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ORIGINAL ARTICLE

Gene-modified T cells for adoptive immunotherapy of renal cell cancer maintain transgene-specific immune functions in vivo

Cor H. J. Lamers · Sabine C. L. Langeveld · Corrien M. Groot-van Ruijven · Reno Debets · Stefan Sleijfer · Jan Willem Gratama

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Abstract

Background We have treated three patients with carboxyanhydrase-IX (CAIX) positive metastatic renal cell cancer (RCC) by adoptive transfer of autologous T-cells that had been gene-transduced to express a single-chain antibody-G250 chimeric receptor [scFv(G250)], and encountered liver toxicity necessitating adaptation of the treatment protocol. Here, we investigate whether or not the in vivo activity of the infused scFv(G250)⁺ T cells is reflected by changes of selected immune parameters measured in peripheral blood.

Methods ScFv(G250)-chimeric receptor-mediated functions of peripheral blood mononuclear cells (PBMC) obtained from three patients during and after treatment were compared to the same functions of $scFv(G250)^+$ T lymphocytes prior to infusion, and were correlated with plasma cytokine levels.

Results Prior to infusion, scFv(G250)⁺ T lymphocytes showed in vitro high levels of scFv(G250)-chimeric receptormediated functions such as killing of CAIX⁺ RCC cell lines and cytokine production upon exposure to these cells. High levels of IFN- γ were produced, whilst production of TNF- α , interleukin-4 (IL-4), IL-5 and IL-10 was variable and to lower levels, and that of IL-2 virtually absent. PBMC taken from patients during therapy showed lower levels of in vitro

C. H. J. Lamers (⊠) · S. C. L. Langeveld · C. M. Groot-van Ruijven · R. Debets · J. W. Gratama Laboratory for Clinical and Tumor Immunology, Department of Medical Oncology, Erasmus MC, Daniel den Hoed Cancer Center, PO Box 5201, 3008 AE Rotterdam, The Netherlands e-mail: c.lamers@erasmusmc.nl

S. Sleijfer

Department of Medical Oncology, Erasmus MC, Daniel den Hoed Cancer Center, PO Box 5201, 3008 AE Rotterdam, The Netherlands scFv(G250)-receptor-mediated functions as compared to pre-infusion, whilst IFN- γ was the only detectable cytokine upon in vitro PBMC exposure to CAIX. During treatment, plasma levels of IFN- γ increased only in the patient with the most prominent liver toxicity. IL-5 plasma levels increased transiently during treatment in all patients, which may have been triggered by the co-administration of IL-2.

Conclusion ScFv(G250)-receptor-mediated functions of the scFv(G250)⁺ T lymphocytes are, by and large, preserved in vivo upon administration, and may be reflected by fluctuations in plasma IFN- γ levels.

Keywords Immunogene therapy · Human T lymphocytes · Single chain chimeric receptor · Renal cell cancer · Clinical study

Introduction

Adoptive transfer of autologous T lymphocytes that have been gene-transduced to express antigen-specific receptors is a potentially attractive therapy to provide tumor-specific immunity to cancer patients [1-4]. We have studied safety and proof of this concept in patients with metastatic renal cell cancer (RCC) [5–7]. To this end we have constructed a single-chain antibody-type (scFv) receptor based on murine monoclonal antibody (mAb) G250 [8]. This mAb recognizes an epitope on carboxy-anhydrase-IX (CAIX) that is over-expressed by RCC cells [9]. Following retroviral introduction of the scFv(G250) transgene into primary human T-cells, the scFv(G250) receptor is expressed on the surface of these cells, which enables them to recognize CAIX and subsequently to exert antigen-specific effector functions in terms of cytokine production and killing of CAIX⁺ RCC cell lines [5, 10].

In a phase I study we have treated three patients with metastatic RCC with autologous $scFv(G250)^+$ T cells and subcutaneous (s.c.) injections of low dose interleukin-2 (IL-2). Unexpectedly, liver toxicity was encountered, after which the study was temporarily put on hold [7]. Examination of liver tissue from patient 1 revealed a discrete cholangitis with T-cell infiltration around the bile ducts and CAIX expression on the bile duct epithelium. The observed liver toxicity, reflected by liver enzyme disturbances, was most likely to be due to specific interactions of the $scFv(G250)^+$ infused T cells with CAIX⁺ epithelial cells lining the bile ducts.

We hypothesized that the liver toxicity reflects a specific immune reaction by the infused $scFv(G250)^+$ T cells and expected that this reaction would be reflected in changes of immune parameters measured in peripheral blood during and after therapy. Therefore, we compared functions of the $scFv(G250)^+$ T cells prior to and post infusion—the latter using peripheral blood mononuclear cells (PBMC) taken from the patients after infusion. We tested scFv(G250)-chimeric receptor-mediated functions, such as killing of CAIX⁺ RCC cell lines and cytokine production upon exposure to CAIX, and determined plasma cytokine levels.

Materials and methods

Treatment schedule

The inclusion and exclusion criteria for patients participating in this study have been reported previously [7]. The main eligibility criteria were that patients had disseminated clear cell RCC not amenable for surgery, and that the primary tumor had been removed by nephrectomy and expressed CAIX.

Autologous T lymphocytes were transduced with the scFv(G250)-chimeric receptor using retroviral transduction [11, 12]. The treatment schedule consisted of i.v. infusions of autologous scFv(G250)-transduced T-cells at escalating doses of 2×10^7 cells at day 1; 2×10^8 cells at day 2; 2×10^9 cells at days 3–5 (treatment cycle 1) and 2×10^9 cells at days 17–19 (treatment cycle 2), combined with s.c. human recombinant interleukin-2 (rIL-2, Chiron, Amsterdam), 5×10^5 IU/m² twice daily administered at days 1–10 and days 17–26. The treatment protocol had been approved by the governmental and institutional medical ethical review boards. Written informed consent was obtained from all patients.

Collection of peripheral blood mononuclear cells and plasma

Peripheral venous blood samples were drawn prior to, during and after treatment. During therapy blood was obtained prior to the first IL-2 injection unless indicated otherwise. PBMC were isolated from sodium heparin-anti-coagulated blood by Ficoll density gradient centrifugation and were cryopreserved in liquid nitrogen. Plasma was harvested from EDTA-anti-coagulated blood by centrifugation for 10 min at $1,000 \times g$ and stored at -70° C.

Flow cytometric analysis of transduced T-cell infusions and patient blood samples

The expression of the scFv(G250) receptor on transduced T-cell cultures was studied by indirect immunofluorescence using G250 anti-idiotype mAb NuH82 [10, 13]. The absolute numbers of scFv(G250)⁺ T-cells in the blood of patients prior to and following infusions of transduced T-cells were assessed as described [14].

Assessment of scFv(G250)-chimeric receptor-mediated functions in transduced T-cell infusions and patient blood samples

Cultures of scFv(G250)-gene modified T cells to be used for adoptive immunogene therapy, as well as patient PBMC obtained prior to and after infusion of gene-modified T cells, were monitored for scFv(G250)-chimeric receptormediated functions, i.e. specific cytotoxic activity and cytokine production, using the CAIX⁺ RCC cell line SKRC17-MW1 (clone 4) [10, 11]. A scFv(G250)-chimeric receptorexpressing T-cell clone (Clone 46) was used as positive control in these assays.

Cytotoxic activity was measured in a standard 4 h ⁵¹Crrelease assay, at 2,500 target cells per well and effector cell/ target cell ratios (E/T) of 80, 40, 20, 10, 5, 2.5, 1.25 and 0.63:1. The activated kill (AK)-activity of the cultured T cells and patient PBMC was blocked by adding a 30-fold excess of non-⁵¹Cr-labelled ("cold") AK-sensitive target cells (i.e. K562) to the test. The specificity of the scFv(G250)-mediated effector functions was verified by blocking CAIX on the target cells by adding G250 mAb (at a saturating final concentration of 25 µg/ml) to the target cells 15 min before addition of the effector cells [10, 11].

The specific cytolytic activity, i.e. specific ⁵¹Cr-release (see Fig. 1, upper panels), was expressed as LU_{20} per 10⁶ T lymphocytes, in which one lytic unit (LU_{20}) is defined as the number of effector cells required to lyse 20% of 2,500 target cells in a standard 4 h ⁵¹Cr-release test [15]. The net CAIX-specific cytolysis is defined as the level of cytolysis on CAIX⁺ RCC cells in the presence of cold K562 targets minus the level of cytolysis on the CAIX⁺ RCC cells in the presence of cold K562 targets and blocking G250 mAb.

For comparison of level of CAIX-specific activities between the T cell infusions of the individual patients the specific cytolytic activity and cytokine production were



Fig. 1 Carboxy-anhydrase-IX-specific cytolysis by scFv(G250)-gene modified T cell cultures used for adoptive immunogene therapy. The *upper panels* show representative experiments from each individual patient. The results are expressed as % ⁵¹Cr-release (4 h assay) at eight effector-to-target (E:T) cell ratios. Target cell were derived from the CAIX⁺ RCC cell line SKRC17-MW1 clone 4, alone (*dotted line*, for patient #2 and #3 only), in presence of a 30-fold excess of 'cold' K562 cells (*closed circles*), and in presence of a 30-fold excess of "cold" K562 cells and 25 µg/ml (final concentration) G250mAb (*open circles*). The net CAIX-specific cytolysis was defined as the level of

expressed per $10^6 \text{ scFv}(G250)^+$ T cells. In addition, to correct for assay variation these activities were related to the positive test control.

Assessment of cytokine levels

The concentrations of IFN- γ , TNF- α , IL-2, IL-4, IL-5 and IL-10 in plasma samples and culture supernatants (see above) were determined by cytokine bead arrays (BD Biosciences, San Jose, CA, USA), according to suppliers' specifications, and expressed in pg/ml. The detection limits of the assay were 2.6 pg/ml for IL-2 and IL-4, 2.5 pg/ml for IL-5, 2.8 pg/ml for IL-10 and TNF- α and 7.1 pg/ml for IFN- γ . Cytokine production by scFv(G250)-gene modified T cells was expressed as ng cytokine produced by 10⁶ T cells per 24 h.

Assessment of transgene copy number by quantitative real-time PCR

Genomic DNA was isolated from aliquots of the transduced T-cell infusions and from blood taken prior to, during and

cytolysis on CAIX⁺ RCC cells (in the presence of cold K562 targets) minus the level of cytolysis on CAIX⁺ RCC cells (in the presence of cold K562 targets and blocking G250 mAb). From these data, CAIXspecific lytic units (LU₂₀) were calculated according to Pross et al. [15]. The *lower panel* shows the LU₂₀-values per 10⁶ scFv(G250)⁺ T cells for the individual T-cell infusions and the mean value (*horizontal line*) per patient. The *open circles* in the *lower panel* indicate the positive test control results. Differences between patients were assessed by Student's *t*-test (two-sided); *P* values ≤ 0.05 are indicated. *Pat* indicates patient, *C* indicates treatment cycle

after therapy using the QIAamp[®] DNA mini kit (Qiagen, Hilden, Germany). The quantitative real-time PCR was performed as described [14], and data is presented as number of scFv(G250) DNA copies per μ l of blood.

Results

Liver toxicity following scFv(G250)⁺ T cell infusions

The three patients initially tolerated the infusions of the gene-modified T-cells well. However, following four to five infusions liver enzyme disturbances developed, reaching CTC grade 4 in patient 1, grade 2 in patient 2 and grade 3 in patient 3 (Fig. 2). The severe liver toxicity in patient 1 also included elevated levels of bilirubin. The severity of the liver toxicity was less during the second treatment cycle than during the first in patient 2. These toxicities required cessation of treatment in patients 1 and 3, corticosteroid treatment for patient 1 and reduction of the maximal T-cell dose to 2×10^8 total T-cells per infusion in patients 2 and 3.



Fig. 2 Kinetics of liver enzyme disturbances in the three patients before, during and following treatment with $scFv(G250)^+T$ -cells. Treatment intervals are indicated by *horizontal bars*; T-ly, daily infusions of retargeted T lymphocytes; IL-2, 0.5×10^6 IU rIL-2 s.c. every 12 h; Corticosteroids, i.v. solumedrol (days 8–10) and oral prednisolon (days 11–36). Treatment was stopped at day 5 after the 4th T-cell infusions as per protocol. For each patient, different linear scales were used for the *y*-axes in order to improve legibility of the panels

Characteristics of transduced T-cells immediately prior to infusion

The ex vivo generation of gene-modified T-cells was highly successful [11, 12]. The proportions of scFv(G250)+ T cells among the infused T cells were 53% for patient 1, 52% and 76% for patient 2 (treatment cycle 1 and 2,

respectively), and 63% for patient 3. Due to the reduced numbers of infused T-cells and early cessation of treatment in patients 1 and 3, the total number of infused scFv(G250)⁺ T-cells varied widely between patients, namely 2.13×10^9 cells for patient 1, 0.85×10^9 cells for patient 2 and 0.38×10^9 cells for patient 3 [7].

In vitro screening of gene-modified T-cells prior to infusion demonstrated high levels of scFv(G250)-mediated cytolysis of a CAIX⁺ RCC cell line (Fig. 1, upper panels). The CAIX-specific cytolytic activity of the scFv(G250)⁺ T-cells of patient 1 [i.e. median value 372 (range 186–433) LU_{20} per 10⁶ scFv(G250)⁺ T cells] was significantly higher than those of patient 2 [i.e. 93 (81–116) LU_{20} per 10⁶ cells] and patient 3 [i.e. 88 (51–143) LU_{20} per 10⁶ cells]. This pattern was also observed when test results were corrected for between-test variation (Fig. 1, lower panel). As a result, the total scFv(G250)-mediated cytolytic capacities of the T-cell infusions varied widely between patients, i.e. 792,204 LU_{20} for patient 1, 78,774 LU_{20} for patient 2 and 33,274 LU_{20} for patient 3 [7].

Upon stimulation with CAIX⁺ RCC cells, the scFv(G250)⁺ T-cells of individual patients showed a characteristic cytokine production pattern (Table 1 and Fig. 3). All T-cell cultures produced predominantly IFN- γ [ranging between the patients from 36.9 to 55.9 ng IFN γ per 10⁶ scFv(G250)⁺ T cells during 24 h], and to lower levels, IL-5 (ranging from 1.2 to 5.4 ng) and IL-4 (ranging from 0.3 to 2.9 ng). Only low levels of TNF- α , IL-2 and IL-10 were produced, although scFv(G250)⁺ T cells of patient 2 produced relatively high amounts of IL-10. These patterns were also observed when test results were corrected for between-test variation (not shown). In the absence of CAIX-triggered stimulation, scFv(G250)⁺ T-cells showed no detectable levels of spontaneous cytokine production (data not shown).

As a consequence of the widely different numbers of infused $scFv(G250)^+T$ cells between patients, the calculated

Table 1 Carboxy-anhydrase-IX-specific cytokine production capacity of infused $scFv(G250)^+T$ cells

•			
Cytokine	Patient 1	Patient 2	Patient 3
IFN-γ	42.0	55.9	36.9
TNF-α	0.3	1.4	1.1
IL-2	nt	0.1	0.0
IL-4	2.9	1.3	0.3
IL-5	5.4	4.9	1.2
IL-10	0.1	1.5	0.0

Cytokine production capacity: ng per 10^6 scFv(G250)⁺ transduced T cells per 24 h

Total number of infused scFv(G250)-transduced T cells: patient 1: 2.13×10^9 , patient 2: 8.45×10^8 , patient 3: 3.79×10^8

nt not tested

100

50

IFN₇ [ng]

IL-4 [ng]

a

Pat 1/C1

Pat 1/C1

p=0.02

Pat 2/C1+2 Pat 3/C1

p=0.01

c

Pat 3/C1

p=0.04

o

Pat 2/C1+2

ΓNFα [ng]

IL-5 [ng]

Fig. 3 Carboxy-anhydrase-IXspecific cytokine production (IFN-γ, TNF-α, Il-2, IL4, IL-5 and IL-10) by scFv(G250)-gene modified T-cells in the individual T cell infusions, expressed as the net amount [ng] of cytokine produced by $10^6 \text{ scFv}(G250)^+ \text{ T}$ cells during 24 h of co-cultivation with the CAIX⁺ RCC cell line SKRC1-MW1 clone 4. For each cytokine, different linear scales were used for the y-axes in order to improve legibility of the panels. See further legend to Fig. 1



1.5

1.0

0.

0.0

Pat 1/C1

IL-10 [ng]

p<0.001

p<0.001

c

Pat 2/C1+2

Pat 3/C1

capacity of scFv(G250)-mediated IFN- γ production of the infused gene-modified T-cells ranged between 89.5 µg of IFN- γ per 24 h for patient 1, 47.2 µg for patient 2 and 14.0 µg for patient 3.

Monitoring of peripheral blood samples

The treatment episodes were associated with transient lymphopenia in all three patients, probably due to the coadministered rIL-2. Circulating T-cell numbers decreased immediately upon start of therapy to reach a nadir at approximately 50% of baseline values on day 4, to normalize again when treatment was stopped (not shown).

Circulating scFv(G250)⁺ T-cells and scFv(G250) DNA copies

ScFv(G250)⁺ T-cells became transiently detectable in the circulation of all three patients from day 3 of therapy onwards, and reached maximum levels between days 6 and 16 (Fig. 4, upper panels, and [14]). Their peak levels during the first therapeutic cycle were higher in patient 1 $[5.3 \text{ scFv}(G250)^+ \text{ T cells/µl}]$ than in patient 2 (2.6 and 1.6 cells/µl in treatment cycle 1 and 2, respectively) and patient 3 (0.8 cells per μ l), which were proportional to the numbers of scFv(G250)⁺ T-cells administered (see above). In parallel scFv(G250)-encoding DNA became transiently detectable in the circulation of all three patients. Here, only limited numbers of samples of patients 1 and 2 (first treatment cycle) were available for testing (Fig. 4, lower panels, and [14]). Again, the number of circulating scFv(G250) DNA copies was clearly higher in patient 1 (i.e. 44 copies/µl on day 10) than in patients 2 (7.1 copies/µl on day 17) and 3 (5.3 copies/µl on day 8).



p=0.03

Pat 2/C1+2

0

Pat 1/C1

p=0.02

o

Pat 3/C1

Peripheral blood mononuclear cells obtained prior to and at 1-4 days following the last T-cell infusion were assayed for in vitro scFv(G250)-mediated cytotoxicity and cytokine (IFN- γ , TNF- α , IL-2, IL-4, IL-5 and IL-10) production upon co-culture with a CAIX⁺ RCC cell line. In all three patients, CAIX-specific activities, i.e. cytolysis of CAIX⁺ RCC cells (Fig. 5, upper row of panels) and IFN- γ production (2nd row of panels) were detected. Virtually no scFv(G250)-receptor-mediated production of TNF- α , IL-2, IL-4, IL-5 and IL-10 was detectable (lower row of panels and data not shown). The ex vivo CAIX-specific activities of PBMC (i.e. cytolysis and IFN-y production) obtained during or immediately after administration of rIL-2 were higher than those of PBMC obtained several days after termination of rIL-2 treatment in patients 1 and 3 (upper and middle rows of panels). In contrast, patient 2 still showed significant levels of these activities at the start of cycle 2 (i.e. 1 week after cessation of IL-2 administration), which were paralleled by high levels of circulating $scFv(G250)^+ T$ cells and DNA copies (see before) and exceptionally high IFN-y plasma concentrations (see below),

Blood cytokine levels

Interleukin-2 became detectable during therapy; levels were typically between 5 and 20 pg/ml in blood samples drawn 12 h after s.c. injection of rIL-2 (Fig. 6, upper row). Relatively high IL-2 plasma concentrations were observed in blood samples taken shortly after rIL-2 injections (marked with asterisks in Fig. 6). The fluctuations in plasma IFN- γ levels varied greatly between patients and as a function of time (2nd row of panels): in patient 1, IFN- γ



Fig. 4 Kinetics of circulating $scFv(G250)^+$ T-cell counts (*upper panels*) and scFv(G250) DNA copy numbers (*lower panels*) in blood samples obtained before, during and following treatment with $scFv(G250)^+$ T cells in patient 1 and 2. Results of patient 3 have been published elsewhere (see Ref. [14]). In the *upper panels, closed circles connected by uninterrupted lines* indicate data points with proportions of $scFv(G250)^+$ cells above the detection limit of 6 in 10,000 CD3⁺ T

Treatment days

cells (see Ref. [14]). In the *lower panels*, the *dashed lines* indicate the detection limit as defined by the 95th percentile of ten blood samples of healthy donors, i.e. 1 scFv(G250) DNA copy per μ l. The mean number of scFv(G250) DNA copies per scFv(G250)⁺ T cell in the T cell infusions were 2.3 for patient 1, and 4.5 and 6.8 for treatment cycles 1 and 2, respectively, of patient 2. *Horizontal bars* indicate treatment intervals, see further legend to Fig. 2

levels increased during therapy to 76 pg/ml at day 4, to decline after day 5 when treatment was stopped. Remarkably, patient 2 had constitutively high IFN-y plasma concentrations prior to start of therapy (73 pg/ml), which transiently declined to 31 pg/ml during the first therapeutic cycle. Prior to the second cycle, IFN- γ levels had increased again to 115 pg/ml, but gradually declined during therapy to levels ~ 20 pg/ml on day 32. The IFN- γ levels of patient 3 remained relatively stable and fluctuated between 7 and 14 pg/ml during the observation period. The yIL-5 plasma levels became detectable in all three patients during therapy, to reach maximum levels following the last rIL-2 administration in treatment cycle 1, i.e. 97 pg/ml at day 5 in patient 1, 50 pg/ml at day 10 in patient 2, and 21 pg/ml at day 6 in patient 3 (3rd row of panels). IL-10 became detectable at low levels (i.e. <10 pg/ml) in all three patients (5th row of panels); this cytokine peaked at day 9 in patient 1 and at day 19 in patient 2. In the latter patient, this pattern coincided with only a minor IL-5 response during the second treatment cycle.

Virtually no TNF- α was detected prior to, during and after therapy except for one observation in patient 1 (4th row of panels). Finally, virtually no IL-4 responses were seen in patients 1 and 2 (lower row of panels); patient 3 stood out by detectable IL-4 levels prior to therapy, which remained at similar levels during and after treatment.

Discussion

We sought in vivo evidence for scFv(G250) receptor-mediated T-cell functions as the infusion of scFv(G250)-retargeted T-cells clearly correlated in time with the induction of liver toxicity. This toxicity is likely to be the result of the specific immune reaction of the retargeted T-cells directed against CAIX expressed by epithelial cells lining the bile ducts [7]. Here, we show that PBMC taken from patients on therapy-during which scFv(G250)⁺ T cells were detectable by flow cytometry and PCR-displayed similar scFv(G250)receptor-mediated functions as did scFv(G250)⁺ T-cells prior to infusion, such as cytokine production after exposure to CAIX and killing of CAIX⁺ RCC cell lines.

We focused on fluctuations in plasma cytokine levels as markers for the in vivo activity of the scFv(G250)⁺ T cells. ScFv(G250)⁺ T lymphocytes prior to infusion produced high levels of IFN- γ and moderate levels of IL-5 upon CAIX stimulation, whereas PBMC taken from patients during therapy produced only moderate levels of IFN- γ and no IL-5 upon ex vivo exposure to CAIX. On the contrary, we observed significant plasma levels of IFN- γ and IL-5 during therapy. The fluctuations in plasma IFN- γ levels varied strongly between patients, with the highest increment in patient 1 with the most pronounced liver toxicity. We have



Fig. 5 Kinetics of scFv(G250)-mediated cytotoxicity (*upper row*) and CAIX-specific cytokine production [IFN- γ (*2nd row*) and IL-5 (*3rd row*)] by PBMC obtained before and after therapy. PBMC were incubated with CAIX⁺ RCC cells (*open triangles*), and CAIX⁺ RCC cells in the presence of G250 blocking mAb (*closed triangles*). Specific cytolysis is expressed as LU₂₀/10⁶ PBMC and cytokine production in pg/ml. The net scFv(G250)-mediated cytolysis (*closed circles*) is defi-

ned as the level of cytolysis of CAIX⁺ RCC cells minus the cytolysis of CAIX⁺ RCC cells in the presence of G250 mAb. The net scFv(G250)-mediated cytokine production (*closed circles*) is defined as the amount of cytokine produced in presence of CAIX⁺ RCC cells minus the amount of IFN- γ produced in presence of CAIX⁺ RCC cells and G250 mAb. See also the legends to Fig. 2

no explanation for the aberrant kinetics of plasma IFN- γ levels in patient 2; an as yet unidentified (auto) immune activation, possibly combined with a 'high IFN- γ genotype' [16] given the elevated IFN- γ level prior to treatment, may have contributed to this pattern. The plasma IFN- γ levels in patient 2 normalized during the second course of treatment, which was paralleled by relatively high CAIX-specific production capacity of the inhibitory cytokine IL-10 and an increment of plasma IL-10 levels in this patient.

To determine whether the increments in plasma IFN- γ and IL-5 levels would reflect in vivo activity of the scFv(G250)⁺ T cells or be related to the IL-2 administrations only, we considered the following aspects.

Adoptive transfer of T cells has often been combined with administration of exogenous rIL-2 in order to promote survival of the transferred T cells, and to support and maintain T-cell activity in vivo [17–19]. Our observation that the highest levels of scFv(G250)-mediated in vitro activity of patient PBMC, in terms of both scFv(G250)-mediated cytotoxicity and cytokine production, were detected during the period of IL-2 administration, is in line with this paradigm.

Only few studies have addressed the effects of low-dose IL-2-based treatment protocols on plasma cytokine levels. Cragun et al [20] performed weekly assessments of serum cytokine levels in patients receiving daily injections of 3×10^{6} IU/m²/day for 6 weeks and showed a gradual increase of IL-5 from week 2 of treatment onwards, whilst levels of IFN-y, IL-4 en GM-CSF did not change. Gene expression profiling of PBMC following 3 h of in vivo or in vitro exposure to IL-2 identified 155 genes, including IL-5 and IFN- γ , whose expression levels were increased >threefold over those in non-exposed samples [21, 22]. In addition, multiplex protein analysis of RCC patient serum showed that IFN- γ and IL-5 levels increased significantly within 3 h following administration of a single high dose of rhuIL-2 (0.72 \times 10⁶ IU/kg). However, IFN- γ levels did not increase further upon repeated IL-2 administrations, in contrast to those of other cytokines such as TNF- α , IL-5 and IL-10 [23]. The discrepancy between the short term (i.e. within 4 h) [21, 22] and long term (i.e. 24 h) [20] effects of IL-2 administrations on IFN- γ and IL-5 plasma levels may be related to the different stabilities of IFN- γ and IL-5 mRNA in T-cells; their half-life is ~ 2 and ~ 4 h, respec**Fig. 6** Kinetics of plasma cytokine levels in blood samples of individual patients obtained before, during and following immuno-gene therapy. During the period of rIL-2 administration, blood samples were drawn just before s.c. rIL-2 injection, excepted for samples marked with *asterisks* (** = 2 h, and * = 4 h after rIL-2 administration); *dashed line*, detection limit. See further the legend to Fig. 2



tively [24, 25]. Given our scheme of IL-2 administrations (i.e. once every 12 h), the relatively long half-life of IL-5 mRNA may explain the gradual increment of plasma IL-5 levels. The shorter half-life of IFN- γ mRNA would preclude such an effect on plasma IFN- γ levels.

On the basis of these combined studies, we propose that the observed increments in plasma IL-5 levels are a 'direct' effect of the IL-2 injections, and probably do not reflect in vivo activity of the infused $scFv(G250)^+$ T cells. On the contrary, the observed increments in plasma IFN- γ levels cannot completely be attributed to the IL-2 injections—due to the short half-life of its mRNA—and therefore would reflect in vivo activity of $scFv(G250)^+$ T cells. This mechanism would especially be apparent in patient 1, which received the highest number of $scFv(G250)^+$ T cells, and developed the most severe liver toxicity.

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