of autoimmunity. Other possibilities for molecules presenting short peptides by granulocytes to ADCC antibodies include selectins and scavenger receptors. The CD66 molecule itself could present peptides although preliminary competition assays could not confirm it (data not shown). Thus, although granulocytes permit us to define linear peptide ADCC epitopes in vitro, their role in vivo in HIV infection is not known. The large population of granulocytes lost in this 5 h ADCC assay suggests neutrophils are predominantly targeted since eosinophils and basophils make up a small proportion of granulocytes. However, further investigation of the specific types of granulocytes (neutrophils, eosinophils or basophils) playing the major role in ADCC would be interesting.

Granulocytes are not targets of HIV infection and not typically viewed as antigen-presenting cells. On the other hand, granulocytes are frequently depleted and have high levels of apoptosis during HIV and SIV infection. An initial role for neutrophils in HIV infection was postulated in the context of increased susceptibility of HIV-infected individuals to bacterial and fungal infections in neutropenic subjects. More recent results suggest additional roles for neutrophils given that neutrophils express several anti-viral factors, such as α-defensins and lactoferrin, which have anti-HIV-1 activity, possibly suggesting a protective role for neutrophils in HIV-1. Studies in mice have shown that neutrophils efficiently transport antigen to draining lymph nodes. Further, neutrophils have been shown to bind bothCCR5 and CXCR4 strains of HIV-1 and efficiently transfer virus to lymphocytes. This would assist enhanced presentation of viral antigens to initiate or expand humoral and cellular immune responses.

Can the association of ADCC peptide epitopes with blood granulocytes be harnessed to improve HIV vaccines? We speculate that it may be possible to engineer more efficient expression of ADCC epitopes (for example, by targeting granulocytes) to stimulate high levels of immunity prior to infection and improve the immunogenicity of ADCC-inducing HIV vaccines. Recently, Duval et al. showed that a bispecific antibody incorporating a broadly reactive anti-gp41 antibody, F240, and an anti-IgA receptor (CD89) antibody is effective at directing neutrophils to destroy HIV. Macaques are more resistant to SHIV infections in neutropenic subjects. More recent results suggest susceptibility of HIV-infected individuals to bacterial and fungal infections in neutropenic subjects. Also, gp140-biotin (kindly supplied by Dr. R. Center) was used in HIV-1 gp140 envelope protein assays; fluorochrome identification was achieved using PE-conjugated streptavidin (BD Biosciences Cat# 550401).

Materials and Methods

HIV-infected subjects. HIV-infected adults with previously described HIV-specific ADCC responses were recruited to donate blood samples. Subjects provided informed consent and the research were conducted under the auspices of the Alfred Human Health Research and Ethics Committee. HIV infected subjects provided sodium heparin anti-coagulated blood for fresh whole blood ADCC assays and blood plasma for the ADCC assays using HIV negative donor cells.

The NK cell activation ADCC assay. The NK cell activation ADCC assay was used as previously described. In brief, for fresh whole blood assays, 200 μl of HIV+ whole blood (or 100 μl HIV negative donor whole blood and 200 μl HIV+ plasma) was incubated with HIV-1 Vpu19 peptide. The incubation to detect intracellular NK activation occurred in the presence of Brefeldin A (final conc. Ten μg/ml, Sigma); time course experiments to detect killing of peptide labeled cells proceeded without addition of Brefeldin A. At the end of the incubation cells were fixed and permeabilized with 1% formaldehyde and Proteinchex (BD) for 30 min at 4°C. At the end of the incubation CD56+CD3- or CD2+CD3- NK lymphocytes were studied for the expression of intracellular IFNγ. Fluorescent antibodies utilized in the NK cell activation ADCC assays were CD3 (BD Biosciences, San Jose, CA, catalog# 347344 PerCP and #641397, APC-H7), CD2 (BD #556611, FITC), CD56 (BD #555156 PE and #335791, PE-Cy7), CD66c (BD #551478, PE), IFNγ (BD #557795 Alexa700 and #554702, APC), CD8 (BD # 2217, PE), CD14 (BD # 557153, FITC), CD20 (BD # 335793, PE-Cy7) and Annexin V (BD #550474 APC).

Antigen conjugated with fluorochrome binding in NK cell activation ADCC assay. The fluorescent-labeled peptides, Vpu-FITC and control peptide were used as peptide antigens within the NK cell activation ADCC assay. Also, gp140-biotin (kindly supplied by Dr. R. Center) was used in HIV-1 gp140 envelope protein assays; fluorochrome identification was achieved using PE-conjugated streptavidin (BD Biosciences Cat# 550401).

Isolation of leukocytes. PBMC were separated using Ficoll density gradient centrifugation as per manufacturers’ protocol. The granulocytes were isolated from peripheral blood by density gradient centrifugation (MACS Miltenyi Biotec method) using Percoll gradient and the granulocyte purity was up to 96% as measured by CELL DYN Emerald analyzer. Briefly, the whole blood was carefully layered on top of the Percoll gradient and then centrifugation at 400xg for 30 min at 20°C in a swinging bucket rotor without brakes. Then granulocytes were carefully aspirated from the cell layer directly above the RBC and used in the assay to add back autologous granulocytes to Ficoll-separated PBMC.

HLA typing by Australian Red Cross Blood Service. HLA class I and II typing was performed on PBMC by the Australian Red Cross Blood Service using the Luminox® SSO (lab type) technique, as per manufacturer’s instructions (see www.onelambda.com).

Conclusion

In summary, we have shown that within the NK cell activation ADCC assay, cells expressing certain HIV antigens are targeted.
for killing by NK cells via ADCC and this mechanism is not dependent on HLA typing. These results provide a rapid and simple way of identifying ADCC epitopes in vitro. Exactly how the interactions are mediated and whether this knowledge can be utilized to improve antigen delivery in vivo and assist in vaccine design requires further investigation.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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