Combined Immunotherapy against Cancer: Limited Efficacy of Transcutaneous Immunization and Low-dose Cyclophosphamide

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Abstract Transcutaneous immunization (TCI) is a novel vaccination strategy with a promising potential for combating tumors or persistent infectious diseases. However, imiquimod-based TCI, we have previously developed, shows only limited effectiveness in terms of tumor protection, partly due to suppression by regulatory T (Treg) cells. To improve the vaccination potency we combined TCI with the cytotoxic drug cyclophosphamide (Cy) that is used for the treatment of tumors and described to mediate inactivation of Treg cells at low doses. Cy only slightly reduced Treg cell numbers in a concentration dependent manner under the chosen conditions, but also enhanced DC activation. Therefore, we used Cy-TCI in a therapeutic tumor assay where E.G7 lymphomas were subcutaneously transplanted and allowed to grow until palpable before the treatments started. Interestingly, the rates of tumor protection in TCI or Cy-TCI treated groups were identical. Towards the underlying mechanisms of the failure of Cy-TCI to provide enhanced tumor protection, we observed increased numbers of monocytic and granulocytic immature myeloid cells after Cy-TCI, partly suppressing TCI-induced immune responses. Taken together, we suggest that Cy-TCI induces inhibitory mechanisms counterregulating TCI enhancing effects, therefore suppressing vaccination-induced immune responses.

Keywords Transcutaneous Immunization, Cyclophosphamide, Regulatory T Cells, Immature Myeloid Derived Cells, Dendritic Cells

1. Introduction

Therapeutic cancer vaccines have to unify several characteristics to successfully combat solid tumors. Beside the generation of adequate T cell responses against tumor cells vaccinations should also target suppressive occurrences like immunosuppressive molecules, secreted by the tumor, or regulatory cells of the immune system, recruited or activated by the tumor \cite{1-3}. In this context we have recently described a transcutaneous immunization (TCI) method, based on the synthetic Toll-like Receptor 7 agonist imiquimod \cite{1-6}, which is the active component of the commercially available Aldara crème. When applied to the skin, imiquimod induces inflammatory responses \cite{7-10} and is also effective in patients with HPV-associated or malignant skin diseases \cite{1-3}. TCI with imiquimod directly addresses the skin associated lymphoid tissue (SALT) to initiate adaptive immunity \cite{4-6}. We and others have shown that a potent and specific primary T cell response is induced upon vaccination with Aldara and a cognate cytotoxic T lymphocyte (CTL) epitope \cite{7-10}. In this situation increased numbers of bone-marrow derived CD11c\textsuperscript{+} dermal dendritic cells (DCs) migrate from the skin in the draining lymph nodes and function as the essential antigen-presenting cells (APCs) \cite{11,12}. However, CTL responses induced by TCI with imiquimod and peptide rapidly fade away, resulting in poor memory formation and only partial tumor protection, shown by us for transplanted EG.7 tumors \cite{13} and also by others in B16 melanoma models \cite{6,14}. Towards the immunological limitations of this vaccination strategy, we have demonstrated that IL-10 and regulatory T cells (Treg) contribute to the suppression of TCI-induced immune responses \cite{2,15}. Another relevant part of Treg-mediated suppression in our context may also be mediated via CTLA4 \cite{16,17}, a major suppressive pathway of Treg cells and also relevant in tumor-induced suppressive environments \cite{18}.

The cytotoxic drug cyclophosphamide (Cy) is a nitrogen mustard-alkylating agent that is used for the treatment of various diseases. Until now, the molecular mechanisms of Cy are not fully understood, but there is a correlation between dose and biological activity \cite{19}. Higher doses of Cy are applied during bone-marrow transplantations and in refractory rheumatoid diseases as immunomodulatory drug \cite{20,21}. Beyond this, also low doses of Cy are effective in
the treatment of melanoma [22,23] and autoimmune disorders [24]. On the cellular level, Cy can lead to enhanced immune responses by increasing for example the differentiation of sensitized CTL precursors into effective CTLs [25-27]. Furthermore, Cy can induce lymphodepletion, subsequently leading to expansion of immature DC in peripheral blood [28], as well as decreasing Treg numbers and also inhibit their suppressive capacities [29].

In this study we combined our TCI protocol with low-dose Cy treatment to prevent suppression by Treg cells and to induce a specific immune response and enhanced tumor protection. Low dose Cy did not diminish vaccination-induced immune responses and led to an enhanced DC activation concurrent with a reduction of Treg cells after vaccination. However, despite an enhanced DC activation and Treg cell depletion, Cy-TCI did not result in an enhanced CTL response or tumor protection compared to TCI alone. Hypothesizing that Cy induces additional inhibitory factors after combined treatment, we detected increased numbers of monocytic as well as granulocytic immature myeloid cells in the spleens of Cy-TCI treated mice along with a reduced CTL activation capacity ex vivo compared to TCI or untreated controls. Taken together, the combination of low-dose Cy with TCI does not diminish vaccination-induced immune responses. Therefore, such combination appears feasible, but we suggest that Cy also induces inhibitory mechanisms that suppress vaccination-induced immune responses and may impair the overall effectiveness of immunotherapeutic treatments.

2. Material and Methods

2.1. Mice

C57BL/6 mice at the age of 6-8 weeks were obtained from the local animal facility of the University of Mainz. DEREG mice on C57BL/6 background were kindly provided by T. Sparwasser [30]. All animal work performed in this study was conducted to the national guidelines and was reviewed and confirmed by an institutional review board/ethics committee headed by the local animal welfare officer. The responsible national authority finally approved the animal experiments, which is the National Investigation Office Rheinland-Pfalz (Koblenz, Germany). The Approval ID assigned by this authority: AZ 23 177-07/G08-1-023 and G13-1-012.

2.2. Transcutaneous Immunizations

Transcutaneous immunizations with Imiquimod were described previously [10]. Briefly, dorsal hair was removed using electric clippers. 100 µg SIINFEKL-Peptid (OVA257-264, kindly provided by Dr. Stevanovic, Department Immunology, Institute for Cell Biology, University of Tübingen, Germany) was added to 50 mg crème containing 5 % Imiquimod (Aldara, Meda) and applied on the dorsal skin of anesthetized mice on days 0 and 1.

2.3. Cyclophosphamide Treatment

Mice were injected i.p. with 25 mg/kg, 50 mg/kg or 100 mg/kg cyclophosphamide one day prior to immunizations.

2.4. Depletion of Regulatory T Cells

Where indicated C57BL/6 mice were depleted of regulatory T cells by i.p. injection of an anti-CD25 Ab (PC61, 400 µg, day -4). For regulatory T cell depletion in DEREG mice received i.p. injections of diphtheria toxin (1 µg) at days -1, 1 and 3.

2.5. Tumor Rejection Assay

Tumors were implanted by s.c. injection of 4 x 10^5 EG.7 thymoma cells (from ATCC) as described previously [13]. Treatments started when tumors get palpable (10-12 days after inoculation). Tumor size was monitored three times per week with a caliper measuring the diameter in two dimensions. Mice were killed when tumor size exceeded 20 mm in both diameters.

2.6. Flow Cytometric Analyses and in vivo Cytotoxicity Assay

The following mAbs were used for analyses by flow cytometry: -conjugated CD8 (clone 53-6.7), -conjugated CD44 (clone IM7), -conjugated CD62L (clone MEL-14), -conjugated CD4 (clone), -conjugated CD25 (clone 7D4), APC-conjugated FoxP3 (clone) all from BioLegend or eBiosciences. Blood samples were collected after tail vein incision and incubated on ice with specific mAbs as indicated after a hypotonic lysis step. For detection of SIINFEKL-specific T cells samples were stained with H2-K^b tetramer. H2-K^b—SIINFEKL^c cells were determined among gated CD8^+ T cells. In vivo cytolytic activity was determined by transfer of carboxyfluorescein diacetate succinimidyl ester (0.4 µM CFSE^low and 4 µM CFSE^high) labelled splenocytes (2 x 10^7 per mouse). The CFSE^low population was additionally loaded with SIINFEKL-peptide. Splenocytes of immunized and control mice were analysed by flow cytometry. All analyses were performed with a LSRII Flow Cytometer and FACSDiva software (BD Pharmingen, Hamburg, Germany).

2.7. Preparation and Staining of Lymph Nodes

C57BL/6 mice were treated as indicated. Inguinal, axillar and brachial lymph nodes were collected, picked and digested for 40 minutes with type 2 collagenase (1 mg/ml from Worthington Biochemicals, Lakewood, NJ) and DNase I (100 µg/ml, Sigma-Aldrich). The reaction was stopped by adding 10 mM EDTA (Sigma-Aldrich). The cells were
labelled with the following mAbs: APC-Cy7-conjugated CD11c (clone N418), Pacific Blue-conjugated MHC II (clone M5/114.15.2), -conjugated CD8 (clone 53-6.7), -conjugated CD4 (clone RM4-5), -conjugated CD80 (clone 16-10A1), -conjugated CD86 (clone GL-1) all from BioLegend or eBiosciences. Absolute cell numbers were calculated using Flow-Count Fluorospheres from Beckmann-Coulter according to the manufacturer’s instructions.

2.8. In Vitro Proliferation Assay

C57BL/6 mice were treated as indicated over a period of three weeks. Spleens of experimental groups were pooled and single-cell suspensions were prepared. After a hypotonic lysis step cells were divided to three parts: i) CD25+ cells were depleted using CD25-specific magnetic beads and MACS sorting, further working with the flow through, ii) Ly6G+ cells were depleted using Ly6G-specific magnetic beads and MACS sorting, further working with the flow through and iii) cells without any depletion (magnetic beads from Miltenyi Biotech, Bergisch-Gladbach, Germany). After incubation of cells with SIINFEKL (10 ng/ml) for 1 hour, cells were washed twice. Stimulator (3 x 10^5 cells/well) and responder OT-1 splenocytes (1 x 10^5 cells/well) were cocultured for three days, pulsed with ^3^H-thymidine (0.5 µCi/ml; Perkin Elmer, Rodgau, Germany) and harvested the following day. ^3^H-thymidine incorporation was assessed with a 1205 betaplate reader (LKB Wallac, Turku, Finland).

2.9. Statistical Analysis

All statistical analyses were performed using GraphPad Prism (version 5.0a for Mac OS X, GraphPad Software, San Diego California USA, www.graphpad.com). Survival analyses were performed by the Mantel-Cox test. For comparison between two groups a two-tailed Student’s t test was used. Comparisons of multiple groups were performed by one-way ANOVA with Bonferroni’s posttest. For all analyses, p < 0.05 was considered as statistically significant.

3. Results

3.1. Low-dose Cyclophosphamide does not Impair Transcutaneous Immunization with Imiquimod

We have previously shown that TCI with imiquimod elicits a potent primary CTL-response, but is diminished by the presence of Treg cells as well as IL-10 [2]. The prevention of Treg-mediated suppression either by reducing Treg-cell numbers or by inhibiting Treg-cell function might be a promising approach to enhance TCI-induced immune responses. Cy treatment is described to impede Treg-mediated suppression [29] and might therefore represent an interesting way to enhance TCI efficacy. To this end, we investigated the effects of various Cy amounts on the TCI-induced CTL response using the TLR7 ligand imiquimod together with the major histocompatibility complex class I-restricted T-cell epitope OVA_{257,264}. One week after vaccination we determined the in vivo cytolytic activity of the induced CTLs. As depicted in Figure 1A, systemic administration of higher doses (50 mg/kg and 100 mg/kg) of Cy slightly decreased lysis of peptide-loaded target cells, whereas the lowest amount of Cy (25 mg/kg) had no impact on cytotoxicity compared to TCI alone. Additionally we assessed the frequency of peptide-specific CD8 T cells (Figure 1B). Again, higher doses (50 mg/kg and 100 mg/kg) reduced induction of peptide-specific CD8 T cell responses compared to TCI alone. Nevertheless, the lowest dose of Cy (25 mg/kg) had no impact on CTL response indicated by equivalent amounts of peptide-specific T cells compared to TCI alone. Analysing the activation status of induced CD8+ T cells (Figure 1C) revealed no difference of CD44 and CD62L expression in both groups, leading to the conclusion that the additional application of low-dose Cy (25 mg/kg) does not diminish TCI induced priming of CTL-responses.

As the combination of low-dose Cy and TCI seemed to be feasible, we evaluated this vaccination protocol in terms of tumor protection (Figure 1D). We inoculated E.G7 thymoma cells, expressing chicken ovalbumin as a surrogate tumor antigen. After tumors became palpable (days 10-12), the mice were treated with TCI with or without injection of Cy (25 mg/kg). Cy treatment alone already enhanced survival of tumor-bearing mice with a median survival of 49 days compared to the untreated group (median survival 32.5 days), which was not significant compared to the untreated group (p=0.15 by Mantel Cox test). TCI alone and Cy-TCI showed identical results (median survival not reached), indicating that Cy neither reduces nor enhances survival compared to TCI alone, also in line with the former experiments concerning induction of CTL-responses.
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Figure 1. Low-dose cyclophosphamide does neither impair TCI-induced CTL-responses nor enhance survival in a therapeutic tumor model.

C57BL/6 mice were injected i.p. with various amounts of Cy as indicated (day -1). Transcutaneous immunizations were performed using Imiquimod-containing Aldara 5 % creme (50 mg) together with OVA257-264 (SIINFEKL, 100 µg) at days 0 and 1. A) Cytolytic activity was determined after adoptive transfer of peptide-loaded splenocytes. B) The frequency and C) activation phenotype of peptide-specific CD8+ T cells in the blood was assessed at day 6 using surface staining and flow cytometry. D) E.G7 cells (4 x 10^6) were injected subcutaneously into the flank of mice. Immunization treatments and Cy injection (25 mg/kg) started after 10-12 days when tumours became palpable. A-C) A cumulative analysis (mean with SD) of two independent experiments for 50 mg/kg Cy or 100 mg/kg Cy (each n=6) and five independent experiments with n=22 for 25 mg/kg Cy, n=21 for TCI and n=14 for control is shown. D) A cumulative analysis of two independent experiments with n=13 for Cy-TCI, TCI and Cy alone and n=12 for untreated is shown. (*) Significant difference (p<0.05) by Mann-Whitney test, n.s not significant.

Figure 2. Low-dose cyclophosphamide does not completely deplete Tregs and has no major impact on their functionality.

Mice were injected i.p. with Cy as indicated (day -1). At days 0 and 1 transcutaneous immunizations were performed. A) Groups received titrated amounts of (Cy 25 mg/kg, 50 mg/kg, 100 mg/kg). Amounts of Treg cells in the spleens were analysed. B) Mice were treated with Cy (25 mg/kg) and TCI or TCI alone with or without the additional injection of an anti-CD25 Ab (clone PC61, 400 µg i.p., day -4). C) FoxP3-diphteria toxin receptor transgenic (DEREG) mice were immunized with Cy (25 mg/kg) and TCI or TCI alone with or without depletion of Treg cells by application of diphtheria toxin (DT, 1 µg i.p., days -1, 1, 3). B,C) In vivo cytolytic activity was assessed after adoptive transfer of peptide loaded and carboxyfluorescin diacetate succinimidyl ester-labelled target cells. A cumulative analysis of two or three independent experiments is shown. (*) Significant difference (p<0.05) by Student’s t test.

3.2. Low-dose Cy Depletes Treg Cells after TCI

One major reason for combining Cy and TCI was its ability to inhibit the suppressive capacity of Treg cells or even deplete them [29]. As the treatment with low-dose Cy in our vaccination protocol did not increase the induction of a specific immune response, we repeated the titration of Cy, immunized mice with TCI and stained for Treg cells in the spleens at day 6 when analysing the CTL-response. The amount of Treg cells (determined as CD4+CD25+FoxP3+ cells) was slightly decreased after the application of 25 mg/kg or 50 mg/kg compared to the untreated control. This effect was more pronounced after treatment with 100 mg/kg Cy. In contrast, TCI alone led to a marginal non-significant increase in Treg cell numbers (Figure 2A).
To determine the remaining suppressive activity of Treg cells after Cy-TCI, we additionally used an anti-CD25 monoclonal antibody (clone PC61, 400 µg, day -4) to deplete Treg cells before immunizations started. Additional depletion of CD25+ cells led to an enhanced immune response shown by increased lysis of peptide-loaded target cells by cytotoxic T cells, independent of additional low-dose Cy application (Figure 2B). Since antibody-mediated depletion may exert secondary effects, we also employed another approach for Treg depletion by immunizing FoxP3-diphtheria toxin receptor transgenic mice (DEREG) with Cy-TCI or TCI alone. By administration of diphtheria toxin (DT) FoxP3+ cells are specifically ablated. Comparable to our published results [2] and in line with the former experiments using a monoclonal antibody, the absence of Treg cells promoted an increased CTL response compared to TCI without prior depletion. The additional application of Cy again had no impact on the detected immune response (Figure 2C). Taken together, these results implicate that either low-dose Cy administration has no sufficient impact on Treg cell numbers and functionality or it induces other suppressive factors leading to CTL responses comparable to TCI alone.

3.3. Enhanced DC activation Phenotype after Cy-TCI

For TCI-induced CTL responses, bone-marrow derived CD11c+ dermal DC are essential for priming of peptide-specific T cells [12]. As Cy is described to mediate expansion of immature DC as a result of lymphodepletion [28], we next investigated the influence of systemic administration of Cy on the local activation of DCs in the skin. We treated mice with TCI in the presence or absence of Cy in the additional presence of FITC on the shaved backs to identify skin-derived DCs in the draining lymph nodes. Three days after immunization the draining lymph nodes were harvested and analysed by flow cytometry (gating strategy Figure 3A). Gating on FITC+DCs we were unable to detect differences in the number of DCs after Cy-TCI compared to TCI alone (data not shown). Interestingly, the phenotype of migrated skin-derived DCs revealed a significantly increased expression of the activation markers CD80 (Figure 3B) and CD86 (Figure 3C) of Cy-TCI-treated mice compared to TCI alone or to FITC treated, but otherwise unimmunized controls. These data clearly show that Cy leads to an enhanced activation status of skin-derived DC.

![Figure 3](image-url)

**Figure 3.** Increased activation phenotype of skin-derived DC after cyclophosphamide application.

Mice were injected i.p. with Cy as indicated (day -1). At days 0 and 1 transcutaneous immunizations were performed. Additionally a FITC-solution was applied on the shaved back skin. Skin-draining lymph nodes were harvested at day 3, digested and analysed by flow cytometry. **A)** Skin-derived DC were determined as living cells with a typical FSC/SSC distribution as MHC class II+CD11c+ and among these as FITC+. Mean fluorescence intensity of **B)** CD80 and **C)** CD86 were analysed. Depicted are representative results with n=4 for 2 independent experiments. (*) Significant difference (p<0.05) by Students t test.
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Figure 4. Monocytic as well as granulocytic immature myeloid cells are induced upon low-dose cyclophosphamide.

Mice were injected i.p. with Cy as indicated (day -1). At days 0 and 1 transcutaneous immunizations were performed. At day 6 samples from A,B) blood (relative amount gated on CD11b+) and C,D) spleen (absolute cell counts) were stained for monocytic (CD11b+Ly-6G-Ly6Chi) and granulocytic (CD11b+Ly-6G+Ly-6Clo) immature myeloid cells and analysed by flow cytometry. Depicted are cumulative results of two independent experiments with n=8 for Cy and TCI, TCI and Cy and n=4 for control. (*) p<0.05 by one-way ANOVA with Bonferroni’s posttest. n.s. = no significant difference. E) Mice were treated with Cy-TCI or TCI alone over a period of three weeks. Afterwards spleens of immunized mice were depleted of Ly6G/C+ or CD25+ cells or left undepleted, loaded with SIINFEKL peptide and used as APCs for OT-1 T cells. Proliferation of T cells was assessed in a [3H] thymidine assay (in triplicate wells). (*) Significant difference (p<0.05) by Students t test.
3.4. Cy-induced Immature Myeloid-derived Cells Suppress CTL Response Post TCI

So far, we demonstrated that the CTL response induced by Cy-TCI was not diminished compared with TCI alone. However, the depletion of Treg cells as well as the DC activation phenotype after Cy suggest that Cy should lead to an enhancement of the CTL response which was not the case. Therefore, we hypothesized that there might also be inhibitory factors induced by Cy treatment. Recently, Sevko and co-workers [31] demonstrated the induction of myeloid-derived suppressor cells (MDSCs) upon systemic injection of low-dose Cy. To elucidate whether such immature myeloid-derived cells were induced upon Cy-TCI, we treated mice with low-dose Cy (25 mg/kg) and TCI, TCI alone or Cy alone and stained for monocytic (CD11b-Ly-6G-Ly6Ch) and granulocytic (CD11b-Ly-6G-Ly-6Clo) immature myeloid-derived cells [32] in blood and spleen, five days after treatment. As depicted in Figure 4A, no induction of monocytic immature myeloid-derived cells could be observed in the blood after Cy treatment, whereas some granulocytic immature myeloid-derived cells (Figure 4B) were detectable. Conversely, the absolute cell numbers in the spleens of treated mice revealed that TCI alone showed no induction of the monocytic subtype (Figure 4C), but a significant induction of granulocytic cells (p<0.05 by one-way ANOVA with Bonferroni’s posttest; Figure 4D). Cy alone showed a comparable amount of monocytic as well as granulocytic cells compared to TCI alone. These results indicate that Cy induces suppressive immature myeloid cells.

To further evaluate whether the induced monocytic and granulocytic immature myeloid-derived cells contribute to the suppression of TCI-induced CTL responses, we immunized mice over a period of three weeks in the presence or absence of Cy. On day 5 after the last treatment, we took the spleens of mice and either depleted Ly6G/C+ cells (targeting immature myeloid cells), CD25+ cells (targeting Treg cells) or left the splenocytes undepleted. Afterwards we co-cultured these splenocytes with TCR-transgenic T cells from OT-1 mice, recognizing the H2-Kb-restricted epitope OVA257-264 to subsequently analyse the induced T-cell proliferation by [%H] thymidine incorporation assay. As depicted in Figure 4E, we detected a strong proliferation of OT-1 T cells using splenocytes from TCI-treated mice as APCs. Notably, this was independent of the cell-type depleted before co-culture. Undepleted splenocytes from Cy-TCI treated induced OT-1 T-cell proliferation in a comparable way to TCI alone. Interestingly, the depletion of either Ly6G/C+ or CD25+ cells from Cy-TCI treated donor cells, significantly increased T-cell proliferation further underlining the idea that Cy induces inhibitory factors that suppress TCI-induced immune responses.

All together, these data demonstrate that Cy treatment leads to the induction of monocytic and granulocytic immature myeloid-derived cells, that an addition to regulatory T cells contribute to the suppression of induced CTL response

4. Discussion

Transcutaneous immunization by the topical application of a TLR agonist together with a CTL epitope might be an attractive way to deliver antigens for the treatment of malignancies or persistent virus infections. Directly targeting skin-resident APCs, TCI elicits a potent primary immune response [10]. As demonstrated by us previously, TCI-induced specific T cells rapidly fade away resulting in poor memory formation and insufficient protection in therapeutic tumor models [6,13,14]. The strength of the T-cell response can be enhanced by additional treatments like CD40 ligation or UV-B irradiation leading to memory formation [12,13] and providing improved vaccination efficacy. Analysing the underlying mechanisms of TCI, we have revealed a suppressive effect of Treg cells, diminishing the induced CTL response [2]. Therefore, a promising advancement could be the combination of TCI with the inhibition of Treg cells. In this context, cyclophosphamide, a chemotherapeutic agent which is used for the treatment of melanoma, lymphomas and solid tumors, could be of interest [22,23,33]. Concerning its mode of action, Cy is described to decrease Treg cell numbers as well as to inhibit their suppressive capacity [29].

Combining low-dose Cy with TCI induced immune responses comparable to TCI alone, whereas higher doses suppressed the vaccination-induced CTL responses, most likely by its well-known lymphotoxic effects.

Working with low-dose Cy therefore seemed to be feasible. Towards the impact of Cy administered systemically on skin-resident DC targeted by TCI, we found that numbers of cells migrated from the skin to the draining lymph nodes are not affected (data not shown). Characterizing the phenotype of migrated DC revealed a more activated pattern, shown in upregulation of CD80 as well as CD86 after Cy-TCI compared to TCI alone. As bone-marrow derived CD11c+ dermal DCs are crucial in skin-associated immunity [12,34,35], an enhanced activated phenotype of skin-derived DC should support the induction of CTL responses which was not the case. Therefore, we suggest that beside Cy mediated enhancing effects, namely Treg cell depletion and DC activation, also inhibiting factors are induced, counteracting the stimulatory effects.

Recently Sevko and coworkers analysed low-dose Cy in a murine melanoma-model. Although this treatment results in reduced Treg cell levels and immunogenic tumor cell death upon Cy treatment, this treatment fails to increase mouse survival [31]. They show that the ineffective anti-tumor immunity is due to the production of various inflammatory mediators, such as GM-CSF, IL-1β, IL-5, IL-10 IFNγ and TNF-α. Importantly, this leads to the expansion and activation of immunoregulatory myeloid-derived suppressor cells (MDSC) that accumulate especially in melanoma lesions. Transferring these findings to our setting, the
induction of immunosuppressive immature myeloid-derived cells could counter-regulate the beneficial effects of Cy and TCI. Indeed, we found increased numbers of monocytic as well as granulocytic immature myeloid cells in the spleens of Cy-TCI treated mice. In addition, utilizing splenocytes from those mice as APCs in co-culture experiments after depleting either CD25⁺ or Ly6G/C⁺ cells revealed a strongly increased proliferation of responder T cells, leading to the conclusion that both cell types suppress CTL responses upon Cy treatment.

5. Conclusions

Taken together, we show that Cy does not diminish TCI-induced CTL-responses. Besides the disadvantage of immature myeloid cell formation and other suppressive influences, low-dose Cy mediates Treg cell reduction and DC activation, both desirable factors for vaccination. Therefore future vaccination protocols on the basis of Cy application may incorporate means to inhibit or modify immunosuppressive factors, e.g. accumulation of immature myeloid cells, to develop new therapeutic options against tumors by combining chemotherapy with immunomodulatory treatments.

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REFERENCES


