Human melanoma-specific CD8+ T-cells from metastases are capable of antigen-specific degranulation and cytolysis directly ex vivo

Yolanda D. Mahnke,1,† Estelle Devêvre,1,† Petra Baumgaertner,1 Maurice Matter,1 Nathalie Rufer,1,3 Pedro Romero1,4 and Daniel E. Speiser1,4, *

1 Ludwig Center for Cancer Research; University of Lausanne; Lausanne, Switzerland; 2 Department of Surgery; University Hospital; Lausanne, Switzerland; 3 Department of Research; University Hospital; Lausanne, Switzerland; 4 National Center for Competence in Research; Molecular Oncology; Epalinges, Switzerland

† These authors contributed equally to this work.

Keywords: interferon gamma, degranulation, cytotoxicity, exhaustion, human tumour immunology

The relatively low frequencies of tumor Ag-specific T-cells in PBMC and metastases from cancer patients have long precluded the analysis of their direct ex vivo cytolytic capacity. Using a new composite technique that works well with low cell numbers, we aimed at determining the functional competence of melanoma-specific CD8+ T-cells. A multiparameter flow cytometry based technique was applied to assess the cytolytic function, degranulation and IFNγ production by tumor Ag-specific CD8+ T-cells from PBMC and tumor-infiltrated lymph nodes (TILN) of melanoma patients. We found strong cytotoxicity by T-cells not only when they were isolated from PBMC but also from TILN. Cytotoxicity was observed against peptide-pulsed target cells and melanoma cells presenting the naturally processed endogenous antigen. However, unlike their PBMC-derived counterparts, T-cells from TILN produced only minimal amounts of IFNγ, while exhibiting similar levels of degranulation, revealing a critical functional dichotomy in metastatic lesions. Our finding of partial functional impairment fits well with the current knowledge that T-cells from cancer metastases are so-called exhausted, a state of T-cell hyporesponsiveness also found in chronic viral infections. The identification of responsible mechanisms in the tumor microenvironment is important for improving cancer therapies.

Introduction

A central issue in tumor immunology is that tumor cells do not express foreign antigens and are therefore not readily fought against by the immune system. Additionally, cancer cells have proven to be experts in immune evasion, making use of a wide repertoire of strategies to elude its adversaries. Nevertheless, tumor Ag-specific T-cells can be readily detected in PBMC of cancer patients, and it has been shown that such T-cells can play a significant role in delaying tumor progression or even eliminate cancer cells in murine model systems,1,2 as well as in cancer patients.3 Ag-specific reactivity is typically demonstrated by IFNγ ELISPOT assays.4 PBMC-derived tumor Ag-specific T-cells have been shown to secrete IFNγ in an Ag-specific manner.5,6 Solid tumors of peripheral tissues are infiltrated with tumor Ag-specific T-cells, and in tumor-infiltrated lymph nodes (TILN, also referred to as metastases hereafter) the tumor cells are also surrounded by immune cells that could potentially attack them. However, it has previously been reported that tumor Ag-specific CD8+ T-cells from melanoma metastases show impaired IFNγ production directly ex vivo, indicating a functional defect or inhibition at the tumor site.5,7 Importantly, IFNγ production is not necessarily representative for other functions of CD8+ T-cells. While IFNγ production was strongly impaired, we found that Perforin production was only marginally reduced and Granzyme B content appeared normal in Ag-specific T-cells from melanoma metastases.5,9 Based on these findings, we were interested in the ex vivo cytolytic capacity of tumor-specific CD8+ T-cells from cancer patients, as the lysis of tumor cells is important for successful tumor defense. Target cell lysis can be achieved through different mechanisms: (1) release of lytic granule contents (Granzymes, Perforin and Granulysin), (2) cell-cell contact (Fas/FasL) and (3) cross-linking of death-receptors (TNFα/TNFRα Type I).10 Since a large proportion of Melan-A (also known as MART-1) specific CD8+ T-cells from cancer patients express Perforin and Granzyme B,11 we hypothesized that their lytic activity would be mainly granule-dependent. Upon granule exocytosis, the lysosomal-membrane associated glycoprotein CD107a (also known as...
LAMP-1) is transferred to the plasma membrane. Cell surface detection of CD107a therefore serves as a read-out for granule exocytosis, and it is closely, though not absolutely, linked to lytic activity.

Hence, to further investigate the ex vivo functionality of tumor-specific CD8⁺ T-cells as well as the apparent functional impairment of TILN-derived cells, we assessed the ex vivo lytic capacity, degranulation and IFNγ production of Melan-A-specific CD8⁺ T-cells from PBMC and TILN of melanoma patients. To this end, we made use of our multiparameter LiveCount Assay (LCA), which was slightly modified (mLCA) to allow the concomitant analysis of lytic activity and surface CD107a, as in the original protocol, as well as intracellular IFNγ. In addition to detecting IFNγ production on a per cell basis, the mLCA does not require pre-purification of Ag-specific cells, thus reducing procedure-related potential influences on their viability and/or functionality.

While circulating T-cells showed strong cytolytic capacity, degranulation and IFNγ production, the results indicate a partial inhibition of T-cell functions at the tumor site, underlining that T-cells sampled from the PBMC do not always reflect functionality at the site of disease. Interestingly, the T-cells from metastases showed considerable lytic activity.

**Results**

Direct ex vivo cytotoxicity by Melan-A-specific CD8⁺ T-cells from PBMC or TILN of vaccinated melanoma patients. Melan-A-specific T-cells were isolated from five PBMC and five TILN samples and tested by ex vivo LCA in triplicates. These T-cells showed strong cytolytic function ex vivo. Though TILN-derived T-cells exhibited lower levels of cytolytic activity than those derived from PBMC (Fig. 1A and B), the difference was not statistically significant. Efficient cytotoxicity by TILN-derived T-cells was remarkable, given previous reports showing hyporesponsiveness of T-cells in metastases. In parallel to the assessment of primary T-cells, we also used a Melan-A-specific CTL clone (clone R11) in each experiment in order to assess inter-assay variability. The clone was consistently used 13 to 16 d after the periodic re-stimulation with

![Figure 1. Melan-A-specific T-cells from both PBMC and TILN of melanoma patients exert strong cytotoxicity directly ex vivo. Melan-A-specific CD8⁺ T-cells were FACS purified from PBMC (A); n = 5) and TILN (B); n = 5) of melanoma patients using pMHC multimers, and incubated for 4 h with an equal number of Melan-A-pulsed T2^{CMN} and HIV-pulsed T2^{CMN} at the indicated E:T ratios. All conditions were measured in triplicates. A semi-paired permutation test yielded no statistically significant difference between the cytolytic capacity of cells from PBMC and TILN (p = 0.2). (C) Pre-incubation with pMHC multimers does not significantly affect the lytic activity of CD8⁺ CTL clones. CTL clones specific for either Melan-A, Flu, CMV or EBV were left untreated (white bars) or incubated with relevant pMHC multimers (black bars) prior to incubating them for 4 h with an equal number of relevant-peptide-pulsed T2^{CMN} and irrelevant-peptide-pulsed T2^{CMN}. E:T = 4 was used for all clones except NM cl.55, which was assayed at E:T = 1. Mean cytolytic activity and SD of quadruplicates are shown for each clone. A paired, two-tailed t-test showed no difference between pMHC multimer treated or untreated cells (p = 0.94).](image)
PHA, such that the clone was always at a similarly high activation state. Indeed, results from clone R11 revealed favorable assay reproducibility (data not shown). In summary, these results show a surprisingly strong and consistent lysis of peptide-pulsed T2 cells by tumor Ag-specific T-cells isolated directly from the patients’ PBMC or TILN, comparable to the lytic activity of a CTL clone with strong cytolytic function.

The modified LiveCount Assay (mLCA) and original LCA provide comparable results. It could be argued that the vigorous cytotoxicity of patients’ T-cells was due to TCR triggering by peptide/MHC (pMHC) multimers used for their sorting by flow cytometry. To address this possibility, we tested CTL clones of different specificities and found similar levels of cytotoxicity, whether they were pre-incubated with pMHC multimers or not (Fig. 1C). Regardless, sorting by flow cytometry may alter cell viability and lytic performance in primary cells. Therefore, we optimised our assay to circumvent prior labeling and sorting of T-cells. This modified LiveCount Assay (mLCA) was validated with CTL clone R11 and produced similar titration curves in terms of specific lysis and CD107a upregulation as the original LCA (data not shown). Characteristics and differences of the two assays are listed in a table (Table S1, which demonstrates the differences between LCA and mLCA). The mLCA was then applied to assess the ex vivo cytotoxicity of Melan-A-specific T-cells from PBMC of three melanoma patients (Fig. 2A). Efficient cytotoxicity was detected, with E:T titration curves comparable to those obtained with the original LCA (Fig. 1A). Thus, considerable cytotoxicity was detected ex vivo, even without prior pMHC multimer based T-cell sorting.

Enhanced activation of Melan-A-specific CD8+ T-cells at low E:T ratios. In addition to the simultaneous assessment of cytotoxicity and degranulation, the mLCA allows to detect cytokine-producing cells (Fig. 2C and F), hence providing a more comprehensive evaluation of T-cell function. Although lysis was not detectable at low E:T ratios, the majority of Melan-A-specific T-cells

![Figure 2](image-url)

Figure 2. The modified LiveCount Assay (mLCA) allows to simultaneously determine cytotoxicity, degranulation and IFNc production. CD8+ T-cells magnetically enriched from PBMC of melanoma patients were incubated with peptide-pulsed T2 target cells (A–C) or the endogenously Melan-A+ melanoma cell line Me 290 (D–F) at the indicated E:T ratios in the presence of fluorescently conjugated CD107a mAb. Brefeldin A was added after 1 h of co-incubation. After a further 3 h of co-incubation, cells were incubated with Melan-A/HLA-A*0201 multimers, CD8 mAb, VIVID and, following fixation and permeabilisation, IFNc mAb. (A and D) % specific lysis. (B and E) Degranulation as measured by CD107a upregulation on the cell surface. (C and F) IFNc production. Results for peptide-loaded T2 and Me 290 melanoma target cells were compared using a semi-paired permutation test, yielding p values of 0.007 (lysis), 0.34 (CD107a upregulation) and 0.50 (IFNc expression). (G) Mean fluorescence intensity (MFI) of cell surface CD107a staining at increasing E:T ratios with peptide-loaded T2 as target cells. The graph shows mean ± SEM of 5 PBMC and 5 TILN samples. (H) The scheme illustrates that CD107a and cytokine expression is reduced on a per cell basis in conditions of limited availability of specific target cells, while the overall specific lysis is increased.
degranulated under such conditions, with a mean of about 70% at E:T = 0.06 (Fig. 2B). With increasing E:T ratios, frequencies of CD107a + cells decreased to values as low as 11% at E:T = 16. The proportion of IFNγ + T-cells mirrored this observation. At E:T = 0.06, an average of 51% of Melan-A-specific T-cells were IFNγ +, but only a mean of 12% produced IFNγ at E:T = 16 (Fig. 2C). Similarly, the mean fluorescence intensity of cell surface CD107a detection, reflecting the average molecular density of this molecule per cell, decreased with increasing E:T ratios and increasing lytic activity (Fig. 2E). This suggests that at elevated E:T ratios, the cells that are participating in killing do not degranulate as much on a per-cell basis as seen at lower E:T ratios (Fig. 2H). Together, our data demonstrate that the majority of Melan-A-specific T-cells are potentially functional, and that different functions are revealed at different E:T ratios. Since cytotoxicity is measured as the result of a collective T-cell action and not on a per T-cell basis, degranulation and IFNγ production are more accurate measurements of activation of individual T-cells.

Successful recognition and killing of melanoma cells. To test the ability of Melan-A-specific T-cells to recognize melanoma cells based on their endogenous production and presentation of antigen, we compared the T-cell functions generated in response to Melan-A+ (Me 290) and Melan-A− (NA8-MEL) melanoma cell lines. Melan-A-specific T-cells directly isolated from PBMC of melanoma patients were able to lyse melanoma cells (Fig. 2D). Successful recognition of melanoma cells in the absence of synthetic peptide was also demonstrated by the T-cells’ ability to degranulate and produce IFNγ (Fig. 2E and F). However, lysis of unpulsed melanoma cells (Fig. 2D) was often less efficient than lysis of peptide-pulsed T2 cells (Fig. 2A), presumably due to lower levels of peptide being presented by tumor cells. This was not statistically significant (p = 0.08) if taking only mLCA data into account, probably due to low sample numbers, but did reach statistical significance when integrating the LCA data from Figure 1A (p = 0.007). Strikingly, stimulating with Me 290 revealed different response patterns generated by PBMC from different patients. While some delivered strong (LAU 1013) or weak (LAU 936) responses in all three parameters analyzed, others showed strong lysis with weak degranulation and IFNγ production (LAU 1146) or vice versa (LAU 1106).

Low IFNγ production by Melan-A-specific CD8+ T-cells from TILN. We compiled the data obtained with T-cells from PBMC and TILN, as assessed with both the LCA and the mLCA. In concurrence with our previous studies,18 the percentage of IFNγ + Melan-A-specific T-cells was consistently lower in TILN compared with PBMC (p = 0.0004, Figure 3A and B). In contrast, T-cells from TILN showed high ex vivo cytotoxicity and degranulation, similar to those displayed by the T-cells from PBMC (Fig. 3B). Comparison of lytic activity was performed at E:T = 4, while degranulation and IFNγ production were evaluated at E:T = 0.5. These E:T ratios were chosen as they yielded high, but not maximal, readings (Figs. 1 and 2), and the standard deviation due to experimental conditions was at a minimum (see Table S1, which shows Comparison of LCA with mLCA data). Moreover, the relative frequency of cells expressing either IFNγ, surface CD107a or both was significantly different between the two organs, with the bulk of TILN-derived cells producing either function being single-positive for CD107a, while PBMC-derived cells were largely double-positive (Fig. 3C). In conclusion, ex vivo assessment of T-cells from metastatic lesions consistently reveals deficient IFNγ responses.

Apparent discrepancy between functional parameters. Even though Melan-A-specific CD8+ T-cells from TILN produced only low levels of IFNγ ex vivo, their IFNγ production correlated with degranulation (CD107a; Figure 4A). A similar trend was observed for PBMC. No correlation was observed between ex vivo IFNγ production and lytic capacity, underlining that IFNγ alone is not sufficient to determine a sample’s reactivity (functionality vs. exhaustion, Figure 4B). Interestingly, an inverse correlation was observed between degranulation and lytic activity (Fig. 4C). This indicates that samples with high ex vivo lytic activity might contain Melan-A-specific cells with high avidity. Thus, only a few (high avidity) cells of the total Melan-A-specific CD8+ T-cell population would be involved in target cell killing, resulting in low frequencies of CD107a + cells. In contrast, more (lower avidity) cells are involved in Ag-specific lysis in samples with lower lytic activity, hence the elevated frequencies, of CD107a + cells.

Increased Ag-reactivity following peptide vaccination. We were able to perform LCAs with longitudinal PBMC samples as well as one TILN from a melanoma patient positive for the cancer/testis Ag NY-ESO-1 (Fig. 5), before and after peptide vaccination.19 Samples obtained pre-vaccination revealed strong ex vivo cytolytic activity, but only very weak degranulation in response to NY-ESO-1–pulsed T2 cells. NY-ESO-1-specific CD8+ T-cells from a PBMC sample drawn after one year of monthly peptide vaccinations had even higher ex vivo cytolytic potential, and a significantly increased degranulation response.

Discussion

In this study we simultaneously determined cytotoxicity, degranulation and cytokine production of human cancer-specific T-cells directly ex vivo, using the original and modified LiveCount Assays. Strong cytolytic activity was observed not only by circulating T-cells, but also by T-cells from melanoma metastases in lymph nodes, a remarkable result given previous reports of functional impairment of cancer-specific T-cells in metastases.14,17,18 T-cells readily killed peptide-pulsed target cells in an Ag-specific manner, to a similar extent as a representative and strongly cytolytic CD8+ CTL clone. We also found efficient recognition and killing of melanoma cells presenting naturally processed antigen. However, killing, degranulation and IFNγ production by T-cells differed considerably between patients, emphasizing the inter-individual differences in T-cell functionality, which possibly reflects, at least in part, differences in TCR avidities14 and cytotoxic mechanisms employed by individual T-cells.

Competent killing activity by T-cells from metastases is in agreement with the notion that cytolytic function often remains intact, despite defects in cytokine production, as observed in
exhausted T-cells in chronic/protracted viral infections in mouse models (45) and in HIV and HCV patients. Recently, we and others have demonstrated that CD8+ T-cells from melanoma metastases are exhausted, as their gene expression profile overlapped significantly with the one of exhausted T-cells in chronic viral infections. T-cell exhaustion was originally described in mice infected with particular strains of lymphocytic choriomeningitis virus (LCMV) strains causing chronic infections. These mice generated LCMV-specific CD8 T-cells that are dysfunctional and cannot clear the virus. T-cell exhaustion is defined by the hierarchical loss of cytokine function of effector T-cells, first with the loss of IL-2, then TNFα and finally IFNγ. While some exhausted T-cells are driven into apoptosis, large numbers remain vital. The question whether exhausted T-cells are capable of cytolysis is still open.

Numerous studies have analyzed T-cell function after prolonged in vitro culture, but such studies do not necessarily reflect in vivo function. In mice, where T-cell functions can be studied in vivo, CD8+ effector T-cells are highly cytolytic against target cells occurring as single loosely associated cells in the tissues. To some extent, CD8+ effector T-cells are also capable of attacking solid tumors, though in vivo imaging studies have shown that target cell lysis in solid tumors may be very slow and that T-cell function may be inhibited by environmental factors such as TGFβ. Indeed, several factors of the tumor microenvironment have been found to inhibit T-cells. Partially deficient T-cell function has been described more than 20 y ago, in the context of murine thymic T-cell maturation. More recently, Ohlen et al. described tumor Ag-specific T-cells with normal effector function but deficient proliferation in a mouse model. Partial T-cell function may be due to deficient TCR signaling and/or co-stimulation, which must be strong for proliferation, intermediate for cytokine production and only weak for target cell lysis. Thus, our findings may be caused by environmental factors such as TGFβ. Indeed, several factors of the tumor microenvironment have been found to inhibit T-cells. Partially deficient T-cell function has been described more than 20 y ago, in the context of murine thymic T-cell maturation. More recently, Ohlen et al. described tumor Ag-specific T-cells with normal effector function but deficient proliferation in a mouse model. Partial T-cell function may be due to deficient TCR signaling and/or co-stimulation, which must be strong for proliferation, intermediate for cytokine production and only weak for target cell lysis. Thus, our findings may be caused

Figure 3. TILN-derived melanoma-specific CD8+ T-cells exhibit decreased IFNγ production, but comparable degranulation to their PBMC-derived counterparts. (A) Representative dot plots for degranulation and IFNγ production by PBMC- (LAU 944) and TILN-derived (LAU 465) Melan-A-specific CD8 T-cells. (B) Frequencies of Melan-A-specific T-cells positive for (IFNγ, CD107a at E:T = 5:1) and % specific lysis at E:T = 4 is indicated. Data are means of triplicates. E:T Data for PBMC and TILN were compared using unpaired, two-tailed t-tests. (C) Pie show the relative proportion of Melan-A-specific CD8+ T-cells positive for either IFNγ and/or cell surface CD107a expressing either combination of these markers.
Figure 4. Apparent discrepancy between functional parameters. Correlation analysis of functional parameters measured following Ag-specific stimulation of Melan-A-specific cells isolated from either PBMC (open circles) or TILN (closed triangles). (A) CD107a vs. IFN-γ. (B) IFN-γ vs. % specific lysis. (C) CD107a vs. % specific lysis. The outliers highlighted in gray have been excluded from the statistical analysis of CD107a vs. % specific lysis, as they had very low lytic capacity.

Figure 5. Peptide-vaccination results in increased degranulation and cytolytic activity of tumor Ag-specific CD8+ T-cells. A longitudinal study was performed with NY-ESO-1-specific CD8+ T-cells from a melanoma patient. PBMC samples were collected 11 and 3 mo before, as well as 12 mo after commencing peptide vaccination. TILN were also obtained 3 mo pre-vaccination. (A) Cytolytic activity and (B) degranulation in response to incubation with NY-ESO-1 peptide-pulsed T2 cells were determined.
by partially deficient TCR and co-stimulatory signals, still sufficient for lysis but not for cytokine production. Furthermore, persistent Ag has been shown to result in impaired nuclear translocation of NFAT proteins, which in turn could negatively impact IFNγ production. In any case, the molecular mechanisms underlying the observed functional deficits of T-cells from metastases need to be addressed, and the responsible tumor-derived suppressive mechanisms identified. IFNγ is known to be essential for tumor defense, and so the lack of IFNγ in TILN is important. Tumor Ag-specific CD8+ T-cells are known to also exist in tumor-free lymph nodes. However, as frequencies of Ag-specific T-cells are very low in such patient samples, we were not able to test whether such cells behave like their counterparts found in TILN or PBMC.

Cytokine production and degranulation assays are complementary to cytotoxicity assays, with the former investigating samples, we were not able to test whether such cells behave like their counterparts found in TILN or PBMC. Our LCA analysis of four such samples (with pre- and post-vaccination time-points) from a single melanoma patient indicated an increase not only in the frequency, but also in the ex vivo detectable degranulation and lytic activity of the tumor Ag-specific T-cells.

Recognition of tumor cells presenting naturally processed tumor antigen is critical, but often not the case, presumably due to low amounts of naturally presented tumor antigen and low avidity of self-specific TCRs. The fact that T-cells were capable of killing melanoma cells in the absence of synthetic peptide suggests that TCR-affinities were relatively high, corresponding to our previous findings. However, tumor cell recognition varied between patients. In contrast to T-cells from patient LAU 1164, T-cells from patient LAU 1106 lysed MHC 290 cells much less efficiently than peptide-pulsed T2 cells, suggesting that the former patient had better TCRs and consequently required lower amounts of antigen for efficient tumor recognition. In the future, comprehensive analyses of TCR repertoire and T-cell function will improve the characterization of T-cell responses. Recently, ex vivo assessment of cytokine production and cytotoxicity has become possible for groups of Melan-A/HLA-A*0201 (see Table S2) and even individual clonotypes, as we have demonstrated in a strong NY-ESO-1-specific T-cell response of a melanoma patient. As T-cell avidity and functionality are key for immune protection, such studies have the potential to reveal correlates of protection against disease, providing the rationale to improve T-cell based immunotherapies.

**Patients and Methods**

Patients and biological specimens. PBMC and TILN were obtained from 14 HLA-A2+ melanoma patients who gave informed consent for this study, approved by the ethical committee of the University of Lausanne and the Ludwig Institute for Cancer Research. PBMC from all patients except patient LAU 50 were analyzed after peptide vaccination with Melan-A peptide, CpG 7909 and IFA. Patients LAU 465 and LAU 969 had not received vaccination at the time of TILN isolation, while LAU 672 and LAU 818 had received Melan-A peptide in IFA and LAU 352 had received Melan-A peptide in AS02. Two PBMC draws and one TILN were obtained from patient LAU 50 before commencing administration of NY-ESO-1, Mage-A10 and Melan-A peptides in Montanide. One additional post-vaccination PBMC sample was also included in this study. Ex vivo mononuclear cells were purified and immediately frozen as described previously. Frequencies of Melan-A/HLA-A*0201 (see Table S2, which shows frequencies and phenotype of Melan-A-specific CD8+ T-cells from PBMC and TILN) or NY-ESO-1/HLA-A*0201 multimer binding CD8+ T-cells were determined by flow cytometry.

CD8+ CTL clones. The CD8+ CTL clones NM c.55 (Influenza-specific), BC25 c.10 (EBV-specific), c.10 (Melan-A-specific), c.48 (both CMV-specific) were isolated from healthy donors, while LAU 465 c.11ES (Melan-A-specific) and LAU 50 c.1 (NY-ESO-1-specific) were isolated from melanoma patients as follows: peptide/MHC multimer binding CD8+ T-cells were purified by FACS from PBMC, cloned by limiting dilution and periodically (every 3–4 weeks)
expanded by stimulation with 1 μg PHA/ml (Sodiagu, 5 × 10^6) irradiated allogeneic feeder cells/ml and 1500U IL-2/ml (Glaxo Welcome). Cells were used 13–16 d after PHA-stimulation.

**Target cell lines.** The T2 cell line, a human TAP-deficient Tdb cell hybrid expressing low levels of surface HLA-A*0201 and virtually no other surface class I MHC molecules, was peptide-pulsed to serve as a target. The melanoma-derived cell lines Me 290 (Melan-A*-A*0201) and NA8-MEL (Melan-A*-A*0201) served as an irrelevant cell line and a target cell line. Me 290 was kindly provided by D. Rimoldi (Ludwig Institute for Cancer Research, Lausanne Branch, Switzerland), and their identity established by testing for the melanoma Ags Melan-A, Ty, gp100 and HMW-MAA. NA8-MEL was a kind gift from F. Jotereau (INERIM); it expresses the common melanoma mutation in BRAF. All cell lines are routinely monitored for morphology, growth characteristics and the absence of mycoplasma. Since they are often used in cytolytic assays, potential alterations in their performance and hence their identity would be readily identified.

**Media and buffers.** Assays were performed in RPMI 1640 containing 100 U penicillin/ml, 100 μg streptomycin/ml, 2 mM glutamine, 0.1 mM non-essential amino acids, 1 mM Na pyruvate (all from Gibco, Invitrogen), 5×10^5 M β-ME (Sigma) and 8% human serum (HS, CTS) (subsequently referred to as RPMI 8% HS). The T-cell clones were maintained in RPMI 8% HS enriched with 150 IU IL-2/ml (Roche Pharma). T2 cells and melanoma cell lines were cultured in RPMI 1640 with 0.24 mM l-arginine, 0.55 mM l-asparagine and 5 mM sodium azide (all from Sigma). 1.5 mM glutamine, 10 mM Hepes, 100U penicillin/ml, 100 μg streptomyacin/ml and 10% FCS (Gibco). Staining buffer was prepared in triplicates. Two series of experiments were performed. The first series was done at E:T 0.25 to 4 or 0.25 to 8 were acquired on a FACS Vantage SE and analyzed with the instrument’s operating software (CellQuest Pro, BD Biosciences). Specific lysis was calculated as originally published.

**Biostatistical analyses.** The areas under the curves between E:T ratios 0.25 to 4 or 0.25 to 8 were contrasted with the Mann-Whitney U test.

**Molecular Probes.** DAPI (358 ex 461 em), or the UV-excitable DNA stain DAPI (358 ex 461 em), and the fluorescent conjugates, were dispensed into the wells of a 96-well plate containing 125 or 250 cells of each target cell population to obtain the indicated E:T ratios. Control wells received only the target cell mixture in order to correct for eventual variations in target cell counts. All conditions were prepared in triplicates. Cells were incubated in 50 μl RPMI 8% HS containing CD107a/C190 mAb and after 3.5 h a fluorescently conjugated CD8 mAb was added. At the end of the 4h co-incubation period, cells were harvested, pelleted and re-suspended in 100 μl staining buffer containing 3 μg/ml DAPI. Samples were acquired on a FACSVantage SE and analyzed with the instrument’s operating software (CellQuest Pro, BD Biosciences). Specific lysis was calculated as originally published.

**Modified LiveCount Assay (mLCA).** T2 cells and/or melanoma cell lines were labeled with 0.125μm CFSE. T2 cells were pulled with Melan-A peptide or left unpulsed. All targets were labeled with 0.125mM CFSE. T2 cells were pulsed with Melan-A peptide or left unpulsed. All targets were pulsed with Melan-A peptide or left unpulsed. All targets were pulse-pulsed. The protein transport inhibitor Brefeldin A was added. At the end of the 4h co-incubation period, the cells were transferred to tubes and incubated with Melan-A/HLA-A*0201 multimers. Samples were washed and re-suspended in 100 μl staining buffer containing ViViD prior to acquisition on a FACSVantage SE.
calculated, and a semi-paired permutation test adapted. This test uses a traditional unpaired t-test, but compares it to a corrected permutation distribution in which pair or the lack thereof is taken into account. To compare the lytic activity generated by pMHC multimer pre-treated vs. untreated CD8⁺ T-cell clones was paired, semi-paired t-test was compared using the permutation analysis in Spice 5.1. The correlation analyses were performed in JMP 8.0 (SAS). All other comparisons were performed using an unpaired, two-tailed t-test.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

Acknowledgments
We gratefully acknowledge the patients’ participation, as well as the hospital staff’s excellent collaboration. We thank A. Krieg, H. Bouzourene, J.-C. Corotini, C. Gnegeler, J. Laurent, H.-A. Lehr, S. Leyvra, O. Michielin, B. Niklaus, and V. Veelken for collaboration, I. Luescher and P. Guillaume for pHMC multimers, Seppic for Montanide ISA-51 (IFA), and Coley Pharmaceutical Group for CpG 7909. Special thanks go to M. Nason for biostatistical support. Excellent technical and secretarial help was received from C. Beavard, C. Geldhof, and N. Montandon.

Y.M., E.D., and P.S. were involved in the conception and experimental design of the study. M.M. and D.S. performed clinical supervision and sampling of patients, P.B. and N.R. performed the experimental work and together with D.S. analyzed the data. Y.M., E.D. and D.S. wrote the manuscript while all authors were involved in manuscript review and editing.

This work was supported in part by the Deutsche Forschungsgemeinschaft (DFG) grant MA2595/2-1 (Y.D.M.), the Lausanne Branch of the Ludwig Institute for Cancer Research (E.D. and P.B.), the University Hospital, Lausanne (M.M.), the Swiss National Science Foundation (SNF) and the National Center of Competence in Research (NCCR) (D.S. and N.R.), and grants from the Fondation Solidar-Immun (D.S.) and from the FWF EU “Cancerimmunotherapy” (P.R.).

Supplemental Material
Supplemental materials may be found here:
www.landesbioscience.com/journals/oncoimmunology/article/198560
37. Bachmann MF, Sebzda E, Kündig TM, Shahinian A, Ohlén C, Kalos M, Cheng LE, Shur AC, Hong DJ,
36. Zou W. Immunosuppressive networks in the tumour
35. Ochsenbein AF, Sierro S, Odermatt B, Pericin M, Karrer
34. Pittet MJ, Grimm J, Berger CR, Tamura T,
33. Breart B, Lemaître F, Celli S, Bousso P. Two-photon
32. Agnellini P, Wolint P, Rehr M, Cahenzli J, Karrer U,
31. dx.doi.org/10.1002/eji.1830260908
30. Carson BD, et al. CD8(+) T cell tolerance to a tumor-
29. Ogg GS, et al. Peptide vaccination for therapy. Proc Natl Acad Sci U S A 2007; 104:12457-
28. Pittet MJ, Grimm J, Berger CR, Tamura T,
27. Walker BD, Fonteneau JF, Cerottini JC, et al. Activation of human
26. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
25. So T, Hanagiri T, Chapiro J, Colau D, Brasseur F,
24. So T, Hanagiri T, Chapiro J, Colau D, Brasseur F,
23. Ayyoub M, Zippelius A, Pittet MJ, Rimoldi D,
22. Ayyoub M, Zippelius A, Pittet MJ, Rimoldi D,
21. Ayyoub M, Zippelius A, Pittet MJ, Rimoldi D,
20. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
19. Blankenstein T. The role of tumor stroma in the
18. Price DA, Brenchley JM, Ruff LE, Betts MR, Hill BJ,
17. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
16. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
15. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
14. Zou W. Immunosuppressive networks in the tumour
13. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
12. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
11. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
10. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
9. Blankenstein T. The role of tumor stroma in the
8. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
7. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
6. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
5. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
4. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
3. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
2. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,
1. Romero P, Dunbar PR, Valmori D, Pittet M, Ogg GS,