Natural Killer Cell Cytotoxicity Is Suppressed by Exposure to the Human NKG2D Ligand MICA*008 That Is Shed by Tumor Cells in Exosomes

Omodele Ashiru1, Philippe Boutet1, Lola Fernández-Messina1, Sonia Agüera-González1, Jeremy N. Skepper2, Mar Valés-Gómez1, and Hugh T. Reyburn1

Abstract

The MHC class I–related chain (MIC) A and MICB ligands for the activating receptor NKG2D can be shed from tumor cells, and the presence of these soluble molecules in sera is related with compromised immune response and progression of disease. Recently, thiol disulphide isomerases and members of the ADAM (a disintegrin and metalloproteinase) gene family were identified as key enzymes in mediating MICA/B shedding from cells. Here, we report shedding of the most frequently expressed MICA allele in human populations (MICA*008) into exosomes, small membrane vesicles that are secreted upon fusion with the plasma membrane. Although similar to other MICA/B molecules in the extracellular domain, the predicted transmembrane and cytoplasmic domains of MICA*008 are quite different, and this difference seemed to be critical for the mode of release from tumor cells. Treatment of natural killer (NK) cells with exosomes containing MICA*008 molecules not only triggered downregulation of NKG2D from the cell surface but also provoked a marked reduction in NK cytotoxicity that is independent of NKG2D ligand expression by the target cell. Our findings reveal a mechanism of NK suppression in cancer that may facilitate immune escape and progression. Cancer Res; 70(2); 481–9. ©2010 AACR.

Introduction

NKG2D is a C-type lectin-like activating receptor expressed on all natural killer (NK) cells, as well as TCRγδ+ and CD8+ TCRαβ+ T cells (1), where it can act to costimulate the activation of naive T cells (2) and even trigger cytotoxicity in the absence of T-cell receptor (TCR) ligation (3). In humans, NKG2D binds to MHC class I–related chain (MIC) A, MICB, and UL16-binding proteins whose expression is restricted or absent on normal tissues but is induced in situations of stress and disease (4). The molecular basis of this altered expression is not well understood, although it has been described that a variety of stresses can upregulate NKG2D ligand (NKG2D-L) expression (5–7). The expression of NKG2D-L has been described in multiple types of tumors (4, 8), and evidence is accumulating that strongly suggests that activation of the immune system, mediated by engagement of NKG2D with its ligands, plays an important role in immunosurveillance of cancer (9, 10). This strong selection pressure seems to have led tumor cells to evolve mechanisms to minimize or avoid the response mediated by NKG2D by shedding NKG2D-L from the cell surface. The shedding of soluble MICA by human tumors not only hinders recognition of the MICA-expressing tumor cells but also leads to downregulation of NKG2D expression on circulating CD8 T cells, NK cells, and γδ T cells, and so the antitumor immune response is impaired (11). Indeed, high levels of soluble MIC molecules in sera correlate strongly with poor clinical outcome in patients suffering from various types of cancer, including colon (12) and prostate cancers (13). The mechanisms of shedding of MICA/MICB molecules have been studied intensively. Proteolytic cleavage of MICA has been shown to depend on the thiol isomerase Erp5 (14) and metalloproteinas (15), specifically the ADAM (a disintegrin and metalloproteinase) family members ADAM10 and ADAM17 [also known as tumor necrosis factor–α–converting enzyme (TACE); refs. 16, 17]. However, no reduction in soluble MICA shedding was observed from HeLa and A375 cells after treatment with metalloproteinase inhibitors (14). One explanation for this discrepancy could be that the extensive polymorphism of MIC molecules affects the shedding process. Here, it is interesting to note that the allele of MICA expressed by HeLa cells is MICA*008, which is by far the most frequently expressed allele of MICA, with an allele frequency...
of 21% to 47% in diverse populations (18–22). Strikingly, although MICA*008 is very similar to other MIC molecules in the extracellular domain, the predicted transmembrane and cytoplasmic domains of this allele are very different. This sequence difference is accompanied by functional differences. Specifically, although surface expression of many MICA molecules is blocked in cells infected by human cytomegalovirus (HCMV) or Kaposi sarcoma–associated herpesvirus (KSHV), MICA*008 seems to be resistant to downregulation by these viruses (23, 24).

The transmembrane and cytoplasmic regions of proteins usually play an important role in their trafficking within the cell. Thus, the different MICA alleles might follow different pathways that could affect their mechanism of shedding. We have recently described that some alleles of MICB can be endocytosed and are found intracellularly in the trans-Golgi network (TGN) and the endosomal system (25). Late endosomes derive from early endosomes through a maturation process that involves gradual change in content as well as the incorporation of material from the plasma membrane and TGN-derived transport vesicles. During this process, late endosomes can accumulate up to hundreds of internal vesicles, which, after fusion of the organelle with the plasma membrane, can be released to the surrounding medium as exosomes (see ref. 26 for review). Here, we describe that full-length, and therefore membrane-resident, MICA*008 proteins can be released from cells in exosomes. In contrast, the MICA*019 allele, which is almost identical to MICA*008 in the extracellular domains but has transmembrane and cytoplasmic tail domains typical of most MICA molecules, was only found in cell culture supernatant as a soluble truncated molecule. Despite this difference in mechanism of shedding, both soluble MICA*019 and exosomal MICA*008 molecules are able to trigger downregulation of NKG2D from the NK cell surface. In addition, exposure of NK cells to MICA*008-containing exosomes triggers a marked loss of cytotoxic function. Thus, exosome release can be an alternative pathway to metalloproteinase-mediated shedding of MICA/MICB molecules for immune evasion.

Materials and Methods

Cells and antibodies. The cell lines HeLa (MICA*008/008; ref. 18), HepG2 (MICA*002/009; ref. 24), and MelJuSo were cultured in DMEM supplemented with 10% FCS, l-glutamine, and sodium pyruvate. HCT116 cells (MICA*001/009; ref. 27) were cultured in RPMI 1640 with the same supplements. Chinese hamster ovary (CHO) cells were cultured in Ham’s F12 medium supplemented with 10% FCS, l-glutamine, and pyruvate.

The isolation and culture of human primary polyclonal NK cells, as well as culture of the cell lines 721.221, RPMI-8866, and Daudi, were as previously described (28). Influenza-specific T-cell lines were derived from peripheral blood lymphocytes of an HLA-A2-positive donor by growth on irradiated autologous B lymphoblasts pulsed with influenza A matrix peptide 58–66 (29). Goat polyclonal antibodies and mouse monoclonal antibody (mAb) specific for MICA/MICB were purchased from R&D Systems, mAb anti-NKG2D was from Santa Cruz Biotechnology, mAb MEM-259 (specific for CD63) was from Abcam, and rabbit anti-caveolin antibody was from Transduction Laboratories.

Molecular biology, plasmids, and transfection. The MICA*019 expression construct has been described previously (30). The MICA*008 expression plasmid was constructed by subcloning cDNA of full-length MICA*008, a gift from Dr. Paul Lehner (University of Cambridge, Cambridge, United Kingdom), into pcDNA3. The 019EC/008TMT chimera was constructed by excising the extracellular domain of the 008 allele by digestion with HindIII and DraIII and introducing the extracellular domain of 019 (also as a HindIII-DraIII fragment). Each MICA expression plasmid was mixed (9:1 ratio) with a vector conferring resistance to puromycin (31), and then CHO cells were transfected with this mixture using Lipofectamine 2000. Stable transfectants were generated by culture of transfected CHO cells in selective medium (8 μg/ml puromycin; Calbiochem) and, where necessary, by cell sorting.

Exosome isolation and characterization. Exosomes were isolated by differential centrifugation as described previously (32). Briefly, adherent monolayers were washed extensively with PBS and recultured in fresh medium (no serum) for 24 h. Cell culture media were centrifuged twice for 10 min at 300 × g to remove cell debris and then centrifuged for 30 min at 10,000 × g and for 2 h at 100,000 × g sequentially. Soluble proteins were recovered from the 100,000 × g supernatant by trichloroacetic acid (TCA) precipitation. The pellets were solubilized in reducing SDS sample buffer and analyzed by Western blot. In some experiments, further purification of exosomes by flotation on a sucrose gradient was performed as described previously (33). Briefly, exosome samples (500 μL) were mixed with 2.5 volumes of 85% (w/v) sucrose in 20 mmol/L Tris-HCl (pH 7.5) containing 150 mmol/L NaCl and 5 mmol/L EDTA (TNE) and placed in centrifuge tubes. The mixtures were layered successively with 4 mL of 60% (w/v), 3 mL of 30% (w/v), and 1 mL of 5% (w/v) sucrose in TNE and centrifuged at 200,000 × g for 18 h at 4°C. Fractions (1 mL) were collected from the top to the bottom of the tube. Aliquots of these fractions were analyzed by Western blot.

Detergent-resistant membrane fractionation. Detergent-resistant membrane (DRM) and detergent-soluble membrane fractions were prepared as previously described (16). Western blot was performed using antibodies specific for MICA/MICB and caveolin. Quantitative analysis of the Western blot data was done using ImageJ software.3

Western blot. Cell lysates were prepared by incubation in TNE buffer containing 1% NP40 and protease inhibitors. After centrifugation to remove nuclei and cell debris, samples were run on 10% or 12% SDS-PAGE gels (for experiments visualizing CD63, the samples were run under nonreducing conditions) and transferred to Immobilon-P (Millipore). The membrane was blocked using 5% nonfat dry milk in PBS–0.1% Tween 20, and then specific antigens were

detected by incubating the membrane with the indicated first antibody followed by horseradish peroxidase–conjugated secondary antibodies. Proteins were visualized using the enhanced chemiluminescence system (Amershams Pharmacia). In some experiments, samples were treated with peptide N-glycosidase F (PNGase F; New England Biolabs) according to the manufacturer’s instructions.

Electron microscopy. Electron microscope examination of exosomes was carried out by floating a carbon-coated 400-mesh Formvar EM grid on top of one drop of freshly prepared exosomes (60 μg/mL in PBS) for ~1 min. The grid was then briefly washed with deionized water and floated on a drop of 2% uranyl acetate. Samples were examined using a Philips CM100 operating at 60 or 80 kV.

NKG2D downregulation. Interleukin-2 (IL-2)–activated human NK cells (1 × 10⁶), 3 d after stimulation with IL-2 (50 units/mL; R&D Systems), were incubated in 96-well flat-bottomed plates for 24 h with supernatants of untransfected CHO cells or CHO cells transfected with either MICA*019 or MICA*008. When using exosome fractions, 40 to 100 ng of total protein were added to the NK cells. NKG2D surface expression was monitored by staining with mAb specific for NKG2D (clone 1D11, 1 μg/10⁵ cells) and flow cytometry using a FACS® cytometer running CellQuest® software (BD Biosciences).

Cytotoxicity assay. Cytotoxicity assays were carried out using a one-step fluorimetric assay based on the use of AlamarBlue (Invitrogen; ref. 34). Effector cells alone, target cells alone, and mixes of effectors and target cells at the indicated E:T ratios were incubated with AlamarBlue in 96-well flat-bottomed plates at 37°C in a humidified 5% CO₂ incubator overnight. Following incubation, the fluorescence of the AlamarBlue was read on a Synergy HT plate reader (BioTek) with excitation at 530 nm and emission at 590 nm at 37°C. The percentage specific lysis was calculated using the following formula:

% Specific lysis = 100 × \[
\frac{\text{(AF of targets alone)} - \text{(AF of mix)}}{\text{(AF of effectors alone)}}
\]

where AF = absolute fluorescence units.

Results

Biochemical analysis of MICA proteins shed from tumor cell lines. In previous experiments, biochemical analysis of MICA/MICB molecules shed from transfectants showed that the shed MICB was a truncated species that remained soluble after centrifugation at 100,000 × g (16). We extended this analysis to study the MICA/MICB molecules shed from a panel of tumor cell lines that express endogenous MICA and MICB molecules (Supplementary Fig. S1). It is not possible to distinguish between MICA and MICB in Western blot experiments due to their high homology, but we could see that soluble MICA/MICB molecules shed from the colon carcinoma cell line HCT116 had a molecular weight similar to the soluble MIC shed from transfectants (16, 17), corresponding to MIC protein cleaved in the stalk region that connects the α3 and transmembrane domains (Fig. 1A). In contrast, analysis of MICA/MICB molecules released from HeLa cells showed that these proteins were not soluble as they pelleted on centrifugation at 100,000 × g. Moreover, their size corresponded to the full-length MICA/MICB protein present in the HeLa cell lysate (Fig. 1A and B). HeLa cells express the MICA*008 allele (18) and very little MICB (Supplementary Fig. S1; ref. 25); moreover, MICA*008 is distinct from other alleles of MICA due to the insertion of a guanine at position 952 in the transmembrane region, which results in expression of a protein with a truncated transmembrane region and a very short cytoplasmic tail (35). For this reason, MICA*008 can be easily distinguished from other MICA and MICB molecules by their molecular weight. Analysis of other tumor cell lines showed that although some of the MICA/MICB molecules released by the cell lines HepG2 and MelJuSo were soluble and comparable in size with those shed by the HCT116 cell line, a proportion of the MICA/MICB molecules shed from the melanoma cell line MelJuSo were present in the 100,000 × g pellet with a molecular weight of 35 kDa after N-glycosanase digestion (Fig. 1C). Cloning...
and sequencing experiments confirmed that this cell line expressed MICA*008 (data not shown); thus, these results show that different MICA/MICB molecules can be released as biochemically different species even in the same cell line. In addition, the data show that MICA*008 is preferentially released as a full-length, membrane-anchored molecule. The simplest interpretation of these data is that MICA*008 was released in exosomes rather than by proteolysis.

**MICA*008 protein is released from HeLa cells in exosomes.** Exosomes are bioactive vesicles formed through the fusion of multivesicular bodies (MVB) with the plasma membrane, leading to the release of 30- to 100-nm lipid bilayer vesicles (36). Physically, exosomes are equivalent to cytoplasm enclosed in a lipid bilayer, with the external domains of transmembrane proteins exposed to the extracellular environment. Although exosome composition varies depending on the cell type of origin, one of the common characteristic features of these vesicles is the presence of tetraspanins such as CD63 (36, 37). The MICA*008-containing pellet obtained by centrifugation at 100,000 × g, but not the supernatant, was enriched for CD63 (Fig. 2A). Previous studies (36) have shown that exosomes display equilibrium buoyant densities between 1.10 and 1.19 g/mL on linear sucrose gradients. So, to confirm that tetraspan proteins and HeLa cell MICA*008 localize to membranes with the same characteristics, exosomes isolated from the culture medium by differential centrifugation were floated into a linear sucrose gradient. The distribution of MICA and CD63 in the gradient was analyzed by Western blotting and shown to overlap substantially (Fig. 2B), providing further evidence that MICA*008 was shed from HeLa cells in exosomes. This conclusion was strengthened by electron microscopy analysis, which showed that the MICA-containing 100,000 × g pellet obtained from HeLa culture supernatant was composed of small membrane vesicles with diameters of 50 to 100 nm (Fig. 2C), typical of exosomes.

**Exosome shedding of MICA*008 is an intrinsic property of this allele.** Analysis of MICA shedding in CHO cells stably transfected with either the allele MICA*019 or MICA*008 confirmed that whereas MICA*019 was found solely as a truncated molecule in the 100,000 × g supernatant, a considerable proportion of MICA*008 molecules were shed in exosomes (Fig. 3).

These data indicated that the incorporation of MICA*008 into exosomes was an intrinsic property of this allele and not of the cell type in which it is expressed. Nevertheless, some soluble MICA*008 was released from the CHO transfectants. Because soluble MICA*008 is not detected on preparations from HeLa cells, which express the protein endogenously, our hypothesis is that detection of the truncated protein is related to the very high level MICA*008 expression by the transfected cells (Supplementary Fig. S2A), which is much higher than the levels of expression reported for MICA expression on cell lines and in vivo (38).

**MICA*008 is found in membrane microdomains.** The best-characterized mechanism underlying recruitment of proteins to MVBs is monoubiquitination and interaction...
with the ESCRT machinery (39). However, this route cannot be relevant for incorporation of MICA*008 to exosomes because this protein does not have lysine residues in its cytoplasmic tail. Another route for cargo recruitment to exosomes depends on the intrinsic physical properties of the protein and preferential segregation into raft-like microdomains (40). Thus, we analyzed whether MICA*008 was present in DRM microdomains. In agreement with observations made with MICB (16), ~25% of MICA*019 molecules are found in low-density detergent-insoluble fractions, whereas >60% of this protein is present in the detergent-soluble fractions (Fig. 4A). In contrast, the majority of MICA*008 molecules are present in low-density detergent-resistant fractions, whereas only 20% are found at the bottom of the gradient (Fig. 4B). MICA*019 and MICA*008 vary by only one amino acid in the extracellular domain, in a region distant from the NKG2D binding site (41), but are very different in the transmembrane region and cytoplasmic tail (Supplementary Fig. S2B; ref. 35). This suggested that the transmembrane/cytoplasmic tail of MICA*008 controlled the localization of MICA*008 to DRMs. This hypothesis was confirmed by the preferential localization in DRMs of a chimeric protein comprising the extracellular domain of MICA*019 and the transmembrane and cytoplasmic tail of the MICA*008 molecule (Fig. 4C and D). Analysis of culture supernatants of CHO cell transfectants expressing the chimeric 019EC/008TMT molecule revealed that this molecule was also shed in exosomes (Supplementary Fig. S3).

Supernatants containing either MICA*019 or MICA*008 can downregulate surface expression of the NKG2D receptor. As well as promoting tumor evasion by reducing the cell surface expression of NKG2D-Ls on tumors, the release of soluble MICA has also been reported to trigger a systemic downregulation of the NKG2D receptor on NK cells and CD8 T cells (11–13). In this context, it was of interest to compare the effect of culture supernatants containing MICA*008 (released in exosomes) or MICA*019 (soluble) on cell surface expression of NKG2D on primary human NK cells. To avoid the confounding effects of factors such as exosomal transforming growth factor-β on lymphocyte proliferation and NKG2D expression (42–44), we chose to compare supernatants of MICA*019 or MICA*008 transfectants with...
supernatants of untransfected CHO cells. A representative example of such an experiment is shown in Fig. 5A. These data clearly indicate that both MICA*019- and MICA*008-containing supernatants can decrease cell surface NKG2D expression. Analysis of multiple experiments shows that these effects are statistically significant (Fig. 5B). Interestingly, although the supernatants from CHO cells expressing MICA*008 shed less MICA than MICA*019-expressing cells, the MICA*008 culture supernatant consistently triggered more downregulation of NKG2D (five of five experiments).

**Exosomal MICA*008 can downmodulate surface expression of NKG2D and compromise NK cell cytotoxicity.** Unlike HeLa cells, the culture supernatant of the CHO cells overexpressing MICA*008 contains some soluble MICA*008 protein. Thus, it was important to test whether purified exosomes containing MICA*008 molecules affected cell surface expression of NKG2D on NK cells. Figure 6A shows that exposure to exosomes purified from MICA*008 transfectants, but not from untransfected cells, triggers a marked reduction in cell surface NKG2D on NK cells. Importantly, whereas pretreatment of NK cells with exosomes prepared from untransfected CHO cells had essentially no effect on the ability of these NK cells to lyse CHO and MICA*019 transfectants, treatment with exosomes containing MICA*008 molecules was associated with a marked reduction in cytotoxicity against both of these target cells (Fig. 6B). These observations

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**Figure 5.** Both soluble and exosomal MICA downmodulate surface expression of NKG2D. Culture supernatants were collected from CHO cells transfected with MICA*019 or MICA*008. Supernatant from untransfected CHO cells was used as control. A, supernatant from CHO cells and MICA-transfected CHO cells was incubated with IL-2–activated human NK cells for 24 h. The amount of NKG2D on the surface of the NK cells was then analyzed by flow cytometry. The result shown is representative of five experiments. B, downmodulation of NKG2D is significant (Student's t test) after treatment with either MICA*019- or MICA*008-containing supernatants. Data are expressed as a percentage of the NKG2D expression observed on NK cells incubated in medium alone.

**Figure 6.** Exosomal MICA*008 can trigger NKG2D downmodulation and compromise NK cell cytotoxicity. Incubation of NK cells with purified exosomes isolated from CHO cells transfected with MICA*008, but not untransfected CHO cells, leads to a marked reduction in NKG2D cell surface expression (A; four experiments) and compromised NK cell–mediated lysis of parental and MICA-transfected CHO cells (B), as well as unrelated “third-party” cells such as 721.221 (C). Note that AlamarBlue measures cell metabolism; thus, negative values for percentage specific lysis indicate target cell proliferation. D, preincubation of CTL, specific for the influenza matrix peptide 58–66 bound to HLA-A2, with either control or MICA*008-containing exosomes, has no effect on specific cytotoxicity (two experiments).
suggested that treatment with MICA*008 exosomes might trigger a general deficit in NK cell cytotoxic function. To test this idea, the effect of exosome treatment on the ability of NK cells to lyse 721.221 cells was tested because recognition of this cell line by NK cells is mediated by multiple activating receptors, including 2B4 (45), NKP44, and NKP46 (46). Strikingly, pretreatment with MICA*008 exosomes markedly impaired NK cell-mediated lysis of 721.221 cells. In light of these observations, we investigated whether treatment with exosomes would also inhibit immune recognition by antigen-specific CTL expressing NKG2D. Although exposure to MICA*008-containing exosomes, but not CHO exosomes, provoked downregulation of CTL surface NKG2D (Supplementary Table S1), this pretreatment had no effect on the ability of these CTLs to recognize and specifically lyse peptide-pulsed target cells (Fig. 6D).

Discussion

MICA*008 is the most frequently expressed allele in nearly all populations studied (18–22). Although similar to other MICA molecules in the extracellular domain, MICA*008 is distinct from other alleles of MICA due to a single-nucleotide insertion in the transmembrane region, which gives rise to a truncated transmembrane region and cytoplasmic tail (35). Here, we show that the particular carboxy terminus of MICA*008 is associated with a different mechanism of release from the cell surface and an altered distribution of the protein to microdomains within the plasma membrane. In contrast to other MICA alleles that are shed as truncated soluble species after proteolysis by ADAM17/TACE (16, 17), MICA*008 accumulates in the supernatant as a full-length molecule present in the membrane of exosomes. This observation probably explains why metalloprotease inhibitors do not inhibit the release of MICA molecules from HeLa cells (14). It is interesting to note that MICA*008 could potentially be released as a soluble protein because CHO cell transfectants with high levels of expression of MICA*008 do release some soluble MIC. However, soluble MICA*008 was not observed in our experiments studying HeLa cells, suggesting that shedding of soluble MICA*008 is an artifact of the ~100-fold higher levels of expression of MICA*008 in the CHO transfectant compared with HeLa cells.

The trafficking of MICA*008 into the MVBs (which give rise to exosomes) seems to depend on the truncated carboxy terminus of this protein because this region confers preferential recruitment of MICA*008 into the particular lipid environment associated with the DRMs enriched in exosomes (40). Further studies are under way to characterize the particular physical properties of the MICA*008 protein that are responsible for this pattern of trafficking. However, it is interesting to speculate that this differential trafficking of MICA*008 is related to the resistance of MICA*008 to the HCMV and KSHV proteins that downregulate surface expression of many other MICA alleles (23, 24). These data also emphasize that sequence variation between different MICA and MICB alleles can be associated with marked differences in their function and confirm that trafficking of MICA/MICB molecules can have a profound effect on the mechanism of shedding of these molecules (16).

The release of MICA/MICB molecules from tumor cells is associated with a marked downregulation in cell surface expression of NKG2D on circulating NK and CD8+ T cells. The current data show that incubation with either soluble MICA*019 or exosomal MICA*008 molecules leads to downregulation of cell surface NKG2D expression, suggesting that both species are able to bind the receptor. However, although the culture supernatant of MICA*008 transfectants contains less MICA protein than that of cells transfected with MICA*019 (Fig. 3), incubation of NK cells with the MICA*008-containing supernatant triggers significantly more NKG2D downregulation than the MICA*019 culture supernatant (Fig. 5). This might be related to differences in how the two MICA species exist in the supernatant: soluble, probably monomeric MICA*019 versus multivalent MICA*008 molecules in exosome membranes. Importantly, purified exosomes containing MICA*008 protein triggered significant downregulation of NK cell surface NKG2D and compromised NK cell function. Strikingly, incubation with exosomes containing MICA*008 also impaired NK cytotoxicity known to be triggered by other activating receptors. The molecular basis of this general loss of NK function is unclear, but this phenomenon is reminiscent of previous descriptions of a global deficit in NK cell cytotoxic function after overnight incubation with NKG2D-L1-expressing cells (47). In contrast, exposure of CTL to MICA*008-containing exosomes, although triggering NKG2D downregulation (Supplementary Table S1), did not significantly affect the ability of the CTL to specifically lyse target cells. CTL recognition of the HLA-A2/matrix peptide 58–66 complex is mediated by an immunodominant high-affinity TCR (48). Thus, it remains possible that treatment with MICA*008-containing exosomes could affect other lower-affinity CTL, but it is interesting to note that these data are consistent with previous observations where constitutive Rae-1: transgene expression triggered local and systemic NKG2D downregulation with generalized defects in NK cell–mediated cytotoxicity but only mild CD8+ T-cell effects (49).

Elevated levels of soluble MICA have been detected in the sera of patients suffering from various types of cancer, including gastrointestinal malignancies, breast and lung tumors, melanoma, prostate cancer, pancreatic carcinomas, hepatocellular cancer, and leukemia (11, 50). Moreover, the levels of soluble MICA and MICB in patient sera can be used as diagnostic markers for cancer progression, where elevated levels of these proteins often correlate with a poor prognosis for the patient (50). This relationship is likely to be related to the known effects of MICA/MICB shedding: a reduction in cell surface density of NKG2D-Ls leading to a reduced susceptibility to NKG2D-mediated cytotoxicity and systemic downregulation of NKG2D on NK cells and CD8+ T cells in cancer patients. These considerations suggest that either blockade of MIC release or neutralization of shed soluble MIC would be a useful addition to immunologic approaches for cancer therapy. In this context, an understanding of the
mechanisms involved in the release of soluble MICA/MICB molecules from tumor cells is crucial for the development of effective strategies to block the shedding of these proteins. ADAM17/TACE has been identified as a key protease mediating proteolytic cleavage of both MICA and MICB molecules (16, 17). However, the data in this article show that MICA molecules can be released from a tumor cell in more than one way. The MICA*008 molecule, that is by far the most frequent allele of MICA in multiple populations (18–22), is not released from cells by proteolysis but rather as a membrane-anchored molecule in exosomes. Thus, effective blockade of the accumulation of soluble MICA in patient sera will require different strategies depending on the MICA alleles expressed by the individual patient.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

We thank S.J. Powis (University of St. Andrews), P. Roda-Navarro, Dr. F. Colucci and his group, and Dr. A. Kelly for helpful discussions; Prof. J. Trowsdale for critical reading of the manuscript; and N. Miller for assistance with cell sorting.

Grant Support

Medical Research Council and Leukemia Research Fund. M. Vañes-Gómez is a recipient of a New Investigator Award Grant from the Medical Research Council. O. Ashiru and P. Boutet were partially supported by The Newton Trust. S. Agüera-González was supported by Fundación Caja Madrid and Ibercaja.

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Received 5/7/09; revised 10/20/09; accepted 11/9/09; published OnlineFirst 1/12/10.

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Exosomally Released MICA*008 Inhibits NK Cell Recognition


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