The Immune System Strikes Back: Cellular Immune Responses against Indoleamine 2,3-dioxygenase

Rikke Baek Sørensen¹, Linda Berge-Hansen¹, Niels Junker¹, Christina Aaen Hansen¹, Sine Reker Hadrup¹, Ton N. M. Schumacher², Inge Marie Svane¹, Jürgen C. Becker³, Per thor Straten¹, Mads Hald Andersen¹*

¹ Center for Cancer Immune Therapy (CCIT), Department of Hematology, Herlev University Hospital, Herlev, Denmark, ² Division of Immunology, The Netherlands Cancer Institute, Amsterdam, The Netherlands, ³ Department of Dermatology, University of Würzburg, Würzburg, Germany

Abstract

**Background:** The enzyme indoleamine 2,3-dioxygenase (IDO) exerts an well established immunosuppressive function in cancer. IDO is expressed within the tumor itself as well as in antigen-presenting cells in tumor-draining lymph nodes, where it promotes the establishment of peripheral immune tolerance to tumor antigens. In the present study, we tested the notion whether IDO itself may be subject to immune responses.

**Methods and Findings:** The presence of naturally occurring IDO-specific CD8 T cells in cancer patients was determined by MHC/peptide staining and ELISpot. Antigen specific cytotoxic T lymphocytes (CTL) from the peripheral blood of cancer patients were cloned and expanded. The functional capacity of the established CTL clones was examined by chromium release assays. The study unveiled spontaneous cytotoxic T-cell reactivity against IDO in peripheral blood as well as in the tumor microenvironment of different cancer patients. We demonstrate that these IDO reactive T cells are indeed peptide specific, cytotoxic effector cells. Hence, IDO reactive T cells are able to recognize and kill tumor cells including directly isolated AML blasts as well as IDO-expressing dendritic cells, i.e. one of the major immune suppressive cell populations.

**Conclusion:** IDO may serve as an important and widely applicable target for anti-cancer immunotherapeutic strategies. Furthermore, as emerging evidence suggests that IDO constitutes a significant counter-regulatory mechanism induced by pro-inflammatory signals, IDO-based immunotherapy holds the promise to boost anti-cancer immunotherapy in general.

Introduction

Recent advances in the understanding of the interplay between cancer cells and cells of the immune system demonstrated the capacity of the immune system to recognize and destroy neoplastic cells; nevertheless, despite the fact that neoplastic transformation is associated with the expression of immunogenic antigens, the immune system often fails to respond effectively to these antigens. Thus, the immune system becomes tolerant towards these antigens [1]. There is a general consensus, that this acquired state of tolerance must be overcome for cancer immunotherapy to succeed. The biological role of Indoleamine 2,3-dioxygenase (IDO) in the immune system is still a subject of active investigation. For example, endogenous IDO has been implicated as one mechanism by which maternal tolerance toward the fetus is maintained [2]. Likewise, IDO can mediate suppression of T-cell immunity to MHC-mismatched liver allografts [3]. Furthermore, IDO control T-cell responses to autoimmune disorders, and regulates the severity of a variety of experimental autoimmune disorders [4,5]. Thus, several lines of evidence indicate that IDO is a major component to maintain the homeostasis of the immune system which, however, also contributes to tumor-induced tolerance. The expression and activation of IDO creates a tolerogenic milieu in the tumor and the tumor-draining lymph nodes (LN) either via direct suppression of T cells by degradation of the essential amino acid tryptophan or via enhancement of local regulatory T-cell (Treg)-mediated immunosuppression. With respect to the former, some of the biological effects of IDO are mediated through local depletion of tryptophan, whereas others are mediated via immunomodulatory tryptophan metabolites [6,7].

IDO can be expressed within the tumor by tumor cells as well as tumor stromal cells, where it inhibits the effector phase of immune responses. In this setting, IDO is believed to inhibit the effector phase of the immune response [8,9]. In a murine model it was observed that tumor cells transfected with IDO become resistant to immune eradication, even in mice in which a fully protective immune response had been established by immunization [9]. Most importantly, in the clinical situation it was repeatedly observed, that expression of IDO in tumor cells is associated with an impaired prognosis [10,11].

Additionally, IDO-expressing antigen-presenting cells (APC) are present in tumor-draining LN, where they are believed to create a
tolerogenic microenvironment. Indeed, IDO-expressing CD19+ plasmacytoid dendritic cells (DC) isolated from tumor-draining LN mediate profound immune suppression and T-cell anergy in vivo [12-14]. Plasmacytoid DC from normal LN and spleen do not express IDO. It should be noted that very few cells constitutively express IDO in normal lymphoid tissue except in the gut. This implies that the DC in tumor-draining LN, which constitutively express IDO must receive a stimulus which is related to the presence of the tumor. This stimulus is believed to be delivered by activated Tregs migrating from the tumor to the draining LN. Tregs have been shown to induce IDO via cell-surface expression of CTLA-4 [15]. The induction of IDO converts the tumor-draining LN from an immunizing into a tolerizing milieu. Indeed, when IDO+ DC are injected in vivo, they create suppression and anergy in antigen-specific T cells in the LN draining the injection site [14,16]. Hence, IDO is a critical cellular factor contributing to immune suppressive and tolerogenic mechanism in cancer. Consequently, IDO has become a very attractive target for the design of new anticancer drugs and several IDO inhibitors are currently being investigated [17,18]. However, to date the possibility to harness the immune system to target IDO-expressing cells has not been explored. This is particular surprising since IDO-expressing cells antagonizes the desired effects of other immunotherapeutic approaches and a combination of IDO- and tumor-targeting immunotherapies should be highly synergistic. In the present study, we provide evidence for the immunogenicity of IDO by demonstrating the presence of spontaneous cytotoxic immune responses against IDO-expressing cells in cancer patients.

Methods

Patients

Peripheral Blood Mononuclear Cells (PBMC) were collected from cancer patients (renal cell carcinoma, melanoma, and breast cancer) and healthy controls. Blood samples were drawn a minimum of four weeks after termination of any kind of anti-cancer therapy. The majority of renal cell carcinoma patients had previously been treated with IL2 and IFN-α, most melanoma patients had received high dose IL2 and IFN-α, while all breast cancer patients were pre-treated with several kinds of chemotherapy, (e.g. epirubicin, docetaxel, cabecitabine), trastuzumab, and/or endocrine therapy. PBMC were isolated using Lymphoprep separation, HLA-typed (Department of Clinical Immunology, University Hospital, Copenhagen, Denmark) and frozen in FCS with 10% DMSO. The protocol was approved by the Scientific Ethics Committee for The Capital Region of Denmark and conducted in accordance with the provisions of the Declaration of Helsinki. Written informed consent from the patients was obtained before study entry.

Peptides

Epitopes from IDO were predicted using the “Database SYFFEITH” [19] in combination with manual examination of the protein sequence for MHC class I anchor residues. Eleven synthetic 9mer and 10mer peptides were produced: IDO1 (IDO54-62: QLKRERVEKL), IDO2 (IDO164-172: FLVSLIVELE), IDO3 (IDO195-203: TLLKALLE), IDO4 (IDO41-49: FIAKHLPDIL), IDO5 (IDO199-207: ALEIEASCL), IDO6 (IDO260-268: VLSKGDGL), IDO7 (IDO283-291: DLKMNFLKT), IDO8 (IDO275-283: VLLIQGQTA), IDO9 (IDO101-109: KVLPRNIAV), IDO10 (IDO104-112: KLNMDSDL), and IDO11 (IDO341-349: SLSRSHHLQIF). The peptides were dissolved in DMSO (final concentration 10 mM) or distilled water (final concentration 2 mM). The HLA-A2 high affinity binding epitope HIV-1 pol476-484 (ILKEPVHG) was used as irrelevant control. The HLA-A2 restricted Epstein - Barr virus peptide EBVEMLF1250-1258 (GLCTHLVAML) was used as control.

Assembly assay for peptide binding to MHC class I molecules

The binding affinity of the synthetic peptides (Genscript) to HLA-A2 molecules, metabolically labelled with [35S]-methionine, was measured in the assembly assay, as described previously [20,21]. The assay is based on peptide-mediated stabilization of empty HLA-molecules released upon cell lysis, from the TAP-deficient cell line T2. Stably folded HLA-molecules were immune-precipitated by using the HLA class I specific, conformal dependent monoclonal antibody (mAb) W6/32 and separated by isoelectric focusing (IEF) gel electrophoresis. Major histocompatibility complex (MHC) heavy-chain bands were quantified using the ImageGauge Phosphoimager program (FUJI Photo Film, Carrolton, TX). The intensity of the band is directly related to the amount of peptide-bound class I MHC complex recovered during the assay. Subsequently, the extent of stabilization of HLA-A2 is directly related to the binding affinity of the added peptide. The recovery of HLA-A2 was measured in presence of 100, 10, 1, 0.1 and 0.01 μM of the relevant peptide. The C50 value was calculated for each peptide as the peptide concentration required for half maximal stabilization of HLA-A2.

ELISPOT assay

The ELISPOT assay was used to quantify peptide epitope-specific IFN-γ releasing effector cells as described previously [22]. In some experiments PBMC were stimulated once in vitro with peptide prior to analysis as described [23] to extend the sensitivity of the assay. After 7 days in culture with peptide and 40 U/ml IL-2 (PeproTech), cells were counted and analyzed in IFN-γ ELISPOT. Briefly, nitrocellulose bottomed 96-well plates (MultiScreen MAIP N45; Millipore) were coated overnight with IFN-γ capture mAb (Mabtech). The wells were washed, blocked by X-vivo medium and the effector cells were added in duplicates at different cell concentrations, with or without 10 μM peptide. The plates were incubated overnight. The following day, medium was discarded and the wells were washed prior to addition of biotinylated secondary Ab (Mabtech). The plates were incubated at room temperature (RT) for 2 hours, washed, and Avidin-enzyme conjugate (AP-Avidin; Calbiochem/Invitrogen Life Technologies) was added to each well. Plates were incubated at RT for 1 hour and the enzyme substrate NBT/BCIP (Invitrogen Life Technologies) was added to each well and incubated at RT for 5–10 min. Upon the emergence of dark purple spots, the reaction was terminated by washing with tap water. The spots were counted using the ImmunoSpot Series 2.0 Analyzer (CTL Analyzers).

Flow cytometry

For tetramer stainings, PBMC from cancer patients and healthy donors as well as tumor infiltrating lymphocytes (TIL) from cancer patients were stimulated once in vitro with peptide, or analysed directly ex vivo. The CD8 T cells were isolated from PBMC using the Dynal CD8 negative isolation kit (Dynal Biotech) at day 7. The resulting T-cell cultures were stained with PE coupled tetramers, followed by antibody staining with an appropriate combination of the fluorochrome-coupled mAbs: CD8allophycocyanin/APC-Cy7, CD3-FTTC, CD45RO-FTTC, CD45RA-PE-Cy5 and CD28-allophycocyanin (BD Bioscience). For comparison, cells were stained with appropriate isotype controls. Tetramers were prepared using the MHC-peptide exchange technology [24] as described in [25]. Tetramer stainings were performed in PBS +2%
FCS, for 15 min., RT, in the dark, whereas antibody stainings were performed in PBS +2% FCS, for 20 min., 4°C in the dark. The MHC tetramer complexes used were: HLA-A2/IDO5 (ALLEIASCL) and HLA-A2/HIV-1 p0447-454s (ILKEPVHGV). The samples were analyzed on BD FACS Aria, using DIVA software (BD Biosciences).

Cancer cells were examined for HLA-A2 expression using flow cytometry. Cells were stained with a fluorochrome-coupled HLA-A2 mAb (BD Biosciences). For comparison, cells were stained with an isotype matched control. The samples were analyzed on BD FACS Aria, using DIVA software (BD Biosciences). Assuming normality, HLA-A2 expression was given by a one-tailed two sampled t-test comparing MFIHALA-A2 and MFIIsotype control, where MFI is the Mean Fluorescence Intensity. For p-values <0.05 (significance level) cells were defined HLA-A2+. The fold of expression was defined as MFIHALA-A2/MFIIsotype control.

Intracellular protein staining
PBMC, DC, and cancer cells were examined for intracellular IDO expression using flow cytometry. After fixation and permeabilization (Cytotox/Cytoperm, BD), cells were stained with anti-IDO antibody (Millipore Corporation). For comparison, cells were stained with an isotype matched control (Millipore Corporation). After incubation, cells were washed twice with Perm/Washbuffer (BD) before staining with a FITC-labeled secondary antibody (DAKO). The samples were analyzed on BD FACS Aria, using DIVA software (BD Biosciences). Assuming normality, intracellular IDO expression was given by a one-tailed two sampled t-test comparing MFIIDO and MFIIsotype control, where MFI is the Mean Fluorescence Intensity. For p-values <0.05 (significance level) cells were defined IDO+. The fold of expression was defined as MFIIDO/MFIIsotype control.

Dendritic cells (DC)
DC were generated from PBMC by adherence on culture dishes at 37°C for 60 min in RPMI-1640 enriched with 10% human AB serum. Adherent monocytes were cultured in RPMI-1640 supplemented with 10% human AB serum in the presence of IL-4 (1000 U/ml) and GM-CSF (250 U/ml) for 6 days. DC were matured by addition of IL-1β (1000 U/ml), IL-6 (1000 U/ml), TNF-α (1000 U/ml), and PGE2 (1 μg/ml).

Establishment of antigen specific T-cell cultures and clones
PBMC from cancer patients were stimulated with irradiated (25 Gy), IDO5-loaded autologous DC (PBMC:DC ratio = 5×10^5:3×10^3), with 5 μg/ml anti-IFN, 20 U/ml IL-12 (PeproTech), and 40 U/ml IL-7 (PeproTech). The cultures were stimulated every 10 days with irradiated autologous DC (2×) followed by irradiated PBMC (2×), with 40 U/ml IL-2 (PeproTech). After one month, growing cultures were tested for specificity for IDO5 in a standard 51Cr-release assay. PBMC from specific cultures were cloned by limiting dilution in the presence of 10^5/ml irradiated (25 Gy) IDO5-loaded PBMC, and 120 U/ml IL-2. Every 3-4 days 50 μl fresh media were added containing IL-2 to a final concentration of 120 U/ml. Growing clones were expanded using IDO5-loaded PBMC (5×10^4 cells/well) and 120 U/ml IL-2. After expansion the clones were tested for specificity and cytotoxic potential in a standard 51Cr-release assay.

Cytotoxicity assay
Conventional 51Cr-release assays for CTL-mediated cytotoxicity was carried out as described elsewhere [26]. Target cells were T2-cells, in vitro generated autologous immature DC (iDC) and matured DC (mDC), in vitro generated allogeneic HLA-A2+iDC and mDC, autologous ex vivo isolated monocytes, T cells and B cells (isolated using CD14+, CD3+ or CD19+ microbeads (MACs)), the natural killer target cell line K562, ex vivo enriched HLA-A2+AML blasts (B and T cells were depleted from the bone marrow of AML patients using CD19+ and CD3+ microbeads (MACs)), the breast cancer cell lines CAMA-1 and MDA-MB-231, as well as the colon cancer cell lines HCT-116 and SW480 (all available at the American Type Culture Collection (ATCC)), and FM55M [27]. Lysis were blocked using the HLA specific mAb W6/32 (2 μg/100 μl) [28]. In some assays, target cells were treated with 100 U/ml IFN-γ for 2 days.

Down-regulation of IDO in cancer cells
Human SW480 cancer cells were transfected with indicated short hairpin RNA (shRNA) plasmids obtained from SuperArray using FuGene6 (Roche) according to manufacturers instructions. Cells were lysed directly in LSB buffer (Sigma). The LSB lysates were boiled for 5 min. and loaded on 10% precast protein gels (BioRad). Proteins were electro transferred to a PVDF membrane (Millipore Corporation) by a semidry transfer method and probed with indicated antibodies according to manufacturers instructions. Blots were developed with the ECL system obtained from Amersham and a CCD camera (LAS-1000, Fujifilm). Following antibodies were used: anti-Cdk7 (MO-1) (Santa Cruz) and anti-IDO (Millipore Corporation).

Statistical analysis
Statistical analysis was performed on data on intracellular IDO staining. Assuming normality, intracellular IDO expression was given by a one-tailed two sampled t-test comparing MFIIDO and MFIIsotype control, where MFI is the Mean Fluorescence Intensity. For p-values <0.05 (significance level) cells were defined IDO+. The fold of expression was defined as MFIIDO/MFIIsotype control.

Results
IDO-derived HLA-A2-restricted T-cell epitopes
Eleven IDO-derived peptides were selected from the main HLA-A2 specific anchor residues and subsequently synthesized [29]. Using the ELISPOT IFN-γ secretion assay, we then examined PBMC from cancer patients and healthy individuals for the presence of specific T-cell responses against these IDO-derived peptides. This approach has previously proved to be highly effective for identifying tumor specific cytotoxic T-lymphocytes (CTL) in cancer patients [22,30,31]. Thus, PBMC from HLA-A2+, late stage cancer patients (breast cancer, melanoma and renal cell carcinoma) were stimulated once with the different peptides in vitro before examination by ELISPOT. This procedure was chosen to extend the sensitivity of the ELISPOT as described [22,32]. ELISPOT responses were detected against IDO2 (IDO164-172; FLVSLLVEI) (Fig. 1b), IDO6 (IDO320-325; VLSKGDGL) (Fig. 1c) and, especially, IDO5 (IDO193-207; ALLEIASCL) (Fig. 1a). To this end, PBMC from a larger number of cancer patients were examined for reactivity against IDO2 and IDO6, since these responses were less frequent compared to IDO5. As control, we examined PBMC from healthy individuals for reactivity against these three IDO derived peptides. No spontaneous responses could be detected against any of the IDO-derived peptides in any of the healthy controls. A BLAST search of the amino acid sequences of these
peptides using the “NCBI database” showed that these motifs are only prevalent in the IDO protein.

Detection of IDO-reactive HLA-A2-restricted T cells in cancer patients

The apparently most immunogenic IDO-derived peptide, i.e. IDO5, was examined for its binding affinity to HLA-A2 by comparison with a HLA-A2 high affinity positive control epitope, i.e. HIV-1 pol476-484 (ILKEPVHG), by the assembly assay (Table 1). For comparison, the HLA-A2 binding affinity of IDO2 and IDO6 were analyzed as well. C50 values were estimated for each peptide as the peptide concentration required for half maximal stabilization of HLA-A2. Notably, IDO5 bound HLA-A2 even better than the high-affinity control epitope, whereas IDO2 and IDO6 bound HLA-A2 with intermediate affinity compared to the control epitope (Table 1). The high binding affinity of IDO5 to HLA-A2 enabled us to make stable HLA-A2/IDO5 tetramers, which were used to detect IDO-reactive CTL by flow cytometry. This analysis clearly confirmed the presence of IDO5-reactive CD8 T cells in the blood of HLA-A2+ cancer patients (Fig. 2). Figure 2a illustrates an example of an IDO5-specific T cell response after in vitro stimulation in a renal cell carcinoma patient with an HIV-1 tetramer-complex used as control. While the frequency of IDO-reactive T cells is markedly increased by in vitro stimulation, IDO-reactive T cells were readily detectable ex vivo in selected patients (Fig. 2b): In the three patients with strongest responses after in vitro stimulation, a respective reactivity was also detected ex vivo. Overall, PBMC from 7 HLA-A2+ healthy individuals and 11 HLA-A2+ patients were analyzed. No IDO-reactive T cells could be detected in any of the healthy donors (Fig. 2b). The ex vivo stainings of IDO-reactive T cells showed that naturally occurring IDO5-specific T cells have a CD54RA-CD28+ central/effector memory phenotype [33]. An example of such an ex vivo phenotype staining of IDO5 tetramer gated cells is shown in figure 2c. As a comparison the sample were stained with isotype matched controls. Next, we examined the presence of IDO5-specific T cells in IL-2 treated TIL cultures from HLA-A2+ melanoma and head and neck cancer patients by tetramer stainings. As illustrated in figure 2d IDO5-specific T cells could readily be detected among the TIL. Overall, 4 of the 5 analyzed patients had detectable IDO5-specific T cells. The percentage of IDO5-specific T cells among the TIL ranged from 0.05 to 0.1% of the total number of CD8 T cells. Likewise, IDO5-specific T cells in TIL cultures from melanoma and head and neck cancer patients could be detected in ELISPOT (data not shown). To control the specificity of the HLA-A2/IDO5 tetramer we stained an IDO5-specific T-cell clone. The HLA-A2/IDO5 tetramer did efficiently stain the IDO5-specific T-cell clone, whereas the T-cell

Table 1. HLA-A2 binding affinity of selected IDO-derived peptides.

| Peptide      | Sequence   | C50 (mmol/L) |
|--------------|------------|--------------|
| HIV-1 pol 476-484 | ILKEPVHG   | 0.3          |
| IDO2 (IDO164-172) | FLVSLLVEI  | 30.0         |
| IDO5 (IDO199-207) | ALLEIASCL  | 0.2          |
| IDO6 (IDO320-326) | VLSKGDGL   | 3.5          |

*The C5 value is the peptide concentration required for half maximal stabilization of HLA-A2.

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clone was not stained by the control HLA-A2/HIV-1 pol476–484 tetramer (Fig. 2e).

Having identified patients hosting responses against the IDO5 peptide, we used PBMC from such patients to generate CTL bulk cultures against this peptide in vitro. PBMC were stimulated with autologous IDO5-pulsed DC. After four rounds of stimulation, the peptide specificity was tested in standard 51Cr-release assays. Bulk cultures were generated from three different cancer patients. Cells from two of these bulk cultures lysed TAP-deficient T2-cells pulsed with IDO5 peptide. To analyze the lytic capacity of IDO-specific T cells in more detail, CTL clones were established from one of these bulk cultures by limiting dilution cloning. After a short expansion period, the specificity of the growing clones was analyzed in standard 51Cr-release assays. Of thirty three T-cell clones displaying an IDO-specific lytic capacity, four clones were selected for further expansion due to a superior growth rate. A representative T-cell clone is depicted in figure 3a: the T-cell clone RBS35 effectively killed IDO5-pulsed T2-cells whereas T2-cells pulsed with an irrelevant peptide (HIV-1 pol476–484 (ILKEPVHG)) were not lysed (Fig. 3a).

**Killing of tumor targets by IDO-specific T cells**

Importantly, RBS35 killed not only peptide pulsed T2-cells but also the HLA-A2+/IDO+ colon cancer cell line SW480 with high efficacy (Fig. 3b). In contrast, RBS35 did not lysed the HLA-A2+/IDO+ colon cancer cell line HCT-116 (Fig. 3b). HLA-restriction of
RBS35 was confirmed by blocking HLA-class I using the HLA-specific mAb W6/32, which completely abolished lysis of the SW480 target cells (Fig. 3b). A number of cancer cell lines were examined for IDO expression by intracellular protein staining followed by FACS analysis [34]. The colon cancer cell line SW480, the melanoma cell line FM55M, the breast cancer cell lines CAMA-1 and MDA-MB231 as well as directly enriched AML-blasts were all IDO+. Only the colon cancer cell line HCT-116 was IDO-. Representative examples of IDO stainings are illustrated in histograms in figure 3c.

Next, we demonstrated that the HLA-A2+/IDO+ melanoma cell line FM55M was killed by RBS35 (Fig. 4a). Cold targeted inhibition assays using unlabeled T2-cells pulsed with the IDO5 (10 μM) peptide confirmed the HLA-A2/peptide-specificity of the killing: The addition of cold (unlabeled) IDO5-pulsed T2-cells completely abrogated the killing of FM55M melanoma cells, whereas the addition of cold T2-cells pulsed with an irrelevant peptide (HIV-1 pol476-484) did not have an effect on the killing of FM55M (Fig. 4a). No cytotoxicity was observed against the NK-cell target cell line K562 (Fig. 4a).

Furthermore, we tested the ability of RBS35 to lyse human AML-blasts enriched directly ex vivo from the bone-marrow of AML patients. For this purpose, we depleted T cells (CD3+) and B cells (CD19+) from the bone marrow of AML patients; the highly enriched AML-blasts (CD3-, CD19-) were subsequently used as target cells in a 51Cr-release assay. We enriched AML blasts from six patients (5 HLA-A2+ patients and 1 HLA-A2- patient). All AML blasts expressed IDO (data not shown). RBS35 efficiently lysed the HLA-A2+ leukemia cells in an HLA-dependent manner, while HLA-A2- leukemia cells were not lysed (Fig. 4b). We did not find a correlation between the amount of IDO expression in the AML blasts and the percentage of lysis (data not shown).

To illustrate the representative killing of tumor targets by RBS35 the killing of SW480 by a polyclonal, IDO5-specific bulk culture as well as by three other T-cell clones (RBS26, RBS31, RBS46) are shown in figure 4c and figure 4d. The T cell clones had distinct TCR as verified by TCR Clonotype Mapping (data not shown). Similar to RBS35, none of the clones (RBS26, RBS31, RBS46) lysed the HLA-A2+/IDO+ colon cancer cell line HCT-116 (Fig. 4d).

Finally, we examined the killing of the HLA-A2+ breast cancer cell lines CAMA-1 and MDA-MB-231. The CAMA-1 cell line was killed by RBS35, whereas MDA-MB-231 was not recognized by RBS35 (Fig. 5a). IFN-γ treatment increased the expression of IDO...
and HLA-A2 in both cell lines. This is exemplified for CAMA-1 in figure 5b and c. In agreement with this, IFN-γ treatment increased the killing by RBS35 of CAMA-1 and introduced killing of the MDA-MB-231 cells (Fig. 5a).

Additionally, using IDO ShRNA we down-regulated IDO protein expression in the human SW480 colon cancer cell line and thereby rescue these tumor cells from being killed by the polyclonal IDO-specific bulk culture, whereas cells transfected with irrelevant control ShRNA were killed (Fig. 5d). This down-regulation was visualized by intracellular protein stainings. These stainings confirmed that the use of IDO ShRNA reduced the level of IDO protein expression in the cells (Fig. 5e).

**Figure 4. Specificity and functional capacity of IDO5-specific T cells assayed by 51Cr-release assays.**

- **(a)** Lysis by the IDO5-specific T-cell clone (RBS35) of the HLA-A2+/IDO+ melanoma cell line FM55M without and with the addition of cold T2-cells pulsed with IDO5 peptide or an irrelevant peptide (HIV-1 pol476-484) (inhibitor to target ratio = 20:1), and NK cell activity of RBS35 examined using the natural killer cell line K562 as target cells. **(b)** Lysis by RBS35 of AML-blasts enriched from 5 HLA-A2+ AML patients and 1 HLA-A2- AML patient. B cells and T cells were depleted from the bone marrow of the AML patients using CD19+ and CD3+ microbeads, respectively. The highly enriched AML-blasts were used as target cells with or without the addition of the HLA-class I specific antibody W6/32. **(c)** Lysis of T2-cells pulsed with IDO5 peptide or an irrelevant peptide (HIV-1 pol476-484), and lysis of the HLA-A2+/IDO+ colon cancer cell line SW480 by an IDO5-specific T-cell bulk culture. **(d)** Lysis of the HLA-A2+/IDO+ colon cancer cell line SW480 and HLA-A2+/IDO+ colon cancer cell line HCT-116 by three different IDO5-specific T-cell clones (RBS26 (white triangle), RBS31 (black triangle), RBS46 (grey triangle)) assayed by 51Cr-release assay. All assays were performed in different E:T ratios. doi:10.1371/journal.pone.0006910.g004

**Killing of dendritic cells by IDO-specific T cells**

IDO expression is not restricted to tumor and tumor stroma cells, but can also be induced in immune cells. Thus, as the next and even more important step we addressed the question whether IDO-expressing DC would also be susceptible to killing by IDO-reactive CTL. To test this notion, we generated autologous DC from the same donors from whom the CTL clones had been generated; the DC were matured by addition of a standard maturation cocktail consisting of IL-1β, IL-6, TNF-α, and PGE2 [35]. RBS35 effectively killed the in vitro matured DC (mDC). In contrast, autologous IDO immature DC (iDC) were not killed by RBS35 (Fig. 6a). Moreover, we examined the recognition of allogenic IDO mDC and IDO iDC from an HLA-A2+ background by RBS35 and RBS46 (black triangle) and RBS46 (grey triangle) (Fig. 6b). RBS46 killed both mDC and iDC when the T cells were stimulated with IL-2. This killing was not observed when the T cells were not stimulated with IL-2 (Fig. 6c).
donor. The allogenic mDC were killed by RBS35 whereas the allogenic IDO-iDC were not killed (Fig. 6a). In figure 6b it is illustrated that mDC express IDO in contrast to iDC. Next, we tested the ability of RBS35 to lyse autologous monocytes, T cells and B cells. For this purpose, we isolated CD14+ monocytes, CD3+ T cells and CD19+B cells directly ex vivo from IDO+ PBMC. The isolated cells were subsequently used as target cells in a 51Cr-release assay. Autologous CD14+ monocytes, CD3+ T cells and CD19+B cells were not lysed by RBS35 (Fig. 6c). IFN-γ treatment introduced killing of the CD14+ monocytes, but not of the T and B cells (Fig. 6c).

Finally, we set up an in vitro model to examine if IDO specific T cells enhance immune responses by depleting IDO-expressing suppressive cells. Hence, cultures of PBMC, with and without IDO specific T cells, were treated with IFN-γ to increase the immune activity as well as IDO expression in the cultures. Five days later we examined the immune reactivity against the HLA-A2 restricted immunodominant epitope from EBV BMLF1280-288 (GLCTLVAML) in the cultures. Although the overall cell number was the same in the cultures the reactivity against the EBV peptide was higher in the cultures with IDO5-specific T cells (Fig. 6d). Next, we scrutinized if the addition of IDO specific T cells increased the immune reactivity to an extent that allowed detection of EBV responses in an ELISPOT with only 10^4 PMBC; far below the normal detection limit. Indeed, we could detect a clear EBV response even at this low concentration of PBMC (Fig. 6d). As expected we could not detect any EBV response at this low cell concentration in the culture without IDO5 specific T cells (Fig. 6d).

**Discussion**

IDO has a critical immunosuppressive function in cancer. In the present study, we set out to examine if IDO itself may serve as target for immune responses, which may be exploited for immune therapy. By following a 'reverse immunology' approach, we identified HLA-A2 peptides within the IDO protein to which
Figure 6. Functional capacity of an IDO5-specific T-cell clone (RBS35) to kill immune cells assayed by $^{51}$Cr-release assays. (a), Left: Lysis of autologous in vitro immatured and matured DC. Right: Lysis of allogeneic HLA-A2$^*$ in vitro immatured and matured DC. All assays were performed in different E:T ratios. (b), Histograms showing intracellular IDO expression in DC. Data are representative of 3 experiments. Intracellular IDO expression was given by a one-tailed two sampled t-test comparing MFIIDO (dark histograms) and MFIIsotype control (light histograms), where MFI is the Mean Fluorescence Intensity. Left: In vitro immatured DC ($p = 0.100$). Right: In vitro matured DC ($p = 0.001$). (c), Lysis of autologous CD14$^+$ monocytes, CD3$^+$ T cells and CD19$^+$ B cells isolated directly ex vivo from IDO$^+$ PBMC, and lysis of autologous CD14$^+$ monocytes, CD3$^+$ T cells and CD19$^+$ B cells after IFN-$\gamma$ treatment. As a control, we used in vitro generated autologous IDO$^+$ immatured DC and IDO$^+$ matured DC. (d), Examples of HLA-A2 restricted T-cell responses against EBV BMLF1280-288 (GLCTLVAML) as measured by ELISPOT in PBMC from a breast cancer patient. Cultures of PBMC were treated with IFN-$\gamma$ for 5 days with autologous freshly isolated CD8 T-cells (left) or with autologous IDOS specific T-cells (at a PBMC:IDOS specific T-cell ratio of 300:1) (right) before examination for reactivity against the HLA-A2 restricted epitope from EBV BMLF1280-288 (GLCTLVAML). Three different PBMC concentrations were examined; $1.5 \times 10^5$ cells, $5 \times 10^5$ cells (two top rows) and $10^4$ cells (bottom two rows). doi:10.1371/journal.pone.0006910.g006
spontaneous T-cell reactivity were detected in patients suffering from unrelated tumor types, i.e. melanoma, renal cell carcinoma and breast cancer but not in healthy individuals. These naturally occurring T-cell responses in cancer patients could be readily visualized by flow cytometry using HLA/peptide tetramers after in \textit{vivo} stimulation but even more remarkable in direct \textit{ex vivo} assays. In this regard, it should be noted that with a few exceptions it is not possible to detect conventional tumor associated antigen specific T cells in PBMC from cancer patients directly \textit{ex vivo}, i.e. without any \textit{in vitro} steps to expand or enrich these cells [32]. Thus, in some cancer patients IDO-specific T cells are present in relative high frequencies. We further reveal that IDO-specific T cells readily can be detected in the tumor microenvironment in lesions from both head and neck cancers and melanoma using tetramer stainings as well as ELISPOT. The combined detection of IDO-specific T cells in blood and tumor lesions indicate that these cells are capable of circulating and homing to the effector site. This is a significant finding, since several clinical reports have indicated the existence of a functional dissociation between local and systemic anti-cancer T-cell responses [36,37].

Furthermore, we confirm that IDO-reactive T cells are indeed peptide specific, cytotoxic effector cells. Hence, an important issue concerning translation of our findings to a clinical setting lies in demonstrating the cytotoxic capacity of IDO-specific T cells against clinically relevant target cells. In this regard, IDO-specific T cells effectively lysed IDO$^+$ cancer cell lines of different origin, such as melanoma, colon carcinoma, and breast cancer. Most importantly, leukaemia cells enriched directly from AML patients were killed \textit{ex vivo} by IDO-specific T cells. IDO expression in blasts of AML patients have been correlated to significantly shorten overall and relapse-free survival [38]. The presence of spontaneous T-cell responses against IDO-derived peptide epitopes in PBMC from patients suffering from unrelated cancer types as well as the killing of cancer cells of different origin by IDO-specific T cells underline the immunotherapeutic potential of IDO. The induction of IDO expression after \textit{in vitro} maturation of DC might be a major problem and an explanation for the lack of success of DC-based immunotherapy. However, even more distinctive was our finding that IDO-specific CTL recognize and kill IDO$^+$ \textit{in vitro} matured DC; hence, IDO-specific T cells are indeed able to kill immune suppressive cells. It is well described that IDO is up regulated in DC in tumor draining LN creating a tolerogenic microenvironment. Furthermore, DC isolated from cancer patients has impaired functionality and possesses an altered phenotype compared to healthy individuals. Hence, IDO vaccination might restore the ability of DC in cancer patients to initiate and/or activate anti-cancer immune responses by killing immune competent DC.

Counter-regulatory responses are important in the immune system as they help to limit the intensity and extent of immune responses, which otherwise could cause dangerous damage to the host. However, with regard to anti-cancer immunotherapy counter-regulatory responses antagonize the ability to create an intense immune response against the tumor. Counter-regulation differs from tolerance in the sense that counter-regulation is a secondary event, elicited only in response to immune activation. By definition most anti-cancer immunotherapeutic strategies irrespective of their molecular targets aim at the induction of an immunological activation and inflammation. Virtually, within the limits of acceptable toxicity as much immune activation as possible is the goal; hence, counter-regulation is not desired. IDO may be highly relevant to the outcome of immunotherapy of cancer as an inflammation-induced counter-regulatory mechanism. Hence, IDO is known to be induced by both type I and II interferons, which are found at sites of immune activation and inflammation [39,40]. In this regard, we demonstrated that that the susceptibility of tumor cells to killing by IDO-reactive T cells is increased by preincubation with IFN-\(\gamma\) although this increased recognition might also be due to higher expression of HLA on the surface after IFN-\(\gamma\) treatment. Furthermore, we included a model where the addition of IDO specific T cells to IFN-\(\gamma\) treated PBMC increased the immune reactivity towards EBV. Hence, our findings suggest that IDO-based immunotherapy would work synergistically with additional therapy that introduces inflammation at the site of the tumor. Additionally, we illustrate that down-regulation of IDO rescue tumor cells from being killed by IDO-specific T cells. In this regard, immunoselection of antigen-loss variants during immunotherapy have been demonstrated in several cases [36,41–44]. Accordingly, down regulation of IDO during IDO-based immunotherapy might save cancer cells and immune suppressive cells for immune-mediated destruction by IDO-specific CTL. This should, however, lead to removal of local immune suppression within the tumor and/or tumor draining LN and thereby enabling circulating effector cells to function or to get activated.

The state of lymphopenia following high-dose chemotherapy appears to provide a window of enhanced responsiveness to immunotherapy [45]. Additionally, certain chemotherapeutic drugs may assist in breaking immune tolerance by preferentially inhibition of Tregs [46,47]. Nonetheless, chemotherapy alone is never sufficient to break tolerance to tumors probably because the original tolerogenic mechanisms rapidly restore tolerance following each cycle of chemotherapy. In principle, therefore, chemotherapy might provide an environment in which IDO-based immunotherapy could have a synergistic effect on established tumors, by preventing the re-acquisition of tolerance. This is supported by the report that the IDO blocker 1-Methyl-Tryptopan (1-MT) work synergistically with different chemotherapeutic drugs in established murine cancers [17].

In conclusion, we demonstrate that IDO is a natural target for CTL in cancer patients. Thus, we describe that although IDO up regulation has an immune suppressive effect in cancer, the immune system – at least in some patients - finds a way to fight back; by the induction of IDO-specific CTL. The immune dysfunction in the draining LN due to local IDO expression in DC is not indicative of a generalized immune suppression in cancer patients. Subsequently, it may well be possible to strengthen the described natural immunity against IDO by active immunotherapy. Since IDO-expressing cells might antagonize the desired effects of other immunotherapeutic approaches targeting IDO-expressing cells by vaccination would consequently be highly synergistic with such therapeutic measures. Finally, the recent ability of genetically modifying lymphocytes has opened possibilities for the \textit{in vitro} creation of specific lymphocytes with appropriate therapeutic properties. Hence, the IDO T-cell receptors isolated from the T-cell clones described in the current manuscript can be introduced into a patient’s normal lymphocytes and administered to lymphodepleted patients as described for other antigens [43,48–50].

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Author Contributions

Conceived and designed the experiments: MHA. Performed the experiments: RBS LBH NJ CAH. Analyzed the data: JCB PtS MHA. Contributed reagents/materials/analysis tools: SRH TNS IMS MHA. Wrote the paper: MHA. Edited manuscript: TNS JCB PtS.
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