Induction of monocyte antitumor response by human cancer cells transduced with TNF-GFP fusion gene: possible implications for immunotherapy of cancer

Jerzy Więckiewicz, Bożenna Mytar, Rafał Szatanek, Kazimierz Węglarczyk, Jarosław Baran

Department of Clinical Immunology, Polish-American Institute of Pediatrics, Jagiellonian University Medical College, Krakow, Poland

Abstract: This study was undertaken to determine how human pancreatic cancer (HPC-4) cells transduced with the TNF-GFP fusion gene (TLG) alter the antitumor response of human monocytes in vitro and whether they could act as an antitumor vaccine. In our model, HPC-4 cells were transduced with retroviral vector harboring TLG gene and designated as HPC-4TLG. The TLG protein expression was confirmed by Western blot and flow cytometry analysis. Monocytes were co-cultured with transduced and control HPC-4 cells. The secretion of TNF, IL-10 and IL-12 was measured by ELISA. The cytotoxicity of monocytes against HPC-4 cells was determined by MTT test. The results show that the HPC-4TLG cells expressed membrane-bound, intracellular and secretory TLG protein. When cultured with HPC-4TLG cells, monocytes released a higher amount of TNF, but IL-10 and IL-12 secretion was inhibited. The pre-exposure of monocytes to HPC-4TLG, but not to HPC-4, cells did not decrease TNF nor increase IL-10 production, thus not leading to monocyte deactivation. Also, the antitumor cytotoxicity of monocytes stimulated with HPC-4TLG was not downregulated, which occurred when non-transduced HPC-4 cells were used. In conclusion, compared to parental HPC-4 cells, TLG gene transduced HPC-4 cells induced stronger antitumor response of monocytes in vitro and prevented deactivation of monocytes. (Folia Histochemica et Cytobiologica 2011, Vol. 49, No. 3, 512–520)

Key words: fusion gene, monocytes, tumor cells, tumor necrosis factor

Introduction

A serious problem with the retroviral system of gene transfer is that it is difficult to maintain a high level of cloned gene expression following transduction. Usually, several passages of transduced cells result in the outgrowth of clones that express much lower levels of the transgene coded protein. The use of the internal ribosome entry site (IRES) elements rather than internal promoters could increase the likelihood of stable expression of the cloned gene of interest. However, even the use of IRES sequence does not completely reduce the number of potential mechanisms, both at the transcriptional and post-transcriptional levels, that could affect the transgene expression [1]. Cloned gene expression also depends on retroviral DNA integration site in the host genome [2, 3]. Cloning of the gene coding for green fluorescent protein (GFP) isolated from the jellyfish Aequorea victoria and its efficient species-independent expression is a new versatile reporter gene system [4, 5].

Autologous cancer cells used as vaccines to enhance antitumor activity have been explored extensively, but their efficiency has proved to be low. There-
fore, genetically engineered tumor cells transfected with various cytokines coding genes have been used [6–8]. However, despite positive effects in murine experimental tumor models, their clinical efficiency in human cancer is marginal, with an objective response rate of 3.3% [9]. Also, human monocytes that exert a direct cytotoxic effect on tumor cells, as mediated by radical oxygen intermediates (ROI) and tumor necrosis factor (TNF), upon arrival at the tumor site form tumor infiltrating macrophages (TIM), which become polarized to M2 macrophages, thus strongly reducing their cytotoxic activity [10]. This occurs due to the tumor cell products, including extracellular matrix components, IL-10, CSF-1 and other cytokines. In general, M2 macrophages have poor antigen presenting capacity, have an IL-12low and IL-10high phenotype, suppress the inflammatory response and Th1 adaptive immunity, and promote angiogenesis, tissue remodeling and tumor progression [10]. Such TIM changes may be responsible, at least in part, for the inefficiency of tumor vaccines.

We have previously observed that human monocytes stimulated with tumor cells produce TNF, IL-10 and IL-12 [11]. However, when monocytes were first pre-exposed to tumor cells, sorted out from co-culture and then re-stimulated with tumor cells, they became M2 polarized cells that produced increased amounts of immunosuppressive IL-10 [11]. Therefore, we wanted to investigate whether the use of TNF gene transduced tumor cells might increase their immunostimulatory potential for monocytes and prevent their M2 polarization.

**Material and methods**

**Reagents and antibodies.** The oligonucleotides were purchased from TIB Molbiol (Poznan, Poland). Polymerase chain reaction (PCR) and enhanced avian reverse transcriptase (RT) kits, agarose (for routine use), polybrene (hexadimethrine bromide), and the antibiotics ampicillin, neomycin sulfate and tylsomin tartrate were from Sigma Chemical Company (St. Louis, MO, USA). The Pwo DNA polymerase with the appropriate incubation buffer was obtained from Roche (Basel, Switzerland). Restriction enzymes, 1 Kb Plus DNA ladder and T4 DNA ligase were from Invitrogen (Carlsbad, CA, USA). The jetPEI cationic polymer transfection reagent was obtained from Obigene (Morgan, Irvine, CA, USA), DMEM and RPMI 1640 media, respectively, supplemented with the following antibiotics: 50 μg/ml neomycin, 100 μg/ml ampicillin, 10 μg/ml tylsomin, and 10% FBS referred to further as complete. Both cell lines were incubated at 37°C in a humidified 5% CO₂ atmosphere. All cell cultures were routinely tested for *Mycoplasma sp.* contamination using a Mycoplasma PCR ELISA kit (Roche Diagnostics GmbH, Penzberg, Germany).

**Cell cultures.** The RetroPack PT67 murine dualtropic cell line was used for retrovirus packaging and production of retroviral pMIGR1 and recombinant pMIGR1-TLG (pTLG) vectors. The established human pancreatic adenocarcinoma (HPC-4) cell line [13] was used for transduction of the TLG fused gene by the recombinant retroviral infection. The PT67 cells and HPC-4 cells were grown in DMEM and RPMI 1640 media, respectively, supplemented with the following antibiotics: 50 μg/ml neomycin, 100 μg/ml ampicillin, 10 μg/ml tylsomin, and 10% FBS referred to further as complete. Both cell lines were incubated at 37°C in a humidified 5% CO₂ atmosphere. All cell cultures were routinely tested for *Mycoplasma sp.* contamination using a Mycoplasma PCR ELISA kit (Roche Diagnostics GmbH, Penzberg, Germany).

**Construction of fusion TLG transgene and retroviral vector preparation.** The cDNA encoding human TNF (26 kDa) was obtained by the RT-PCR method using specific oligonucleotide primers and the total RNA sample isolated from human blood monocytes stimulated in *vitro* for 2 h with LPS (1 μg/ml). The synthesis of the first strand CDNA was carried out with enhanced avian RT and anchored oligo-dT primers according to the manufacturer’s instructions. The PCR amplification of the first strand of TNF-cDNA was performed using the Pwo DNA polymerase and the following primers: BglFtnf sense 5'-GGCGAGATCTATGACCTGAAAGGGTATGAT-3' which provided BglII site (bolded) and XhoRtnf antisense 5'-CATATATCTCGAGCAGGGAATGATCCCAA-3' which provided XhoI site (bolded). The PCR amplification of the first strand of TNF-cDNA was performed using the Pwo DNA polymerase and the following primers: BglFtnf sense 5'-GGCGAGATCTATGACCTGAAAGGGTATGAT-3' which provided BglII site (bolded) and XhoRtnf antisense 5'-CATATATCTCGAGCAGGGAATGATCCCAA-3' which provided XhoI site (bolded). The TNF-cDNA was cleaved with BglII and XhoI restriction enzymes and ligated to the BglIII-XhoI linearized vector pMIGR1 to give the recombinant pMIGR1-TNF plasmid. The LIN23-cDNA sequence coding for a linker of (GGC)₄TCT as the template and the primers: LinSXho sense 5'-ATAATATCTCGAGGCGGCGG- GC-3' which provided XhoI site (bolded) and LinRNeo an-
Production of retroviral particles. The pTLG plasmid transfer was performed using the jetPEI transfection reagent according to the manufacturer’s protocol. Transfected PT67 cells expressing the TLG protein were isolated by sorting with the use of FACS Vantage with DiVa option (BD Biosciences Immunocytometry Systems, San Jose, CA, USA) equipped with an aerosol protection system (Flexoduct International, Greve, Denmark) using DiVa software. The ion laser Innova Enterprise II (Coherent, Santa Clara, CA, USA) operating at 488 nm was used as a light source. Sorted PT67 cells were propagated in complete DMEM medium until cell culture reached confluence and sorted again. Double-sorted PT67 cells producing retrovirus were plated at 60–80% confluency in the desired number of culture vessels. The supernatants were harvested at 24 h intervals until cells were no longer viable. The supernatants were collected, spun at 500 × g for 10 min and stored at –70°C until use.

Transduction of human HPC-4 cells. The target HPC-4 cells were cultured in complete RPMI 1640 medium for 24 h at 1–2 × 10^6 per 60-mm Petri dishes (Becton Dickinson Labware, Franklin Lakes, NJ, USA). Then the medium was replaced with 2.5 ml of thawed supernatant from transfected PT67 cells diluted 1:1 with fresh complete RPMI 1640 medium supplemented with polybrene (final concentration 5 μg/ml). The retroviral infection of the HPC-4 cells was performed three times sequentially at 12 h intervals. Then the HPC-4 cells were cultured for 48 h and sorted twice sequentially on the basis of the TLG protein fluorescence as described above. The purity of the sorted TLG positive cells (HPC-4_{TLG}) was checked by flow cytometry re-analysis. As the control, HPC-4 cells transduced with the original pMIGR1 retroviral vector and designated as HPC-4_{GFP} were used.

Western blot analysis of TLG fusion protein expression. HPC-4 cells and their genetically modified clones, HPC-4_{GFP} and HPC-4_{TLG} cells were lysed in M-PER protein extraction reagent (Pierce Biotechnology, Rochford, IL, USA) containing protease and phosphatase inhibitors (Sigma). The extracted proteins (30 μg) were loaded on 4% loading gel, separated in 12% SDS gel and transferred to the poly(vinylidene fluoride) membrane (Immune-blot PVDF, 2 μm, BioRad, Hercules, CA, USA). GFP was detected with anti-GFP mAb IgG₁ (Molecular Probes) and with horseradish peroxidase-conjugated goat anti-mouse IgG (Santa Cruz Biotech, Santa Cruz, CA, USA), as a secondary antibody. The membranes were developed with the SuperSignal West Pico Chemiluminescent Substrate (Pierce), dried and subsequently exposed to HyperFilm (Amersham Life Science, Little Chalfont, UK).

Intracellular detection of GFP and TLG fusion protein. Intracellular staining for the detection of GFP and TLG fusion protein was performed after cell fixation and permeabilization, as previously described [14]. Briefly, HPC-4 cells and their genetically modified clones were fixed and permeabiled with Cytofix/Cytoperm solution (BD Pharmingen) for 20 min at 4°C. Then, the cells were washed twice in Perm/Wash solution (BD Pharmingen) and pelleted cells were stained (30 min at 4°C) for intracellular detection of GFP and TLG fusion protein using Alexa Fluor 594-conjugated anti-GFP rabbit IgG and PE-conjugated mAb against human TNF. Appropriate matched isotype controls were used in parallel.

Flow cytometry analysis. Samples were analyzed in a FACS Canto flow cytometer (BD Biosciences) using DiVa software. The list mode data of 50,000 events from HPC-4 cells in a ‘live gate’ mode was acquired. The cells were gated according to FSC and SSC parameters. Intracellular GFP and TLG proteins were determined by green fluorescence measurement in channel FL1. Intracellular and membrane-bound TLG fusion proteins were also detected according to FL4 (Alexa Fluor 594-conjugated anti-GFP rabbit IgG) and FL2 (PE-conjugated mAb against human TNF) emission.

Isolation of cell populations. Human monocytes were separated from peripheral blood mononuclear cells of healthy donors by counter-flow elutriation, using a JE-6B elutriation system (Beckman Instruments, Palo Alto, CA, USA), as previously described [15]. They were 90–98% pure, as judged by staining with anti-CD14 mAb. Monocytes were suspended in RPMI 1640 medium with 5% FBS at the concentration of 1 × 10^5 cells/ml for in vitro cultures. In some experiments, monocytes isolated from a co-culture with parental or transfected HPC-4 cells (pre-exposure) at the ratio 1:0.3 for 4 h, were used. For this purpose, after staining with PE-labeled anti-CD14 (BD Biosciences) mAb, CD14⁺ cells were isolated by FACS sorting (FACS Vantage with DiVa option, Becton Dickinson Immunocytometry Systems). The purity of the sorted cells was checked by flow cytometry, and exceeded 95%. As control, CD14⁺ cells were sorted out from monocytes cultured in medium alone (‘dummy sorting’).

Determination of cytokine release. HPC-4, HPC-4_{GFP} and HPC-4_{TLG} cells (1 × 10^5/well) were incubated in RPMI 1640 medium with 5% FBS for 18 h in flat-bottom wells microtiter plates (Nunc, Roskilde, Denmark). Then, the supernatants were harvested and the concentration of TNF was...
measured by ELISA. Also, monocytes (1 × 10^5/well) were incubated either in the medium alone or with parental HPC-4, or HPC-4_GFP or HPC-4_TLG (3 × 10^5/well) for 18 h (as above) and the level of TNF, IL-10 and IL-12 was determined. In some experiments, isolated CD14^+ cells were cultured in the medium alone or with parental HPC-4 or their genetically modified clones (as above), and the level of cytokines was measured. All the cytokine level measurements were performed in duplicates using commercial ELISA kits (BD Pharmingen).

**Cytotoxicity assay.** Monocytes (5 × 10^4/well) were cultured for 6 h in the medium alone or with γ-irradiated (2500 cGy) HPC-4 or HPC-4_GFP or HPC-4_TLG cells (5 × 10^3/well), and then non-irradiated HPC-4 cells (2 × 10^4/well) were added and cultured for a further 18 h. The culture medium was removed and 100 μl of MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) (2 mg/ml, Sigma) dye solution was added for the last 2 h. Cell proliferation was assessed by the reduction of MTT. The data was expressed as the percentage of cytotoxicity calculated, as previously described [16].

**Statistical analysis.** The one-way analysis of variance ANOVA and post hoc Tukey test was used to calculate the statistical significance between individual treatments. Differences were considered as significant at p < 0.05.

**Results**

**TLG-transduced tumor cells**

The cDNA coding for human 26 kDa TNF was fused to the 5' end of the cDNA coding enhanced version of the GFP protein. The short sequence of cDNA — CTC GAG [(GGC)4 TCT]4 GCC — coding for a small flexible linker was inserted between TNF-cDNA and GFP-cDNA to obtain fused gene designated as TLG. This gene was cloned in the retroviral vector pMIGR1 [12] to obtain the final retroviral vector pTLG (Figure 1). The pTLG expression vector was used as a delivery system to HPC-4 tumor cells. The effectiveness of transduction ranged from 0.1% to 0.5%. Transduced HPC-4_TLG cells following FACS sorting, yielded 1–10% TLG^+ cells. After propagation of these selected cells and re-sorting, the purity of TLG^+ cells exceeded 95% (Figure 2). In parallel, the HPC-4_GFP cells mock-transduced with the original empty pMIGR1 vector were also more than 95% GFP^+ (not shown). The HPC-4_TLG cells demonstrated stable fluorescence of the TLG fusion protein, even after 30 days of maintenance in vitro in cultures with nonselective medium. The mean fluorescence intensity (MFI) of the HPC-4_TLG cells was about 4,000 versus 200 MFI for the nontransduced cells. The percentage of TLG positive cells decreased to about 60–70%

**Figure 1.** Linear scheme of the retroviral vector pTLG (B) construction using the parental pMIGR1 vector (A).

The retroviral fragment of the vectors contain the following sequences: TNF — human pre-TNFα-cDNA (719 bp); LTR — retroviral Long Terminal Repeats; ψ2 — sequence necessary for packaging viral RNA into virus capsids; IRES — sequence Internal Ribosomal Entry Site from EMCV virus; GFP — gene coding enhanced form of Green Fluorescent Protein; LIN23 — sequence coding for linker [(Gly)4Ser]4 Ala. The schemes of pMIGR1 and pTLG vectors do not show the prokaryotic fragment of the entire plasmid. The schemes are not drawn to scale.

**Figure 2.** TLG positive HPC-4 cells detected by flow cytometry. Left dot plot — three days after transduction; middle — three days after propagation of FACS sorted initially TLG positive cells; right — three days after propagation of FACS re-sorted TLG^+ cells. Data from one representative experiment out of five performed is shown. The numbers show percentage of TLG positive cells in a region set according to green fluorescence (FL1).
by day 30 post-transduction. However, the subsequent FACS sorting allowed selection of stable transductants expressing the TLG fusion protein for an additional 2–3 months in culture.

**TNF expression in the TLG-transduced tumor cells**

The amount of TNF secreted from HPC-4$_{TLG}$ cells was 2,428 ± 466 pg/ml (mean ± SEM), while mock-transduced HPC-4$_{GFP}$ and parental HPC-4 cells did not produce TNF (Figure 3). No IL-6, IL-10, MIP1α, or RANTES were found in the supernatants of HPC-4$_{TLG}$ cells (data not shown).

As membrane bound TNF is regarded as important in the cellular interactions [17], in the next step we analyzed whether synthesized TLG fusion protein was expressed as the membrane-bound form. For this purpose, transduced HPC-4$_{TLG}$ cells were stained with anti-TNF PE-conjugated mAb. However, taking into consideration significant spillover of green fluorescence from enhanced GFP into PE channel (data not shown), we took an advantage from anti-GFP antibodies conjugated with Alexa Fluor 594, the dye which did not interfere with PE emission spectrum. Anti-GFP antibodies were used in parallel with anti-TNF mAb for cell staining. After staining, the cells were analyzed by flow cytometry according to FL4 (anti-GFP) and FL2 (anti-TNF) fluorescence. In comparison to control HPC-4 and HPC-4$_{GFP}$ cells, two-color flow cytometry analysis of HPC-4$_{TLG}$ cells showed that 30.9% of cells stained with anti-TNF mAb, 31.3% were GFP+ while 19.5% were both TNF and GFP positive (Figure 4). Intracellular staining of HPC-4$_{TLG}$ cells with anti-TNF and anti-GFP Abs showed 40.6% TNF+ and GFP+ 95.3% cells (Figure 5A). Anti-GFP Ab stained 84.3% of HPC-4$_{GFP}$ cells and none of parental HPC-4 cells.

Figure 3. TNF secretion by: (A) HPC-4, (B) HPC-4$_{GFP}$ and (C) HPC-4$_{TLG}$ cells. Data is based on five independent experiments, two replicates per treatment, and expressed as pg/ml/1 × 10$^5$ cells/18 hours. Mean values ± SEM are shown; HPC-4$_{TLG}$ vs. HPC-4 and HPC-4$_{GFP}$, p < 0.001

Figure 4. Expression of membrane-bound TLG fusion protein by HPC-4, HPC-4$_{GFP}$ and HPC-4$_{TLG}$ cells. Cells were stained with PE-conjugated anti-TNF mAb and anti-GFP Alexa Fluor-594 conjugated Ab, and were analyzed by flow cytometry according to FL2 (PE) vs. FL4 (Alexa Fluor-594) fluorescence. Representative data from one out of four experiments performed is shown.
The intracellular co-expression of TNF and GFP in HPC-4 TLG cells was confirmed by Western blot analysis of lysates from TLG transduced cancer cells (Figure 5B).

**Induction of cytokines in monocytes stimulated with TLG-transduced tumor cells**

Monocytes were cultured for 18 h with HPC-4 TLG or HPC-4 GFP or parental HPC-4 cells to test whether they stimulated differently TNF, IL-10 and IL-12 production in monocytes. In comparison to parental HPC-4, HPC-4 TLG cells induced significantly higher release of TNF, while the induction of IL-10 was significantly lower, and no IL-12 was detected (Figure 6). Interestingly, monocytes stimulated with mock-transduced HPC-4 GFP cells produced significantly less TNF and no IL-10 or IL-12 compared to those stimulated with parental HPC-4 cells.

**Figure 5.** Expression of intracellular TLG fusion protein by: (A) flow cytometry and (B) Western blot analysis.

**Figure 6.** Secretion of TNF, IL-10 and IL-12 by monocytes cultured in the medium (Med) or with: (A) HPC-4 or (B) HPC-4 GFP or (C) HPC-4 TLG cells for 18 h. No cytokines were released by tumor cells except HPC-4 TLG, which secreted 900 ± 180 pg/ml of TNF/3 × 10⁶ cells/ml (not shown).

The results of five different experiments, two replicates per treatment (mean ± SEM) are shown; for TNF: HPC-4 vs. HPC-4 TLG p < 0.01, HPC-4 GFP vs. HPC-4 TLG p < 0.01; for IL-10: HPC-4 vs. HPC-4 TLG p < 0.01, HPC-4 vs. HPC-4 GFP p < 0.001; for IL-12: HPC-4 vs. HPC-4 GFP and HPC-4 TLG p < 0.01.
We have previously shown that short pre-exposure of monocytes to tumor cells deactivated them, as evidenced by decreased TNF and enhanced IL-10 production [11]. In order to check whether transduced HPC-4 cells act similarly, monocytes were cultured for 4 h with parental HPC-4 or transduced HPC-4GFP or HPC-4TLG cells. CD14+ cells were then isolated by FACS sorting and stimulated with HPC-4 cells for 18 h. Figure 7 shows that pre-exposure of monocytes to parental HPC-4 cells inhibited TNF and IL-12 and enhanced IL-10 production. In contrast, HPC-4TLG cells neither caused inhibition of TNF and IL-12 synthesis nor enhancement of IL-10 production. Pre-exposure of monocytes to mock-transduced HPC-4 GFP cells caused similar production of TNF, IL-10 and IL-12 as those pre-exposed to parental cells.

**Cytotoxicity of monocytes pre-exposed to γ-irradiated HPC-4TLG**

Monocytes/macrophages possess significant spontaneous cytotoxicity against tumor cells [16], which is decreased by the pre-exposure to tumor cells. This decrease is a part of the monocytes’ deactivation. To check whether HPC-4TLG cells may also prevent inhibition of cytotoxicity, monocytes were preincubated with γ-irradiated HPC-4 or HPC-4GFP, or HPC-4TLG cells, and cytotoxicity against HPC-4 tumor cells was measured. As expected, monocytes pre-exposed to HPC-4 or HPC-4GFP cells demonstrated lowered cytotoxicity (Figure 8). In contrast, there was no inhibition of cytotoxicity when monocytes were pre-exposed to HPC-4TLG cells.

**Discussion**

For a successful cancer gene therapy, it is fundamental to obtain as many cells transduced with the therapeutic gene as possible. Thus, knowing the actual gene transfer rate is critical. The GFP has made a breakthrough in gene transfer technology. It is a small protein that fits most of the currently used vectors and can be fused with other proteins to yield therapeutic fusion proteins [5, 18, 19]. The use of GFP fusion proteins provides enhanced sensitivity and resolution compared to antibody staining techniques and permits kinetic studies of a fusion protein localization [4, 20]. The construction of the fusion protein often requires the linking of two proteins or protein domains by an appropriate polypeptide linker. Many studies have suggested that the flexibility and hydrophilicity of the linker were very important in preventing the disturbance of the functions of both domains [21, 22].

In the present study, the TLG fusion gene was constructed by joining the TNF-cDNA and enhanced GFP-cDNA with the LIN23-DNA, coding for a short, flexible and hydrophilic linker Leu Glu [(Gly)4 Ser]4 Ala, which allowed retention of GFP and TNF expression. The retroviral vector pTLG carrying the TLG fusion gene was used for transduction of HPC-4 cells. The TLG gene transduced cells exhibited unchanged morphology and proliferation during in vitro culture for more than six months as compared to the parental nontransduced cells. The TLG gene transduced cells secreted a substantial amount of TNF, the presence of which was observed on their mem-
bran and intracellularly. One could argue that only a part of the GFP+ cells express TNF at the same time (Figures 4 and 5). In the preliminary set of experiments, we compared membrane and intracellular expression of TNF in HPC-4TLG cells without anti-GFP staining. Under such conditions, the amount of double positive cells was only around 50–60% of GFP+ cells (for both membrane and intracellular staining — data not shown). Such a relatively small amount of double positive cells could be a result of a spillover of green fluorescence from enhanced GFP to the PE channel, which could not be fully compensated during FACS analysis. To resolve this issue, we changed our detection method and introduced staining with anti-GFP Ab conjugated with Alexa Fluor 594, a dye which does not interfere with PE. This approach, however, did not improve the expression ratio, neither in surface nor intracellular staining. Thus, the relatively low number of double positive HPC-4TLG cells during membrane staining was probably due to secretion of TNF, cleaved rapidly from the TLG protein, to the medium. In the case of intracellular staining, the high amount of TLG present in the cytoplasm is probably neither properly folded nor fully accessible for both antibodies used for FACS analysis. Such an explanation is supported by Western blot analysis of HPC-4TLG cell lysates showing intracellular expression of the fusion protein (Figure 5B). It shows that there is also a band of free GFP present, which was probably left after the cleavage of TNF from TLG and its secretion. However, the exact trafficking and processing of the TLG fusion protein in transduced HPC-4 cancer cells has not been investigated in the present paper, and should be addressed in our future studies.

It has been previously shown that monocytes respond to tumor, but not normal, cells by producing several toxic mediators: TNF, ROI, radical nitrogen intermediates (RNI) and increased cytotoxicity [16, 23, 24]. On the other hand, clinicopathological evidence suggests that TIM, of which monocytes are precursors, may enhance the growth of some tumors [25]. Our previous data showed that although the first contact of monocytes with tumor cells induces anti-tumor activity, secondary contact leads to their deactivation (diminished TNF, IL-12 and ROI production, decreased cytotoxicity and enhanced IL-10 production), i.e. they became M2 polarized monocytes [25].

In the present experiments, we have investigated how the contact with tumor cells expressing TNF changes the activity of monocytes. In comparison to parental cells, HPC-4TLG cells induced higher production of TNF by monocytes and lower IL-10. Contrary to what could be expected, we found that HPC-4GFP cells, compared to parental HPC-4 cells, caused a decrease in the secretion of all tested cytokines by monocytes. Although we have no formal proof, this phenomenon may be due to more accelerated apoptosis of GFP transduced cells [26], which are rapidly phagocytosed by monocytes during their co-culture. Phagocytosis of apoptotic cells by monocytes, as evidenced by us and others, does not lead to secretion of TNF, IL-10 or IL-12. The only cytokine released by monocytes/macrophages during phagocytosis of apoptotic cells is TGFβ, which was not analyzed in our study. Another concern may arise when analyzing TNF secretion in the co-culture of monocytes and HPC-4TLG cells. In this case, the question is whether the TNF release comes from the transduced cells (GFP-TNF), or from the monocytes alone. Indeed, in this situation both cell populations may be a source of TNF; however, as indicated in the legend of Figure 6, the amount of TNF secreted by HPC-4TLG cells (900 ± 180 pg/ml of TNF/3 × 10^5 cells/ml) is far less compared to the level secreted by monocytes, thus having little effect on the total level of this cytokine in the co-culture supernatants.

The CD14+ monocytes sorted out from the co-culture with HPC-4 cells and restimulated with HPC-4, showed decreased TNF and IL-12 production. However, CD14+ cells isolated from the co-culture with HPC-4TLG produced a higher level of TNF and IL-12 and a diminished level of IL-10. This may suggest that, in the presence of TNF on the cell surface of HPC-4TLG or released by them, no deactivation of monocytes occurred. The observation that cytotoxic activity of monocytes pre-exposed to HPC-4TLG cells was not diminished also confirmed this suggestion.

The ability of TNF to activate monocytes/macrophages has been widely reported [17, 27, 28]. Activated monocytes/macrophages produce several toxic mediators such as RNI, ROI and TNF, which are probably directly involved in tumor cell killing in vitro. TNF may be also involved in indirect tumor cell killing by inducing RNI and ROI [29]. The inhibition of tumor growth after transduction of cancer cells with DNA construct coding for TNF has been observed, although different experimental models and divergent results concerning the activity of membrane-bound vs. secreted TNF have been reported [6, 17, 30]. It has been suggested that in immunocompetent mice, tumor regression was T cell-dependent [6, 30], while in immunodeficient mice the effect was macrophage-dependent [6]. Membrane bound TNF is regarded as important in the cellular interactions, e.g. with murine macrophages [17]. In keeping with this, HPC-4TLG cells effectively stimulated human monocytes for TNF release, while decreasing IL-10 and IL-12 secretion.

In summary, the present data indicates that human cancer cells transduced with TNF-cDNA release TNF and express its membrane-bound form. These
cells induced a higher TNF production by monocytes, and prevented their deactivation, thus giving a rationale for tumor cell based immunotherapy of cancer.

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References