Modulation of Microenvironment Acidity Reverses Anergy in Human and Murine Tumor-Infiltrating T Lymphocytes

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Abstract
Stimulating the effector functions of tumor-infiltrating T lymphocytes (TIL) in primary and metastatic tumors could improve active and adoptive T-cell therapies for cancer. Abnormal glycolysis, high lactic acid production, proton accumulation, and a reversal of an intracellular pH gradient are thought to help render tumor microenvironments hostile to roving immune cells. However, there is little knowledge about how acidic microenvironments affect T-cell immunity. Here, we report that lowering the environmental pH to values that characterize tumor masses (pH 6–6.5) was sufficient to establish an anergic state in human and mouse tumor-specific CD8⁺ T lymphocytes. This state was characterized by impairment of cytokytic activity and cytokine secretion, reduced expression of IL-2R (CD25) and T-cell receptors (TCR), and diminished activation of STAT5 and extracellular signal–regulated kinase (ERK) after TCR activation. In contrast, buffering pH at physiologic values completely restored all these metrics of T-cell function. Systemic treatment of B16-OVA–bearing mice with proton pump inhibitors (PPI) significantly increased the therapeutic efficacy of both active and adoptive immunotherapy. Our findings show that acidification of the tumor microenvironment acts as mechanism of immune escape. Furthermore, they illustrate the potential of PPIs to safely correct T-cell dysfunction and improve the efficacy of T-cell–based cancer treatments.

Introduction
Several lines of evidence suggest that T-cell immunity may play a role in controlling tumor development. Tumor-infiltrating T lymphocytes (TIL) with an effector/memory phenotype are associated with a more favorable prognosis in patients with cancer (1) and mediate tumor regression when adoptively transferred after ex vivo activation (2). However, TILs are reported to acquire functional defects at the tumor site and enter a state of reversible anergy, mostly attributed to suboptimal T-cell receptor (TCR) ligation, a lack of TCR and CD8 colocalization, or inhibitory signaling through CTLA4 (3–5). This may, at least in part, explain the clinical observation that TILs can be abundantly detected in progressing cancer lesions (6).

An issue frequently neglected relates to the metabolic features of the tumor microenvironment, which represents a potentially hostile milieu due to the hypoxic and acidic conditions that occur as a consequence of inadequate blood flow, inflammation, enhanced glycolysis (the so-called Warburg effect), and the production of metabolic acidosis (7). In addition, excessive CO₂ release caused by altered glycolysis leads to increased tumor expression of carbonic anhydrase IX (CA-IX), thus contributing to further acidification of the extracellular tumor environment (8, 9). Hypoxia also activates the hypoxia-inducible factor (HIF) pathway, which in turn upregulates glucose transporters and CA-IX and leads to additional exacerbation of tumor acidosis (8). As a result, the extracellular pH can drop to values of 6.0 or less and is on average 0.2 to 1.0 units lower than that of normal tissues (10, 11). This biochemical shift has been shown to provide a selective growth advantage to tumor cells (7) at the expense of infiltrating host cells, including immune cells. While the impact of these metabolic alterations on TILs is presently unknown, the clinical evidence that metabolic acidosis is often associated with immunodeficiency (12), and that both leukocyte activation and the bactericidal capacity of leukocytes are generally impaired at reduced pH (13), suggests that T cells could be extremely sensitive to pH variations.
Here, we report that relatively minor changes in the extracellular pH promote reversible anergy associated with impaired effector functions in both human and mouse tumor-specific CD8+ T lymphocytes. Buffering low pH at the tumor site by in vivo administration of the proton pump inhibitor (PPI) esomeprazole improved TIL efficacy and delayed cancer progression in tumor-bearing mice. These data highlight tumor acidity as a novel mechanism of immune escape that could be targeted for rescuing effective tumor immunity and achieving disease control in patients with cancer.

Materials and Methods

Phenotypic and functional studies of human TILs
TILs were isolated and cultured as previously described (14), stained with the indicated monoclonal antibodies (mAb; BD Biosciences) and analyzed with a FACSort (BD Biosciences) and FlowJo Software (Tree Star Inc.). TILs were assessed for tumor recognition by IFNγ ELISpot assays (15). TIL were stimulated with Dynabeads CD3/CD28 T Cell Expander (1:20 ratio; Invitrogen Dynal AS) for 72 hours at the indicated pH and then assessed for proliferation by carboxyfluorescein succinimidyl ester (CFSE; CellTrace CFSE Cell Proliferation Kit; Molecular Probes). For cytokine secretion analysis, TILs were cultured at the indicated pH for 3 days and then stimulated with CD3/CD28 T Cell Expander beads (1:20 ratio) or with autologous tumor cells (8:1 ratio). Twenty-four-hour supernatants were tested using FlowCytomix Human Th1/Th2 multiplex kit (Bender MedSystems). Data were analyzed by FlowCytomix Pro 2.2 software (Bender MedSystems). The IL-2R α-chain was analyzed using an anti-CD25 mAb (BD Biosciences) on TILs stimulated for 24 hours with Dynabeads CD3/CD28 beads (1:20 ratio), CD3-ζ-chain and perforin levels (detected after 6 hours of incubation with autologous tumor cells, 8:1 ratio) were evaluated by flow cytometry. For Annexin-V/propidium iodide (PI) studies, TILs were cultured for 3 days at the indicated pH and then stained with the rhAnnexin V-FITC Kit (Bender MedSystems). For intracellular phospho-protein analysis, TILs were stimulated with an anti-CD3 mAb (2 μg/mL, 5 minutes; OKT3, Orthoclone; Janssen-Cilag) or IL-2 (150 ng/mL, 30 minutes; Proluken, Chiron Corp.). Cells were then fixed, permeabilized, and stained with anti-ERK (pT202/pY204) mAb. Data were analyzed by FlowJo Software (Bender MedSystems).

No detrimental effect of low pH on either mAb binding activity or cytokine detection was observed. Melanoma cell lines, authenticated and characterized according to UKCCCR guidelines (16), were obtained from patients after informed consent and ethical committee approval.

Animals and tumor cell lines
C57BL/6 (Charles River) and C57BL/6-Tg(TcraTcrb) 425Cbn/J (OTI) female mice (17) were treated in accordance with the European Community guidelines. B16F1 melanoma (American Type Culture Collection; ATCC) and RMA thymoma cells (18) were maintained in RPMI-1640 plus 10% heat-inactivated fetal calf serum (FCS). For B16-OVA cells (19), the medium was supplemented with hygromycin (100 μg/mL).

Immunization procedures
Dendritic cells (DC; ref. 20) were incubated for 60 minutes at 37°C with 2 μg/mL of OVA257–264 (21), TRP-2180,188 (ref. 22; Proimmune), or STEAP186–195 (23), and injected intradermally (5 × 10⁷/mouse). Esomeprazole was administered intraperitoneally (i.p.; 12.5 mg/kg; AstraZeneca).

OTI cells
Day 5 RAG−/− OTI CD8+ T blasts (20), which were cultured for the previous 24 hours at pH 5.5 to 7.4, were stimulated with phorbol myristate acetate (PMA; 60 ng/mL) and ionomycin (1 μg/mL) and assessed for intracellular cytokine production (20) or for cytolytic activity by ⁵¹Cr release assay (19). Lytic units (LU) were determined as the number of effector cells capable of killing 50% (LU50) of the target cells. Cells cultured at pH 7.4 were also CFSE-labeled (20) and injected i.v.

STEAP-specific T cells
Splenocytes from C57BL/6 mice vaccinated with STEAP186–195-pulsed DCs were specifically restimulated in vitro and, after an additional day of culture at pH 6.5 or 7.4 (day 5), were either counted by trypan blue exclusion or assessed for intracellular cytokine production (20).

Tumor implantation, processing, and flow cytometric analyses
Mice were challenged s.c. with 2 × 10⁷ B16-OVA or 5 × 10⁴ B16F1 cells. When needed, B16 melanoma–bearing mice were sublethally irradiated (600 rad) and, the day after, were infused (i.v.) with 60 × 10⁶ splenocytes derived from female donors presensitized 1 week before against tyrosinase-related protein 2 (TRP-2) antigen (21). In survival experiments, tumor size was evaluated in vivo by measuring 2 perpendicular diameters by a caliper; animals were sacrificed when lesions reached >10 mm diameter. To analyze immune cell infiltrate, animals were killed and tumors were excised, skinned and wet weighed (g) before processing. They were disaggregated and digested in collagenase D for 1 hour at 37°C to obtain single-cell suspension. Live cells were counted by trypan blue exclusion and stained for flow cytometric analysis. Data concerning the absolute numbers of different cell subsets analyzed were reported as referred to grams of tumor wet weight. Single-cell suspensions were assessed for phenotype, whereas intracellular cytokine production was evaluated after stimulation with PMA/ionomycin. For phospho-protein cytofluorometric analysis, TILs were stimulated 20 minutes with PMA (200 ng/mL), fixed, permeabilized, and stained with anti-ERK (pT202/pY204) mAb. Data were analyzed by FlowJo software (Tree Star Inc.), gating on low physical parameters that select for lymphocytes. An additional gate was generated within the CD8−CD4+ T cells, and cells were next analyzed for the expression of CFSE and/or IFNγ. The absolute cell number of each subset in the serial gates was calculated by multiplying the number of gated cells by their percentage in the total cell population. In some experiments, CD8+ cells were purified with anti-mouse CD8 MicroBeads (Miltenyi Biotec) and intracellular IFNγ production was assessed in the presence of target
cells unpulsed or pulsed with increasing concentrations of OVA257–264 peptide.

**In vivo magnetic resonance spectroscopy**

**In vivo** measurements of tumor pH were conducted by $^{31}$P magnetic resonance spectroscopy (MRS) using the exogenous cell-impermeable pH reporter 3-aminopropyl phosphonate (3-APP). The extracellular pH (pHe) value was measured from the chemical shift difference between the 3-APP resonance and that of $\alpha$-ATP at $-7.57$ ppm. B16-OVA–bearing mice underwent MRI/MRS examination once the tumor reached a volume of 300 µL. The 3-APP probe was administered i.p. (128 mg/kg) immediately before MRI/MRS analysis. Esomeprazole was administered i.p. (12.5 mg/kg), and examinations occurred at times ranging from 20 minutes to 6 hours. Data analysis and technical procedures have been previously described in detail (10).

**Statistics**

Statistical analyses were conducted using the unpaired Student $t$ test or the log-rank test (survival curves) with Prism 5 software (GraphPad Software). $P$ values less than 0.05 were considered statistically significant.

**Results**

**Acidic pH induces reversible anergy in TILs**

To test the effects of pH alterations on the proliferative and functional properties of tumor-specific T cells, we selected TILs (ref. 14; Supplementary Table S1) able to specifically recognize autologous melanoma cells (Supplementary Fig. S1). To mimic pH conditions at the tumor site, TILs were cultured at pH ranging from 7.4 to 6.5 for 3 days, as previous experiments showed that longer exposure was associated with significant apoptosis. After 3 days of culture, Annexin-V/PI staining showed no major change in the percentage of early or late apoptotic cells at pH 7.0 or 6.5 relative to pH 7.4 (Fig. 1A). TILs proliferative activity, evaluated by CFSE dye dilution following CD3/CD28 stimulation, was significantly decreased at pHs lower than physiologic levels in all cases, with a mean inhibition of 30% (with a range of 14%–54%) and 56% (with a range of 40%–90%) at pH 7.0 and 6.5, respectively (Fig. 1B). Similar data were obtained using CD3+ T cells from healthy donors or Ag-specific CD8+ T lymphocytes from patients with melanoma, suggesting this is a more general response of T cells to unfavorable pH conditions rather than a specific TIL feature (Supplementary Fig. S2).

Impaired proliferation in the absence of apoptosis at acidic pH levels suggested the onset of anergy. Indeed, IL-2 secretion in response to mitogenic stimuli, a feature typically tested for assessing T-cell anergy (24), was strongly affected in TILs maintained for 3 days at an acidic pH (Fig. 2A). Interestingly, IL-2 production (Fig. 2A), as well as proliferative activity (data not shown), were restored when TILs first cultured in pre-conditioned medium (pH 6.5), were then returned to physiologic pH (7.4) for 24 hours before the assay, suggesting that the phenomenon is reversible at least under these experimental conditions. CD25 expression, upon CD3/CD28 stimulation, was also progressively affected by culturing TILs at decreasing pH levels, and expression was similarly reversed by physiologic pH buffering for 24 hours (Fig. 2B).

Anergic T cells may have compromised JAK3/STAT5 pathway signaling (25) and reduced activation of extracellular signal–regulated kinase (ERK; ref. 26). Correspondingly, STAT5 and ERK phosphorylation in response to IL-2 or OKT3 activation, respectively, was completely abrogated in TILs following 3 days of culture at pH 6.5, whereas a full recovery was obtained by pH buffering for an additional 24 hours (Fig. 2C). A tendency to decrease CD3 and $\zeta$-chain expression, another feature of T cells in patients with cancer (27), was in addition observed upon TIL culturing at a low pH (Supplementary Fig. S3).

Perforin degranulation and IFN-$\gamma$, TNF-$\alpha$, and IL-2 release in response to autologous tumor cells were significantly impaired at pH 6.5, although these functions were restored with additional 24-hour incubation at physiologic pH (Fig. 3).
The detrimental impact of low pH on T-cell activity was not due to the effects of pH on the tumor target cells, as these cells did not change their surface expression of human leukocyte antigen (HLA) class I, tumor antigens (Supplementary Fig. S4A), or the HLA/peptide complexes recognized by the Ag-specific T cells under low pH conditions (Supplementary Fig. S4B). Taken together, these data show that acidic pH conditions resembling those observed in tumor lesions promote a reversible anergic state in human TILs, probably due to downregulation of the high affinity
IL-2 receptor (CD25) and perturbation of the STAT5 and ERK pathways.

**Effector functions of murine T lymphocytes are also affected by low pH**

To investigate the sensitivity of murine T lymphocytes to acidic pH, in vitro primed OTI cells (20) were cultured for 24 hours at pH ranging from 5.5 to 7.4 and assessed for their viability. At pH less than 6.5 most of the cells died, whereas more than 80% survival was observed at pH 6.5 (Fig. 4A). Upon antigen stimulation, cytolytic activity, as well as IFNγ and IL-2 release (Fig. 4B and C), were substantially reduced. As for human TILs (Figs. 2 and 3), the low pH-associated reduction in cytokine secretion by OTI cells was reversed by pH buffering (Fig. 4C). CD3 and TCR expression were also significantly downregulated in OTI cells exposed to pH 6.5 (Fig. 4D) and again fully recovered upon pH adjustment, corresponding with the trend observed in human TILs (Supplementary Fig. S3). Also in line with the natural data, acidity-mediated functional impairment could be reproduced as well in spleen T cells from mice sensitized against the natural tumor antigen STEAP (Supplementary Fig. S5).

**High-dose esomeprazole mediates buffering of tumor pH, improving TIL effector functions**

To investigate whether tumor pH buffering could result in improved T-cell function in vivo, we first evaluated the pH value of tumor masses by MRS in mice challenged s.c. with B16-OVA melanoma cells. Mice were then treated with a high dose of esomeprazole (12.5 mg/kg), a PPI used as a standard therapy for neutralization of gastric acid that has been shown to increase tumor pH in human melanoma xenografts (10). The impact of PPI treatment on tumor pH was then evaluated.

MRI T2-weighted images of a B16-OVA s.c. xenograft show homogeneity in the vast majority of the tumor mass and in all sections examined, with absence of hyperintense areas or hypointense areas (Fig. 5A). Under these conditions it is reasonable to assume that pH is also homogeneous. At baseline, the tumor pH value was approximately 6.5, whereas PPI therapy caused a rapid increase in the tumor pH value, reaching 7.0 within 60 minutes (Fig. 5B) and maintaining this value at least up to 6 hours (data not shown).

To evaluate the effects of pH buffering by PPI treatment, we first assessed leukocyte infiltrate using an active immunotherapy approach. Mice bearing 7-day-old B16-OVA tumors were vaccinated with DC-OVA257–264, treated at the peak of the vaccine-induced immune response (i.e., day 3 postvaccine) with either PBS or one single PPI administration, and sacrificed 24 hours later. Leukocytes infiltrating the tumor mass were predominantly macrophages (CD11b+), and to a lesser extent CD8+ and CD4+ T cells, B cells (CD19+B220+) and immature myeloid-derived cells (CD11b+Gr1+; Fig. 5C). Most CD4+ and CD8+ T cells were CD44+, confirming exposure to antigen (Fig. 5D and E). Treatment with PPI did not substantially modify the inflammatory infiltrate in this experimental setting (Fig. 5C–E). Because the limited number of endogenous CD8+ T-cells infiltrating tumor lesions did not allow any functional analysis, a second experimental approach based on adoptive immunotherapy was introduced.

To this aim, mice bearing a 12-day-old melanoma (B16-OVA) were infused with activated CFSE-labeled CD44+ OTI T cells, 24 hours later treated with PPI or PBS and sacrificed at day 14, according to the schedule depicted in Fig. 6A. As shown in Fig. 6B (left), a tendency toward a higher infiltration of CD8+ T cells was observed in tumor lesions from mice receiving PPI with respect to PBS. However, the differences between the 2 groups
were much more evident when the TIL activation state was assessed. Indeed, the number of CD44+ CD8+ IFNγ+ T cells per gram of tumor was statistically higher in PPI-treated animals (Fig. 6B, right). We then dissected the contribution of adoptively transferred versus endogenous T cells in this phenomenon by analyzing IFNγ production in CSFE− or CSFE+ CD44+ CD8+ T cells, respectively. As depicted in Fig. 6C, both the percentage and the absolute number of IFNγ+ CD44+ CD8+ TILs staining positive for CSFE and thus representing injected OTI cells, was enhanced in PPI with respect to PBS-treated mice (Supplementary Fig. S6B). Interestingly, a potentiated activation state could be also detected in CSFE− TILs from PPI-treated mice (Fig. 6E), indicating that endogenous TILs could benefit from a buffered pH milieu as well.

Because PPIs are prodrugs activated by protonation at low pH (28), their effect on resident lymphocytes should be selectively triggered by acidic environments, such as that displayed by tumor lesions (Fig. 5A). Accordingly, no sign of increase T-cell activation was detected in both CSFE− and CSFE+ CD8+CD44+ T cells subsets isolated from spleen (Fig. 6F), lung, and kidney (data not shown) from PPI-treated mice.

Finally, to evaluate in vivo effects of PPI on TIL functions also in a model of active immunotherapy, naive OTI cells were adoptively transferred in mice, subsequently challenged with B16-OVA cells and vaccinated with DC-OVA257–264 with or without PPI treatment, according to the schedule detailed in Supplementary Fig. S7A. TIL analysis conducted in mice sacrificed at day 12 showed that treatment with PPI increased the ability of CD8+ T cells, including OTI and endogenous CD8+ T cells, to recognize OVA257–264 peptide with higher affinity (at least one log difference) with respect to the same cells derived from PBS-treated mice (Supplementary Fig. S7B).

**PPI treatment increases the therapeutic potential of adoptive immunotherapy**

To investigate the therapeutic potential of the combination of PPI treatment and immunotherapy, mice bearing an 8-day B16 or B16-OVA melanoma were treated according to the schedules detailed in Fig. 7. Esomeprazole was used at 12.5 mg/kg (Fig. 7A), which in dosing experiments (ranging from 1.25–600 mg/kg) was the highest tolerated dose, whereas activated OTI cells were administered in a single treatment at numbers ranging from 1 to 6 × 10^6 cells, representing the best condition for antitumor effect (data not shown). Preliminary experiments were also conducted to identify the appropriate time schedule for PPI treatment after adoptive transfer. As reported in Supplementary Fig. S8, PPI treatment synergized with the adoptive transfer of lymphocytes when given either at the time of lymphocyte infusion or within the following 3 days. Later on the synergy diminished and was lost by day 9, likely due to the loss of adoptively transferred T cells (29). Hence, in a first set of experiments PPI was given at the time of adoptive transfer and repeated in the following days. Combined treatment with PPI and OTI cells at the doses mentioned earlier induced a statistically significant increase in animal survival, with a doubling in the overall survival rate relative to mice treated with PBS alone (Fig. 7A). Notably, equal tumor growth inhibition was achieved when PPI was administered before OTI adoptive transfer (Fig. 7B). The specificity of the treatment was confirmed by the finding that OTI cells and PPI did not impact on the survival of mice bearing B16 not expressing OVA (Fig. 7A, inset).

To test whether pH tumor buffering could improve the efficacy of nontransgenic T cells, splenocytes from mice previously sensitized against TRP-2 (19) were adoptively transferred into B16 melanoma-bearing mice (Fig. 7C).
Mice were pretreated with nonmyeloablative total body irradiation (TBI; 600 rad) to favor in vivo expansion of the adoptively transferred T cells (2). PPI administration was applied as described in the treatment schedule (Fig. 7C). Interestingly, survival was significantly prolonged in mice receiving both TRP-2–specific T cells and PPI treatment as compared with mice treated only with adoptive T-cell transfer (Fig. 7C).

Discussion
Here, we provide evidence that tumor acidity negatively regulates CD8+ tumor-specific effector T cells in both human and murine experimental settings, and that systemic administration of esomeprazole restores physiologic pH at the tumor site and promotes a more efficient antitumor T-cell activity in melanoma-bearing mice. TILs cultured at pH values most frequently found in the tumor microenvironment (10, 11) reproducibly displayed reduced effector functions, coupled with an inability to secrete IL-2, upregulate CD25, and activate STAT5/ERK signaling upon TCR activation. This is compatible with the induction of T-cell anergy (25, 26). Together with defects in perforin degranulation and cytokine secretion (30), these features are reminiscent of those detected in TILs analyzed ex vivo or in circulating CD3+ T cells of advanced disease patients (3). Thus, acidity might indeed contribute to the dysfunction of tumor T-cell immunity observed in both mice and humans (31).

Microenvironmental acidosis, a near-universal property of solid cancers, is due in part to the upregulation of glycolysis and increased glucose consumption (7). Recent evidence has shown that mutations in crucial oncogenes, such as KRAS and BRAF, trigger the activation of glucose-related genes, leading to enhanced glycolysis in turn driving the acquisition of further mutations (32). These metabolic conditions, promoting acidity and additional biochemical alterations from the very initial steps of malignant transformation (33–35), could then act as selective forces in tumor microenvironment (7, 33–35).
Our results support the hypothesis that acidity reduces T-cell performance by unbalancing the biochemical equilibrium required for physiologic activities, including proliferation, exocytosis, and secretion. In this scenario, specific cellular pathways might be more sensitive to pH variations than others. Perforin activation, for instance, is known to depend on short C-terminal peptide cleavage within acidic lysosome-like granules (36). As reported herein, microenvironmental acidity appears to perturb plasma membrane and microtubule mobility, leading to a less efficient association of different TCR components with CD8 or other coreceptors thereby contributing to T-cell anergy (4). It could be hypothesized that additional hypoxia-driven metabolic dysfunctions, causing extracellular adenosine accumulation (37), could act in synergy with acidic pH in dampening T-cell function through A2A adenosine receptor–driven cyclic AMP intracellular generation (38).

In our in vitro experiments, acidity-related T-cell anergy appeared to be reversible upon pH buffering, although longer exposure or lower pH values caused permanent damage and T-cell apoptosis. This implies that a portion of T-cell immunity might be lost at tumor sites characterized by extreme metabolic alterations. MRS analysis indicated that pH buffering could be transiently achieved in vivo by proton pump inhibition. Similar results were previously attained in human melanoma xenografts (10), where the pH modulating effect of PPI was shown to persist no longer than 24 hours. Spatial heterogeneity of pHe within the tumor has been reported in human solid tumor xenografts and is expected to occur in a metabolically complex tumor environment (7, 11). B16-OVA tumors displayed a homogeneous structure, at least at the examined time points, suggesting that pHe might as well be rather homogeneous. However, we cannot exclude spatial distribution of pHe in localized regions which were not detectable with the settings used.

By exploiting the adoptive transfer of antigen-specific T cells, we found that in vivo pH buffering by PPI was reproducibly associated with ameliorated TIL recruitment and effector functions in melanoma lesions. Notably, this effect was extended also to endogenous T cells, suggesting that PPI-mediated pH restoration might contribute to the recovery of spontaneous immunity as well. Tumor lesions from PPI-treated mice receiving OTI cells did not show a dramatic increase of T-cell infiltration, but rather a strong boost in the activation state. Nevertheless, in all survival
experiments, pH buffering was able to significantly improve the therapeutic efficacy not only of adoptive immunotherapy but also of cancer vaccines, even based on natural melanoma antigens such as TRP-2. It could be speculated that PPI in combination with immunotherapy may enable effector T cells to locally trigger lymphocytes with different specificities through antigen spreading contributing to tumor control (39). This has been suggested to occur in human setting such as in patients with melanoma, where clinical efficacy of cancer vaccines can be achieved even in the presence of apparently low frequency of TIL in tumor regressing lesions (40). The buffering of tumor acidity may be also associated with other beneficial effects. Indeed, in addition to the boost of IFNγ release, T cells infiltrating PPI-treated tumors may improve granzyme and CD40L expression as well as pathways associated with therapeutically effective T-cell responses (41–43). Furthermore, PPI might directly correct biochemical alterations known to occur at TCR or MHC/peptide complex level (44, 45), thus strengthening T-cell/target interaction and improving tumor recognition and killing. In line with this hypothesis, we report that the TCR affinity for the cognate OVA peptide is improved in T cells collected from PPI-treated lesions. We can also easily foresee that buffering tumor acidity could improve the activity of other effector cells such as natural killer (NK) or natural killer T (NKT) cells, or reduce the negative impact of
regulatory populations (T_{reg} and MDSC), as presently under investigation.

Finally, PPI might represent a rather tumor selective immunomodulator, as they do not apparently affect T cells infiltrating normal tissues lacking acidic conditions, required for the prodrug transformation into its active sulfonamide form. The specificity of PPI for the low pH milieu might explain why these drugs can be administered at very high doses without significant toxicity, as occurs in the treatment of patients with Zollinger–Ellison syndrome (28) or in the currently ongoing clinical trials testing the combination of high-dose PPI with chemotherapy in patients with cancer (www.clinicaltrials.nih.gov).

Previous reports have shown that the hypoxic and/or metabolic pathways in cancer cells may contribute to tumor immune escape by rendering the tumor microenvironment hostile for normal cells (46, 47). The present study shows that acidity per se may represent a novel mechanism of immune escape, facilitating disease progression and invasion. In this view, pharmacologic tools restoring physiologic acidity hostile for normal cells (46, 47). The present study shows that acidity per se may represent a novel mechanism of immune escape, facilitating disease progression and invasion. In this view, pharmacologic tools restoring physiologic metabolic conditions (22, 23) or in the currently ongoing clinical trials testing the combination of high-dose PPI with chemotherapy in patients with cancer (www.clinicaltrials.nih.gov).

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

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doi:10.1158/0008-5472.CAN-11-1272

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