CD47-targeted cancer immunogene therapy: Secreted SIRPα-Fc fusion protein eradicates tumors by macrophage and NK cell activation

Magdalena Billerhart,1 Monika Schönhofe,1 Hemma Schueffl,2 Wolfram Polzer,1 Julia Pichler,1 Simon Decker,1 Alexander Taschauer,1 Julia Maier,1 Martina Anton,3 Sebastian Eckmann,1 Manuel Blaschek,1 Petra Heffeter,2 Haider Sami,1 and Manfred Ogris1

1University of Vienna, Faculty of Life Sciences, Department of Pharmaceutical Sciences, Laboratory of MacroMolecular Cancer Therapeutics (MMCT), Althanstrasse 14, 1090 Vienna, Austria; 2Institute of Cancer Research, Department of Medicine I, Comprehensive Cancer Center, Medical University of Vienna, 1090 Vienna, Austria; 3Institutes of Molecular Immunology and Experimental Oncology, Klinikum rechts der Isar, Technische Universität München, Ismaninger Straße 22, 81675 Munich, Germany

INTRODUCTION

The immune system is a key component in growth control and eradication of cancer cells. Immune surveillance of tumors is divided into three phases, designated elimination, equilibrium, and escape.1 Tumor escape is promoted by the occurrence of an immune-suppressive microenvironment, which can also be triggered by hypoxia followed by localized expression of immune-suppressive cytokines like transforming growth factor-beta. Attraction of immune cells with suppressive functions, like regulatory T cells or M2 macrophages, then leads to the inhibition of natural killer (NK) and CD8+ cytotoxic T lymphocyte function.2 Body cells can overexpress surface proteins with immuno-modulatory functions: Programmed Cell Death-Ligand 1 (PD-L1) expression in heart prevents T cell-mediated inflammation. However, PD-L1 overexpression on tumor cells and its interaction with programmed cell death 1 (PD-1) receptor on T cells suppresses their inflammatory and hence antitumoral response.3 CD47 (integrin-associated protein [IAP]), a ubiquitously expressed transmembrane glycoprotein, acts as a “marker of self” and plays an important role in hematopoiesis. Upon binding to CD47 ligand signal regulatory protein α (SIRPα), expressed on macrophages and dendritic cells, phagocytosis of target cells is inhibited.4 Also on erythrocytes CD47 prevents their phagocytosis by macrophages, and loss of CD47 during senescence triggers their elimination. In hematopoietic cancers and also many solid tumors, CD47 is found to be upregulated and correlates with a poor prognosis.5 Also in cancer stem cells, the transmembrane protein level is increased.6 Therapeutic antibodies have proven to be an effective approach by specifically blocking a target on tumor cells and at the same time attracting and activating immune cells by the exposed Fc part triggering antibody-dependent cell-mediated cytotoxicity (ADCC). Cetuximab, for example, blocks the epidermal growth factor receptor (EGFR) and subsequent downstream signaling, whereas its immunoglobulin G (IgG1)-derived Fc successfully triggers both ADCC and complement activation (complement-dependent cytotoxicity [CDC]).7 As EGFR is often overexpressed on cancer cells...
both are based on IgG4. They show reduced complement activation and are being evaluated in clinical trials or are already approved, and CD47-binding antibodies like Hu5F9-G4 (magrolimab) and SRF231 of the CD47 antibody, and it can trigger severe anemia as a side effect.5

Several approaches are on the way to find a balance between the binding affinity for CD47 and triggering ADCC and/or CDC. High-affinity CD47-binding antibodies like Hu5F9-G4 (magrolimab) and SRF231 are being evaluated in clinical trials or are already approved, and both are based on IgG4. They show reduced complement activation and ADCC but also fewer side effects.5,6 As an alternative, fusion proteins have been developed that block CD47 with lower affinity but induce more potent immune activation. TTI-621 is a fusion protein constant for CD47 with TTI-621 is significantly lower than that of anti-CD47 antibodies (5F9: kD = 8 nM; 10 human SIRPα fusion protein ALX148 was developed, which combines a high-affinity SIRPα blockage and Fc functionality, and other control plasmids for deciphering the presence of combined/dual functionality.13 Recently, Trillium Therapeutics also developed an alternative, TTI-622, with similar a SIRPα domain but an IgG1 Fc domain, in an attempt to improve tolerability of the protein (ClinicalTrials.gov Identifier: NCT03530683). Weiskopf and colleagues designed a high-affinity variant of SIRPα, termed CV1, with a binding constant of 11 pM for human CD47.7 When a CV1-hlgG1 Fc fusion protein was applied, a significantly decreased hematocrit in NSG mice was observed. Full antitumoral activity in the absence of toxicity was achieved by application of the CV1 monomer combined with, e.g., rituximab in CD20-positive lymphoma xenografts. The fusion protein ALX148 was developed, which combines a high-affinity SIRPα version with an IgG Fc part lacking any CDC or ADCC activity.13 In syngeneic and xenograft tumor models, the combination of ALX148 with PD-L1 or CD20-blocking antibodies was necessary for potent antitumoral activity. In non-human primates, ALX148 alone was non-toxic up to the highest dose tested, i.e., 100 mg/kg. Both TTI-621 and ALX148 are already evaluated in clinical trials to treat a rare T cell lymphoma13 or advanced head and neck cancer in combination with pembrolizumab (Kim et al., 2019, Am. Soc. Hematol., abstract). We and others have demonstrated that the tumor-restricted expression of secreted therapeutic proteins can be a vital approach, which also allows a potent bystander effect on adjacent, non-transfected cells in the near vicinity. With a synthetic gene delivery system, systemic application of plasmid vector resulted in significant tumor necrosis factor alpha (TNF-α) expression in implanted tumors.10 Interleukin-12 (IL-12), a highly potent cytokine with antitumoral activity, has been expressed in tumors after local delivery and is being investigated already in several clinical trials.17 For our present study, we designed a gene vector encoding for a secreted fusion protein having high-affinity SIRPα variant CV1 coupled to a IgG1 Fc part with maximized CDC and ADCC activity. Secretion is accomplished by the IL-2 secretion signal, which resulted in a potent bystander effect even at very low transfection efficiencies. This enables selective blocking of CD47 in cancer cells and finally leads to immune cell recruitment and tumor cell eradication (Figure 1).

**RESULTS**

**Design and cloning of a gene vector for secreted sCV1-hlgG1 fusion protein**

We have designed a set of plasmids, namely therapeutic plasmid pSCV1-hlgG1 with dual/combined functionality of CD47 blockage and Fc-mediated cytotoxicity, single-component control plasmids, and transfection control plasmid (for plasmids maps see Figure S1). All of these plasmids are based on the CpG-free backbone from pCpG-hCMV/SCEP-mCherry, to ensure optimal transgene expression in vitro and in vivo.18 As further controls to the therapeutic plasmid (pSCV1-hlgG1), plasmids were cloned that encode for either secreted hlgG1-Fc only (pshlgG1) or secreted SIRPα CV1 variant only (pSV1), or as a transfection control the fluorescent reporter mCherry (pCpG-hCMV/SCEP-mCherry, short: pmCherry) was used. All plasmids could be propagated as low-copy plasmids in *E. coli* DB3.1 λpir or GT115. With the designed plasmid at hand, we could produce ~3 mg of plasmid per liter of culture volume.

**sCV1-hlgG1 exhibits combined CD47 blockage and Fc functionality and potent bystander effect in vitro**

Before investigation of pSCV1-hlgG1 transfection and fusion protein functionality, expression of CD47 was measured by flow cytometry on human lung and triple-negative breast carcinoma cell lines A549 and MDA-MB-231 and their metastatic variants 231/LM2-4 and 231/LM2-4-EGFPLuc, respectively (Figure S2). All of them were clearly positive for CD47, with MDA-MB-231 cells exhibiting a higher level compared to A549. The screened cell lines were then transfected with plasmid pSCV1-hlgG1, encoding for the secreted fusion protein combining CD47 blockage and Fc functionality, and other control plasmids for deciphering the presence of combined/dual functionality. We measured both blocking of CD47 (with a competitive anti-CD47 binding assay) and the presence of human IgG1-Fc domain (with an anti-IgG1 antibody on the cell surface) as shown in Figure 2. With both readouts we observed an effect on all measured cells, i.e., a complete shift of the whole cell population. When stained for the Fc part of
the fusion protein, a strong shift in signal was observed on psCV1-hlgG1-transfected cells, whereas no signal above background was observed for psCV1- and pmCherry-transfected cells. Blockage of CD47 binding was most pronounced on psCV1-hlgG1-transfected cells, with an ~96% signal reduction. Of interest, on psCV1-transfected 231LM2-4 cells a partial block of 76% was observed, whereas on A549 cells CD47 access was blocked to a similar extent on psCV1-hlgG1- and psCV1-transfected cells (Figure S3). Using mCherry as a reporter plasmid, we observed under similar transfection conditions only 1%–2% mCherry-positive cells 48 h after transfection of 231LM2-4-EGFPLuc cells when analyzed by flow cytometry (Figure S4), whereas A549 cells could be transfected with an efficiency of >28% (data not shown). In view of the apparently strong bystander effect and to further substantiate the specificity, we next evaluated CD47 blockage and fusion protein detection as a function of the amount of transgene expressed and the effect of diluted supernatant (from psCV1-hlgG1-transfected cells) transferred to non-transfected recipient cells (Figure 3). CD47 blockage was evaluated similarly as in Figure 2, whereas cell surface-bound scV1-hlgG1 protein was measured by flow cytometry with fluorescently labeled protein A (which binds to the Fc domain). With a 1:2 diluted supernatant from transfected cells, CD47 access was strongly blocked in all recipient cells, whereas with a 1:10 diluted supernatant a minimal effect was seen, and none was seen with the 1:50 dilution (Figure 3A). Similar as for the anti-IgG Fc staining, protein A also staining allowed sensitive detection of Fc-positive cells even when treated with a 1:30 diluted supernatant. From a dot blot assay (Figure 3B) and a respective calibration curve, we estimated a concentration of scV1-hlgG1 in the supernatant corresponding to ~0.24 ng/μL. IgG protein, which corresponds to a concentration of 3.1 nM, produced within 48 h after initial transfection. The amount secreted was also sufficient to be detected by immunostaining of transfection supernatant-treated, untransfected cells (Figure 3C). Since the CV1 sequence was initially optimized for binding to human CD47, we also transfected a murine cell line to evaluate the protein for its binding affinity for murine CD47 (Figure S5), as Weiskopf et al. also demonstrated cross-reactivity with mouse CD47. Similar as for the two human cell lines evaluated, we could also detect efficient surface binding of fusion protein in the murine colon carcinoma cell line CT26 when staining the Fc part with fluorescently labeled protein A. Next, we studied the time course of in vitro transgene expression for scV1-hlgG1 within two different experimental setups, one over a time frame of 4–48 h and the other from 2 to 12 days. Earliest detectable CD47-blocking activity could be observed 6 h after transfection, although after 48 h it was most pronounced (Figure 4A; Figure S6). Surface-bound scV1-hlgG1 could already be detected at 6 h after transfection, and the signal increased up to 48 h (Figure 4B; Figure S6). When psCV1-hlgG1-transfected cells were tracked over days, CD47 blocking was still measurable after two 1:8 passaging steps (Figure 4C, p1 and p2). With an antibody directed against IgG-Fc, detection of bound scV1-hlgG1 protein was even possible also at passage 3, i.e., after 12 days of transfection (Figure 4D). This is noteworthy considering an initial transfection rate below 1.5% (see Figure S4) and with far less than 1% of cells expressing scV1-hlgG1 protein over the investigated duration (Figures 4C and 4D).

**Therapeutic activity in vivo**

Next we evaluated the effect of scV1-hlgG1 gene therapy in an orthotopic xenograft model of MDA-MB-231 human triple-negative breast cancer in CB17-severe combined immunodeficiency (SCID) mice (Figure 5). When implanting the highly malignant metastatic variant 231LM2-4-EGFPLuc percutaneously into the 4th pair of inguinal nipples, we observed, besides fast onset of local growth, axillary and
sentinel lymph node infiltration being positive for CD47 (Figures 5A and 5B). In two independent experiments with the same settings, 231/LM2-4-EGFPluc were transiently transfected in vivo 48 h prior to implantation with pmCherry, psCV1, or psCV1-hlgG1 or left untransfected and thereafter implanted. Tumor growth was semiquantitatively followed by bioluminescence imaging (BLI), and tumors were explanted at the end of the experiment. Whereas in experiment I measurements were continued until day 34, when animals in the untransfected tumor group reached endpoint criteria (reduced well-being, which was also due to metastasis), in experiment II the study was terminated and tumors explanted on day 25 after implantation. In the case of untransfected cells, the BLI signal doubled approximately every 4 days starting with implantation. In both experiments, psCV1-hlgG1-transfected tumors showed a significant, on average 50% reduced signal for >2 weeks after implantation, compared to the untransfected control group (Figures 5C and 5D). In individual tumors, this reduction was down to 0.1% of the starting signal. After day 15, re-growth started in some tumors, whereas at the end of the experiment (day 25 or 34 after implantation, respectively), only four of six tumors (8 of 12 in total) explanted were significantly smaller than the pmCherry-transfected tumors. The implan-}

---

**Figure 3. Bystander effect and fusion protein secretion in vitro**  

231/LM2-4 cells were transfected for 48 h, and the supernatant was harvested thereafter and either analyzed for fusion protein content or transferred onto untransfected 231/LM2-4 cells. (A) Cells treated with transfection supernatant for 1 h (diluted with complete medium at the indicated ratio) and thereafter stained with anti-CD47 antibody (B6H12.2) (secondary antibody goat anti-mouse IgG H&L, Alexa Fluor 647, right) or protein A (Alexa Fluor 488 conjugated, left). Representative histogram blots shown. Red line, cells incubated with supernatant from psCV1-hlgG1-transfected cells; blue line, cells incubated with supernatant from mock-transfected (buffer only) cells. (B) Dot blot analysis (2 μL/dot) with anti-human IgG Fc HRP antibody. Top (dots in duplicates): 48-h transfection supernatant diluted with complete medium at the indicated ratio, BSA solution (2 mg/mL), complete medium. Bottom: indicated amounts per dot of human IgG1 isotype. (C) Cells treated with 48-h transfection supernatant for 1 h

---

The growth curve of the untransfected cells showed a slightly higher size after harvesting was correlating (Figure 5F). However, the pmCherry transfection control group (Figure 5D). Also, the tumor size after harvesting was correlating (Figure 5F). However, the growth curve of the untransfected cells showed a slightly higher growth tendency compared to pmCherry and psCV1 tumors, which could be due to the intrinsic growth retardation effect of the ex vivo transfection process in the latter case. Nevertheless, after termination of the experiment pmCherry-transfected tumors (6/6 per experiment, 12/12 in total) were all large (diameter 10–15 mm) and showed signs of core necrosis, whereas untransfected tumors were exceeding a diameter of 20 mm (Figures 5E and 5F). We also quantified the volume of explanted tumors (Figure S7). In Exp I (terminated on day 34 after implantation), pmCherry-transfected tumors were significantly smaller than the untreated control and psCV1-hlgG1-transfected tumors were significantly smaller than the pmCherry-transfected tumors (Figure S7A). When terminating the experiment on day 25 (Exp II), the size of pmCherry-transfected tumors was slightly, but insignificantly, smaller compared to untransfected tumors (Figure S7B). Nevertheless, only sCV1-hlgG1-transfected tumors were significantly smaller than pmCherry-transfected tumors. The implantation of ex vivo transfected tumor cells could demonstrate that tumor-localized expression of sCV1-hlgG1 had a therapeutic effect. Furthermore, we were also interested in whether the plasmid vector is in principle suitable for a local therapeutic approach by inducing localized immune reaction and CD47 blockade. When forming plasmid-linear polyethyleneimine (LPEI) polyplex particles with a positive charge excess (N/P ratio 9) in a low-ionic buffer (5% glucose, 20 mM HEPES pH 7.4 [HBG]), we usually obtained particles sized ~120 nm in diameter (mode) and ζ-potential of approximately +30 mV, which were then injected intratumorally in 231/LM2-4-EGFPluc tumors. In a first series, tumors were explanted 24, 48, and 72 h after in vivo transfection. When comparing the different time points, at 24 h after transfection we observed areas of blocked CD47 in immunohistochemical (IHC)-stained slides, whereas such blockade was not seen in the control transfected tumors (Figure 6). Of interest, no blockade was observed at 48 and 72 h after transfection with psCV1-hlgG1. In a further in vivo series tumors were explanted 12, 17, and 24 h after intratumoral polyplex injection, and macrophage infiltration was qualitatively and semiquantitatively analyzed by staining for the murine macrophage antigen F4/80 (Figure 7). Although the level of macrophage infiltration was higher at all time points in psCV1-hlgG1-transfected tumors compared to the
pmCherry-transfected control group, only at 17 h after transfection did psCV1-hIgG1-transfected tumors show significantly elevated macrophage infiltration. This early time point is also in line with the in vitro findings, where secretion of sCV1-hIgG1 was detected already after 6 h of transfection (Figure 4).

**NK cell activation**
To detect potential infiltration of tumors with NK cells, sections were stained with anti-natural cytotoxicity triggering receptor 1 (NCR1) antibody, which is exclusively expressed on NK cells. Unfortunately, we could neither verify nor falsify this hypothesis in our model, as all NK cell antibodies tested in our laboratory resulted in high background staining, which did not allow detection of any differences. Therefore, to study the effect of sCV1-hIgG1 on human NK cells in vitro, we developed a luciferase-based in vitro assay that should allow monitoring of NK cell-mediated target cell lysis in vitro. NK cell-mediated lysis of luciferase-expressing tumor cells co-transfected with desired plasmid allowed investigation of the role of NK cells in CD47 blockade and opsonization via sCV1-hIgG1. In a preceding experiment, the lysis activity of the NK cell line against pNLuc-transfected 231/LM2-4 cells was evaluated at increasing E:T ratios (Figure S8). NK cell-mediated lysis at E:T ratios of 1:1 and 2:1 was studied to investigate the role of NK cells in CD47 blockade and opsonization with sCV1-hIgG1 (Figure 8). Although psCV1 or psIgG1 transfection did not induce any cell lysis above the level of transfection control (pmCherry), psCV1-hIgG1-transfected 231/LM2-4 cells were efficiently lysed at E:T ratios of 2:1 and 1:1.

**DISCUSSION**
Our approach here was to combine “the best of both worlds,” i.e., maximized binding affinity for human CD47 combined with full CDC and ADCC activity. Although this would be almost impossible when applying systemic protein therapy, we developed a gene therapy approach for tumor-directed transgene expression and secretion of the therapeutic protein directly by the tumor cells. We evaluated sCV1-hIgG1 expression, secretion, and binding to CD47, as well as its bystander effect in vitro, and confirmed the efficacy of this therapeutic plasmid in vivo.

In our plasmid design, we utilized the CpG-free shuffled CMV-EF1 promoter (SCEP), which combines elements of the strong constitutive cytomegalovirus (CMV) promoter and the human elongation factor-1-alpha promoter and, when used in combination with the hCMV enhancer element within a CpG-free plasmid backbone, enables high and sustained transgene expression levels. The high-affinity CV1 consensus sequence based on human SIRPα1 was fused with an Fc variant based on human IgG1 carrying a point mutation (E333A), which boosts both ADCC and CDC and exhibits increased binding affinity to Fc gamma receptor (FcγR)IIIa. Secretion was ensured with a 20-amino acid secretion signal sequence derived from human IL-2 with intracellular cleavage of the signal after serine at position 20. As a transfection reagent we utilized LPEI with a molecular weight (MW) of 10 kDa. This polymer allows the formation of nanosized, positively charged polyplexes, which enable cellular uptake by adsorptive endocytosis and promote endosomal release of the payload by a proton sponge mechanism. Both SCEP promoter and LPEI polyplex result in non-selective transfection and transgene expression. Here, tumor-specific delivery and tumor cell-specific transgene expression can be achieved, for example, by de-targeting approaches using microRNA (miRNA)-binding sites, which restrict transgene expression to cells lacking otherwise ubiquitously expressed miRNAs, e.g., miRNA 143, 16. Similarly, the polyplex approach could be improved by shielding the positive charge with polyethylene glycol and attaching a cell-binding ligand, e.g., an EGFR-binding peptide, which promotes selective uptake of polyplexes in EGFR-overexpressing tumor cells. Prior to transfection studies, all cell lines were first
screened for CD47 expression (Figure S2). Interestingly, the MDA-
MB-231-derived subline 231/LM2-4, which was initially isolated
from a lymph node metastasis after orthotopic tumor implantation
into SCID mice,23 did not show any further elevated CD47 expression
levels compared to the cell line it was derived from. For both A549
and MDA-MB-231, elevated CD47 levels have been reported and
are considered to be a driver for metastasis for the respective tumor
type (non-small cell lung cancer [NSCLC], A549 cells; triple-negative
breast cancer, MDA-MB-231 cells).23,24 After transfection, CD47
blockage was determined with the anti-CD47 antibody clone
blocks its interaction with CD47, 25 and the presence of sCV1-
expressing the transgene (as shown by mCherry as reporter), points to
the efficient secretion of the protein due to the IL-2 secretion sequence26
and a potent bystander effect, when treating untransfected
cells with transfection supernatant (Figure 3C). We also estimated the
total amount of fusion protein secreted over time (Figure 3B). The
fusion protein has a MW of 38.5 kDa (calculated at https://web.expasy.org/compute_pi/), with
the monomeric CD47-binding SIRPz variant CV1 (13 kDa)25 fused to the IgG1 Fc region
comprising the CH2 and CH3 domains of the IgG heavy chain and the hinge region
(25.5 kDa). Because of the cysteine residues in the hinge region, the protein dimerizes and ex-
hibits a final MW of 77 kDa. In relation to the concentration of IgG1 (~150 kDa), this would
correspond to a concentration of 0.12 μg/mL.

It should be noted that this amount of secreted scV1-hlgG1 is in
addition to the amount already bound to CD47 on cells expressing
the fusion protein; thus the total amount can be expected to be
considerably higher. Nevertheless, compared to protein therapy,
this amount appears rather low. In a preclinical setup, NSG mice xen-
ografted with CD47-positive tumors were treated intravenously with
a daily dose of 7.5 mg/kg CV1-hIgG4.13 This corresponds to an
average blood concentration (assuming 8% blood content) of
~90 μg/mL (i.e., >1 mM). With the therapeutic anti-CD47 antibody
Hu5F9-G4, plasma levels of ~1 mg/mL were achieved with dosages
ranging from 20 to 45 mg/kg.2 Antibody accumulation in solid tu-
mors is typically rather low, ranging from 0.001% to 0.01% of the in-
jected dose per gram of tumor. Thurber and Weissleder developed a
theoretical model to estimate absolute antibody concentration. Even
at saturating concentrations, where >5,000 mg would be needed for
a 70-kg patient to target a highly expressed antigen, intratumoral an-
tigen concentration could be as low as 100 nM.27 Even then, the most
rate-limiting step is extravasation and diffusion within the tissue.
Hence, we assume that a relatively low dose of fusion protein contin-
uously expressed in the tumor could in principle be sufficient for a

Figure 5. psCV1-hlgG1 ex vivo transfected tumor cells show significantly reduced growth in vivo
(A and B) CD47 expression in a 231/LM2-4-EGFPLuc lymph node metastasis 34 days after orthotopic implan-
tation with (A) huCD47 staining and (B) isotype control staining; 4x magnification images with zoomed-in 40x
magnification images. Scale bars: 200 μm (4x magnification) and 20 μm (40x magnification). (C–G) 231/LM2-4-
EGFPLuc were transfected ex vivo with the indicated plasmid and 48 h thereafter orthotopically implanted into the
4th pair of inguinal ripples (2 sites/animal) in female CB17-SCID mice. Two experiments [Exp I (C and E) and
Exp II (D, F, G)] were conducted. Exp I was terminated 34 days after implantation and Exp II after 25 days. 3 ani-
mals/group were used, and the BLI signal at each injection site was quantified within a separate ROI (i.e., n = 6 ±SD per group). (C and D) Tumor growth followed by BLI. (E and F) Size of explanted tumors after termination. (G) Repre-
sentative BLI pictures (color coded, overlaid onto a re-
flected light picture) of animals in Exp II 18 days after im-
plantation. (C, E, and G): n = 6 per experiment, **p ≤ 0.01, *p
≤ 0.05; psCV1-hlgG1 transfected versus pmCherry
transfected; U test (Mann-Whitney). Scale bars in E and F: 2 cm.
potent antitumoral effect while at the same time preventing side effects in other organs. Although protein therapy relies on repeated dosing, local expression can be a continuous source of therapeutic protein. Evaluating both early and late time points after transfection for CD47 blocking activity and Fc detection (Figure 4; Figure S6), we observed both a fast onset (detectability of the Fc part 6 h after transfection) and prolonged duration (12 days and three passages after transfection; Figure 4) of expression. Therefore, one could at least estimate that a single plasmid transfection could last for >2 weeks, as experiments with electroporation of human tumors also revealed that transgene expression is found for >3 weeks.17

For in vivo evaluation, we have selected a xenograft model in CB17-SCID mice. With the EGFP-Luc labeled, orthotopically implanted 231/LM2-4-EGFPLuc cells we observed a fast onset of tumor growth (Figure 5), which is in line with observations by Munoz et al., where tumors of size >300 mm³ developed within 3 weeks and showed metastases in various areas, including lymph nodes and mesenterium.22 Similarly, we could demonstrate axillary lymph node infiltration being positive for CD47 (Figure 5A). In contrast, the parental cell line MDA-MB-231, also when labeled with EGFPLuc and implanted orthotopically, shows a very slow onset of tumor growth and can be followed via BLI for several months (M. Ogris et al., unpublished observations). In both experiments, psCV1-hIgG1-transfected tumors exhibited a rapidly decreasing BLI signal after implantation. Considering the BLI sensitivity, loss of BLI signal and absence of 4 out of 12 tumors after explantation also points to a complete tumor eradication, as with a similar BLI imaging setup even single-digit numbers of luciferase-expressing cells can be detected.28 Here, caliper-based size measurements would not be applicable to measure the presence of tumor cells. Also, BLI only detects metabolically active tumor tissue without considering a necrotic core. Still, pmCherry-transfected tumors appeared smaller then untreated tumors, which was significant in Exp I when measuring the size of explanted tumors (Figure S7). It has been reported that polycations like LPEI or cationic dendrimers can exert an intrinsic effect on delayed tumor cell growth, as observed by us (M. Ogris et al., unpublished observations) and others.29 In Exp II, psCV1-transfected tumors gave BLI signals similar to pmCherry-transfected tumors and were only insignificantly smaller. Apparently the CD47 block alone is insufficient for tumor growth retardation, which is in accordance with the findings of Weisskopf et al.: while applying the CD47-blocking protein CV1 monomer alone, no reduction of tumor growth in vivo was observed. Only with co-application of rituximab (in an EGFR-positive xenograft model), tumor growth was reduced and led to complete tumor eradication and long-term survival (>250 days) in 25% of treated animals.31 Thus, we conclude that also with a locally produced CD47-blocking protein, a functional Fc part for potential macrophage and/or NK cell activation is quite crucial and needed.

In principle, a syngeneic tumor model would also be applicable, as we also demonstrated sCV1-hIgG1 binding to murine CD47 on CT26 cells (Figure S5). This is in line with previous results, where ALX148, a fusion protein based on a high-affinity SIRPα version, acts antitumorally in combination with anti PD-L1 on implanted CT26 tumors in mice.14 Although murine SIRPα (from C57BL/6 mice) binds with similar affinity (kD ~5 μM) to human and murine CD47, a mutation in the SIRPα sequence in non-obese diabetic (NOD)/SCID mice exhibits an exceptional affinity for human CD47 with a kD of 0.08 μM.30 Interestingly, human SIRPα binds human CD47 with a kD of 0.3–0.6 μM, i.e., with a >8-fold higher affinity compared to mice. This points to a potentially higher impact of the SIRPα-CD47 axis in human physiology and oncology compared to mice and makes it necessary to use high-affinity CD47 binders to unleash the full therapeutic potential. Although in vitro transfected cells were used for proof of principle, local or systemic application of gene vectors is inevitable in a therapeutic setting. Also, intratumoral application of plasmid DNA is a vital option for the treatment of unresectable tumors or as a neoadjuvant treatment. For example, image-guided injections of LPEI polyplexes were used for the treatment of pancreatic ductal adenocarcinoma,21 where, in the case of larger tumors, multiple injections can also be necessary and are clinically feasible.17,31 Twenty-four hours after intratumoral transfection, we observed local blockage of CD47 surrounding the injection site in the millimeter range as shown by immunohistochemistry (Figure 6). This localized CD47 blockage can be explained by several means. First, we expect sCV1-hIgG1 to be quickly absorbed by transfected and adjacent tumor cells, as they express high amounts of CD47 (Figure 5A), limiting a further spread. Second, this can also be due to limited total amount of sCV1-hIgG1 fusion proteins produced by tumor cells after transfection, as our in vitro measurement on transfected 231/LM2-4 cells gave a value in the single nanomolar range.
We did not distinguish between different subsets of CD47-blocking antibody applied in an osteosarcoma xenograft literature, as recruitment of macrophages and their in vivo role of macrophages after sCV1-hlgG1-mediated CD47 blocking and observed significant macrophage infiltration at 17 h after transfection (Figure 7). This would be in line with the literature, as recruitment of macrophages and their infiltration into tumor tissue appears rather fast after CD47 blockade: with a CD47-blocking antibody applied in an osteosarcoma xenograft model, significant macrophage infiltration was measurable 24 h after treatment. We did not distinguish between different subsets of macrophages, as we only stained for the pan-macrophage marker F4/80. Lin and colleagues demonstrated that CD47 blockade ex vivo with TTI-621 (a fusion protein of an engineered SIRPα protein and human IgG1-Fc) on lymphoma cells triggered their phagocytosis by a broad subset of macrophages, including both the M1 and the M2 subtype. Nevertheless, Mohanty and colleagues observed mainly M1 macrophage (F4/80+/CD80+) and not M2 macrophage (F4/80+/CD260+) infiltration after CD47 blockade in vivo. Still, as the quantitative macrophage infiltration was not as long-lasting as one could expect from the sustained reduction in tumor growth, other effects or immune cells might also be involved. Hayes and colleagues demonstrated that blocking the cis-interaction between CD47 and SIRPα both expressed on macrophages plus blockage of CD47 on target cells leads to a kind of hyper-phagocytosis, which could explain the prolonged antitumoral effect of macrophages due to higher activity. Neutrophils were found to destruct antibody-opsonized cancer cells not by ADCC but rather by trogocytosis, and blocking the CD47-SIRPα interaction further potentiates this activity. Although CB17-SCID mice lack functional T and B cells, NK cells expressing SIRPα and CD47 are present. Blocking CD47 on NK cells has been demonstrated to increase their antitumoral activity in a syngeneic melanoma model, while blocking CD47 on leukemia cells or major histocompatibility complex (MHC)-I-deficient cells increased the killing capacity of NK cells. As we could not demonstrate NK staining in vivo, we utilized an in vitro assay with a genetically modified NK cell line (NK92) overexpressing a polymorphic variant of FcγRIIIa/CD16a, which exhibits the most potent ADCC effect. For optimal sensitivity, target cells were transfected with the more stable luciferase NanoLuc (nLuc), which gives a bright bioluminescence signal. Lysis activity peaked at E:T ratio 5:1 and 10:1 with 50% lysis (Figure S8). This is in line with observations by Shenouda et al., where the parental cell line of 231/LM2-4, namely MDA-MB-231, was efficiently lysed at E:T ratios of 5:1 and above by activated NK cells isolated from healthy donors as well as from breast cancer patients. Only psCV1-hlgG1-transfected cells were lysed to a significant extent by NK cells (Figure 8). Apparently, a (partial) block of CD47 after transfection with psCV1 (see Figure 2B) is not sufficient to induce NK-cell mediated lysis, and it needs the Fc part bound to the cell surface to trigger FcγRIII binding and subsequent NK cell activation. In our experimental setup, CD47 blockade was occurring on the target cells, but due to the secretion of sCV1-hlgG1, some CD47 blockage on NK cells is possible. Hence, an additional beneficial effect of sCV1-hlgG1 secretion could occur, further increasing antitumoral activity of NK cells.
In conclusion, it can be stated that within a cancer gene therapy approach the tumor-restricted expression of a secreted, highly effective CD47-blocking protein in conjunction with an Fc part for maximized ADCC and CDC can result in a potent antitumoral response concomitant with activation of both macrophages and NK cells. Shortcomings due to low transfection rates could be easily overcome by the potent bystander effect. Recently, Huang and colleagues published a related gene therapy approach but utilizing an oncolytic adenoviral vector. Here, the SIRPζ-Fc fusion protein was not combined with an IL-2 secretion signal; hence the system relies on the necessity of uncontrolled cell burst and fusion protein release. Another advantage of our approach is the combination of maximal CD47 blockade and immune activation: Although the CV1 variant of SIRPζ binds human CD47 with an affinity of 11 pM (i.e., 50,00-fold higher affinity compared to wild-type human SIRPζ), it could not be used as a systemically applied IgG1-Fc fusion protein because of exuberated toxicity. In contrast, the therapeutic fusion protein TTI-621 shows a lower affinity for CD47 (and preferentially binds clustered CD47, hence sparing red blood cells) but comes with a highly active human IgG1-Fc part. Therapeutic antibodies are currently evaluated in clinical trials (Hu5F9-G4 and SRF231) and have a high CD47 affinity (8 nM for monomeric and 12 pM for dimeric CD47 in the case of Hu5F9-G4) but are of a human IgG4 isotype, which exhibits a 10-fold reduced affinity for human FcγRI and weak ADCC and CDC activity. Currently, Hu5F9-G4 is now also successfully combined with rituximab (anti CD20, IgG1 type) in lymphoma treatment. Our approach is currently designed for localized, intratumoral application. Its use within a systemically applied, targeted gene delivery system would be the next logical step, where specificity of delivery (e.g., using cell-targeting ligands) and/or specificity of expression (using tumor-specific promoters or miRNA-based de-targeting approaches) is further pursued. For preclinical studies, syngeneic tumor models need to be evaluated (e.g., CT26 colon cancer in BALB/c mice). This is important as the adaptive immune system, especially CD8+ cytotoxic T cells, is also involved in tumor eradication after CD47 blockage. Taken together, this gene-based approach can be seen as a platform that should enable blocking of other checkpoint inhibitors with high-affinity and -efficacy fusion proteins, avoiding side effects on non-tumorous tissue.

**MATERIALS AND METHODS**

**Cells**

MDA-MB-231 (HTB-26, ATCC, Manassas, VA, USA), 231/LM2-4, kindly provided by R.S. Kerbel, Sunnybrook and Women’s College Health Sciences Centre S-217, Toronto, ON, Canada, and CT26 (CRL-2638, ATCC) were maintained in DMEM, high glucose (Sigma-Aldrich, Vienna, Austria) and A549 (CCL-185, ATCC) in RPMI-1640 (Sigma-Aldrich) with 1% penicillin-streptomycin, and 10% fetal bovine serum (FBS; Biowest). CT26 cells were co-transfected with plasmid pNluc (40 μg) and the indicated plasmid (psCV1-hlgG1, psCV1, pshlgG1, pmCherry; 40 μg) in T-75 flasks for 48 h, transferred to 48-well plates, and 24 h thereafter co-cultured with NK92 cells at the indicated E:T ratio for 4 h. Cells treated with medium containing 0.1% Triton X-100 served as positive control (100%) and cells without co-culture as negative control (0%). Nluc activity was quantified in the supernatant. n = 9 + SD. ****p < 0.0001, ns p > 0.05; U test (Mann-Whitney).

**Plasmids and cloning**

To generate plasmid psCV1-hlgG1, the sequence IL2-CV1-hlgG1 (1,331 bp) was designed by placing the CV1 sequence into the multiple cloning site (MCS) of plasmid pFUSE-hlgG1e4-Fc2 (Invivogen, Toulouse, France) situated between the IL-2 secretion signal sequence (s) and the hlgG1e4-Fc sequence (hlgG1). The full sequence IL2-CV1-hlgG1 containing the IL-2 secretion signal sequence, the CV1 sequence, and the hlgG1e4-Fc sequence (hlgG1) was further codon optimized for human expression and to avoid certain cis-acting sequence motifs (e.g., internal TATA boxes, RNA instability motifs, and cryptic splice donor and acceptor sites). Synthesized, sequenced and cloned into plasmid pMK-RQ by GeneArt (Thermo Fisher, Regensburg, Germany). Furthermore, the IL2-CV1-hlgG1 sequence...
was cloned into pCPg-hCMV/SCEP-mCherry using XbaI and NheI replacing the mCherry cDNA. Plasmid pscCV1 was generated by excising the IL2-CV1 sequence from psCV1-hIgG1 with BglIII and replacing the mCherry cDNA in pCpG-hCMV/SCEP-mCherry. Plasmid pshIgG1 was generated by excising the CV1 sequence from psCV1-hIgG1 with Eco32I and MbiI. pCpG-hCMV/SCEP-mCherry (for short: pmCherry) was used without further modification. All plasmids were propagated in E. coli DH5a with kanamycin.

### Polypelex generation

Polypelexes were generated with 10-kDa LPEI at an N/P ratio (molar ratio nitrogen in LPEI versus phosphate in pDNA) of 9 in principle as described previously. Equimolar volumes of a pDNA-LPEI-containing solution were mixed by flash pipetting and incubated for 5 min at room temperature. For *in vitro* experiments, nanoparticles were generated at a final pDNA concentration of 20 μg/ml (of the desired plasmid) in HBS buffer (HEPES-buffered saline, 20 mM HEPES pH 7.4, 145 mM NaCl) and for *in vivo* experiments at 200 μg/ml in HBG buffer (HEPES-buffered glucose, 20 mM HEPES pH 7.4, 5% glucose w/v). Particle size and ζ-potential were measured on a Malvern NS500 system (Malvern, UK) as described previously.

### In vitro transfection and bystander effect

3.95 × 10⁶ cells were seeded into T-75 flasks (Sarstedt, Nümbrecht, Germany) 24 h prior to treatment. After removal of medium, polypelexes (40 μg pDNA/lask) were added in 15 ml of basal medium (DMEM, high glucose, serum free) for 4 h, and thereafter 15 ml of complete medium (DMEM, high glucose, 1% L-glutamine, 1% penicillin-streptomycin, 10% FBS) was added. Different polypelex treatment groups were performed depending on the desired plasmid needed for transfection, namely pscCV1-hIgG1 (therapeutic plasmid), pshIgG1 (control plasmid; for secreted IgG1-Fc), pscCV1 (control plasmid; for secreted SIRPα), and pmCherry (transfection control plasmid). Transfection supernatant was stored frozen at −80°C until further use. Bystander effect was determined by transferring 30 ml of freshly thawed supernatant (or diluted with complete medium to the indicated ratio) from 48-h-transfected cells to similarly seeded but untransfected cells. These untransfected cells were incubated with supernatant for 1 h at 37°C, and thereafter cells were further processed for analysis. To determine transgene expression over a duration of 12 days, 1.32 × 10⁸ cells were seeded in T-25 flasks, transfected with 13.2 μg plasmid/lask for 48 h (p0), and thereafter further passaged at a ratio of 1:8 after 5 (p1), 9 (p2), and 12 (p3) days into new T-25 flasks with fresh complete medium. Remaining cells were analyzed accordingly for transgene expression.

### Flow cytometry

Cells were detached with TrypLE (A549) or Versene (MDA-MB-231, 231/LM2–4) for 4 min at 37°C, neutralized with complete medium, and washed with phosphate-buffered saline (PBS). Cell count and viability were measured by a flow cytometer (MACSQuant Analyzer 10, Miltenyi Biotec, Bergisch Gladbach, Germany) by adding 1 μg/ml propidium iodide (PI). 150,000 live cells were transferred into 96 V-bottom plates (FM9561, Greiner CELLSAR, Greiner Bio-One, Kremsmünster, Austria). All antibodies were diluted in 0.5% (w/v) bovine serum albumin (BSA) in PBS (pH 7.4) and indicated amounts in 20 μl dilution added per 150,000 cells. For huCD47 staining, 0.03 μg of mouse-anti-huCD47 (ab3283 clone B6H12.2, Abcam, Cambridge, UK) or purified mouse IgG1, κ isotype control (Cat 557273, BD Biosciences) was added for 30 min at 4°C, followed by three washing steps with PBS and 1-h incubation with 1:2,000 diluted goat anti-mouse IgG H&L (Alexa Fluor 647 [AF647] conjugated, preadsorbed; ab150119, Abcam). For Fc detection, cells were incubated with 1:2,000 diluted protein A-AF488 conjugate (P11047, Thermo Fisher Scientific, Vienna, Austria), 1:500 diluted goat anti-human IgG Fc (DyLight 650 conjugated, preadsorbed; ab98622, Abcam), or 1:1,000 goat IgG isotype control AF647 conjugate (Bs_0294P-A647, Bioss Antibodies, Worcester, MA, USA) for 1 h at 4°C, washed three times with PBS, and diluted in 0.5% BSA-2 mM EDTA-PBS for the measurement performed on a MACSQuant Analyzer 10 Flow Cytometer (Miltonyi Biotec). Live/dead cell staining was performed with 4,6-diamino-2-phenylindol (DAPI) (1 μg/ml) and cells gated for live cells. Data analyses were done with FlowJo software (version 1.10).

### Dot blot analysis

Transfection supernatant (2 μl/dot) was dropped onto a nitrocellulose membrane (sc-201706, 0.45 μm; Santa Cruz Biotechnology, Heidelberg, Germany) and air-dried; a standard curve was generated similarly with human IgG1 isotype (ALX-804-133-C100; Enzo Biochem, Farmingdale, NY, USA). BSA (Sigma-Aldrich; 2 mg/ml) and complete medium were dotted as negative controls. The membrane was washed twice with TBS-T (Tris-buffered saline with 1% [w/v] Tween 20) and incubated for 1 h with anti-human IgG Fc-horseradish peroxidase (HRP) (1:10,000 dilution in TBS-T; ab98624, Abcam) at room temperature. After three washing steps with TBS-T, the blot was incubated for 1 min with enhanced chemiluminescence (ECL) substrate (50 μl luminol, 22 μl p-coumaric acid, and 3 μl H₂O₂ [33%] in 10 ml 100 mM Tris pH 8.5) and chemiluminescence was recorded on a Molecular Imager Gel DocTM XR System (Bio-Rad, Vienna, Austria). The integrated signal density was evaluated via ImageJ (Image) 1.53a and data evaluation performed with Microsoft Excel (version 2013) and GraphPad Prism (version 6.0).

### Confocal laser scanning microscopy

25,000 cells/well were seeded into chambered slides (Nunc Lab-Tek II 8-well slides; Thermo Fisher Scientific) 72 h prior to treatment and then washed with PBS, and 400 μl of 48-h transfection supernatant (1:2 dilution in complete medium) was added for 1 h at 37°C. After PBS washing (3 times) cells were fixed with formaldehyde (4% w/v,
in HBS) for 40 min, PBS washed (3 times), and stained for 1 h with goat anti-human IgG Fc (DyLight 650, preadsorbed; ab98622) diluted 1:200 in PBS + 0.5% BSA. After PBS washing and DAPI staining (1 µg/mL), cells were imaged on a Leica TCS SPE microscope (Leica, Wetzlar, Germany) with a 40 × oil immersion objective with laser excitation (405 nm for DAPI, 635 nm for DyLight 650) and appropriate emission filters. Z-scans with a vertical resolution of 0.1 µm were conducted and central sections shown. Data acquisition and analysis was done with LAS X software (version 3.1.2.16221).

**NK cell co-culture cell lysis assay**

3.95 × 10⁶ 231/LM2-4 cells were seeded in T-75 flasks and 24 h thereafter co-transfected with polyplexes containing 40 µg of pNLuc and 40 µg of the indicated plasmid (psCV1-hIgG1, psCV1, or psIgG1 or pmCherry) for 4 h in basal medium; thereafter, complete medium was added and after a total of 48 h cells were harvested and transferred to 48-well plates (Greiner Bio-One; 100,000 cells/well). After a further 24 h of growth, medium was removed and NK92 cells were suspended in 200 µL/well serum-reduced medium at E:T ratios of 10:1, 5:1, 2:1, and 1:1 or medium only (w/o co-culture) added for 4 h at 37°C. The supernatant was harvested and centrifuged for 5 min at 300 × g, and nLuc luciferase released into supernatant was measured in white light (nLuc (w/o coculture)).

**In vivo studies**

Female CB17-SCID mice (CB-17/Icr-Prkdcsid/scid/Rj; Janvier Labs, Le Genest-Saint-Isle, France) were kept in individually ventilated type 2 cages with food and water provided ad libitum. All animal procedures were approved by local ethics committee and are in accordance with the Austrian law for the protection of animals and EU Directive 2010/63/EU. 1 × 10⁶/30 µL of 48-h-transfected 231/LM2-4-EGFP-Luc cells resuspended in sterile PBS was percutaneously implanted into the 4th pair of inguinal nipples (two implantation sites), and tumor growth was evaluated by BLI as described below. In a separate experiment, intratumoral injection into the middle of the untransfected tumor core (two puncture sites per tumor) was performed on day 13 after implantation (50 µL polyplex solution, 200 µg/mL pDNA) with a 26 g needle and a 2-mL syringe (Braun, Melsungen, Germany). BLI signal was detected after injection of d-luciferin (potassium salt; Intrase Medical, Lausanne, Switzerland; 30 mg/mL in DPBS; dosage: 120 mg/kg) with an IVIS Spectrum-CT imaging system (PerkinElmer) in principle as described previously²⁰ and quantified by placing appropriate regions of interest (ROIs) onto tumor sites.

**Immunohistochemistry**

Tissues were fixed for 22 h in 4% formaldehyde in HBS and further processed as described previously.²⁰ For CD47 staining, sections were pre-warmed for 2 h at 55°C, deparaffinized, and dehydrated, and heat-induced epitope retrieval (HIER) was performed for 30 min in Tris-EDTA buffer (10 mM Tris, 1 mM EDTA, pH 9). Slides were treated with normal goat serum (30 min) and avidin/biotin blocking reagent (BUF016, Bio-Rad) for 15 min each, followed by incubation with rabbit-anti-HuCD47 antibody [EPR21794] (ab218810, Abcam; 1:2,000) or Rabbit IgG, monoclonal [EPR25A]-Isotype Control (ab172730, Abcam) in PBS + 2% BSA overnight at 4°C. Endogenous peroxidase was blocked with 0.3% H₂O₂-PBS for 15 min and signals detected via Vectastain ABC HRP Kit (Peroxidase, Rabbit IgG, PK-4001; Vector Laboratories, Burlingame, CA, USA). For F4/80 staining, slides were incubated in 0.3% H₂O₂-PBS for 10 min, heated in DAKO antigen retrieval solution (pH 6.0, #1699, Agilent, Santa Clara, CA, USA) for 1 h, and blocked for 5 min with 10% goat serum-0.1% Tween 20 in PBS. Incubation with rat anti-mouse F4/80 (#123102, BioLegend, San Diego, CA, USA) diluted 1:25 in 1% goat serum-PBS overnight at 4°C was followed by labeling with biotinylated goat anti-rabbit antibody (#BA9401, Vector Laboratories) and staining with the Vectastain Elite ABC Kit (#PK6100, Vector Laboratories) (30 min each) and staining with 3,3′-diaminobenzidine (DAB). Slides were counterstained with hematoxylin and mounted in Entellan new. Numbers of F4/80-positive cells/field were evaluated by Definiens analyses (Definiens, Munich, Germany). Images were acquired on an Olympus BX53 microscope.

**Statistical analysis**

Statistical analysis was performed with GraphPad Prism 6 and an online tool (https://www.socscistatistics.com/). For the comparison of two treatment groups a U test (Mann-Whitney) was performed. Every effort was made to keep testing consistent across related experiments.

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.omto.2021.09.005.

**ACKNOWLEDGMENTS**

This work was in part co-funded by a grant from the Austrian Research Promotion Agency (FFG).

**AUTHOR CONTRIBUTIONS**

M. Billerhart: investigation, methodology, formal analysis, writing – original draft, writing – review & editing; M.S.: formal analysis, investigation; H.S.: formal analysis, investigation, methodology; W.P.: formal analysis, investigation; J.P.: formal analysis, investigation; S.D.: formal analysis, investigation; A.T.: resources; J.M.: resources; W.P.: formal analysis, investigation; M. Blaschek: formal analysis, investigation; P.H.: methodology, supervision, resources, writing – review & editing; H.S.: conceptualization, formal analysis, investigation, methodology, supervision, validation, writing – original draft, writing – review & editing; M.O.: conceptualization, funding acquisition, writing – review & editing, project administration, supervision.
DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES

1. Swann, J.B., and Smyth, M.J. (2007). Immune surveillance of tumors. J. Clin. Invest. 117, 1137–1146.
2. Li, M.O., Wan, Y.Y., Sanjabi, S., Robertson, A.K., and Flavell, R.A. (2006). Transforming growth factor-beta regulation of immune responses. Annu. Rev. Immunol. 24, 99–166.
3. Ricklefs, F.L., Ayalo, Q., Krenzin, H., Mahmoud, A.B., Speranza, M.C., Nakashima, H., Hayes, J.L., Lee, K., Balaj, L., Passaro, C., et al. (2018). Immune evasion mediated by PD-L1 on glioblastoma-derived extracellular vesicles. Sci. Adv. 4, eaar2766.
4. Barclay, A.N., and Van den Berg, T.K. (2014). The interaction between signal regulatory protein alpha (SIRPa) and CD47: structure, function, and therapeutic target. Annu. Rev. Immunol. 32, 25–50.
5. Sikic, B.I., Lakhani, N., Patnaik, A., Shah, S.A., Chandana, S.R., Rasco, D., Colevas, A.D., O’Rourke, T., Narayan, S., Papadopoulos, K., et al. (2019). First-in-Human, First-in-Class Phase I Trial of the Anti-CD47 Antibody Hu5SF9-G4 in Patients With Advanced Cancers. J. Clin. Oncol. 37, 946–953.
6. Cioffi, R., Trabulo, S., Hidalgo, M., Costello, E., Greenhalf, W., Erkan, M., Kleeff, J., Sainz, B., Jr., and Heeschen, C. (2015). Inhibition of CD47 Effectively Targets Pancreatic Cancer Stem Cells via Dual Mechanisms. Clin. Cancer Res. 21, 2325–2337.
7. Garcia-FonciLlas, J., Sunakawa, Y., Aderka, D., Wainberg, Z., Ronga, P., Witzler, P., and Stintzing, S. (2019). Distinguishing Features of Cetuximab and Panitumumab in ColoRectal Cancer and Other Solid Tumors. Front. Oncol. 9, 849.
8. Peluso, M.O., Adam, A., Armet, C.M., Zhang, L., O’Connor, R.W., Lee, B.H., Lake, A.C., Normant, E., Chappel, S.C., Hill, J.A., et al. (2020). The Fully human anti-CD47 antibody SRF231 exerts dual-mechanism antitumor activity via engagement of the activating receptor CD32a. J. Immunother. Cancer 8, e004134.
9. Petrova, P.S., Viller, N.N., Wong, M., Pang, X., Lin, G.H., Dodge, K., Chai, V., Chen, H., Lee, V., House, V., et al. (2017). TTI-621 (SIRPaFc): A CD47-Blocking Inactive Immune Checkpoint Inhibitor with Broad, Antitumor Activity and Minimal Erythrocyte Binding. Clin. Cancer Res. 23, 1068–1079.
10. Lou, J., Wang, L., Zhao, F., Tseng, S., Narayan, C., Shura, L., Willingham, S., Howard, M., Prohaska, S., Volkmer, J., et al. (2015). Pre-Clinical Development of a Humanized Anti-CD47 Antibody with Anti-Cancer Therapeutic Potential. PLoS ONE 10, e0137345.
11. Rodriguez, P.L., Harada, T., Christian, D.A., Pantano, D.A., Tsai, R.K., and Discher, D.E. (2013). Minimal "Self" peptides that inhibit phagocytic clearance and enhance delivery of nanoparticles. Science 339, 971–975.
12. Jalil, A.R., Andrechak, J.C., and Discher, D.E. (2020). Macrophage checkpoint blockage: results from initial clinical trials, binding analyses, and CD47-SIRPa structure-function. Antib. Ther. 3, 80–94.
13. Weiiskopf, K., Ring, A.M., Ho, C.C., Volkmer, J.P., Levin, A.M., Volkmer, A.K., Ozkan, E., Fernthoff, N.B., van de Rijn, M., Weissman, I.L., and Garcia, K.C. (2013). Engineered SIRPa variants as immunotherapeutic adjuvants to anticancer therapies. Science 341, 88–91.
14. Kauder, S.E., Kuo, T.C., Harrabi, O., Chen, A., Sanghaling, E., Doyle, L., Roche, S.S., Bollini, S., Han, B., Sim, J., et al. (2018). ALX148 blocks CD47 and enhances innate and adaptive antitumor immunity with a favorable safety profile. PLoS ONE 13, e0201832.
15. Johnson, L.D.S., Banerjee, S., Kruglov, O., Viller, N.N., Horwitz, S.M., Lesokhin, A., Zain, J., Querfeld, C., Chen, R., Okada, C., et al. (2019). Targeting CD47 in Sézary syndrome with SIRPaFc. Blood Adv. 3, 1145–1153.
16. Su, B., Cengizzeroğlu, A., Farkasova, K., Viola, J.R., Anton, M., Ellwartz, J.W., Haase, R., Wagner, E., and Ogris, M. (2013). Systemic TNFα gene therapy synergizes with liposomal doxorubicin in the treatment of metastatic cancer. Mol. Ther. 21, 300–308.
17. Bhatia, S., Longino, N.V., Miller, N.J., Kulikauskas, R., Iyer, J.G., Ibrani, D., Blom, A., Byrd, D.R., Parvathaneni, U., Twitty, C.G., et al. (2020). Intratumoral Delivery of Plasmid IL-12 Via Electroporation Leads to Regression of Injected and Noninjected Tumors in Merkel Cell Carcinoma. Clin. Cancer Res. 26, 598–607.
18. Kopp, F., Schnoedd, M., Haase, R., Wagner, E., Rosid, A., and Ogris, M. (2013). De-targeting by miR-143 decreases unwanted transgene expression in non-tumorigenic cells. Gene Ther. 20, 1104–1109.
19. Taschauer, A., Geyer, A., Gehrig, S., Maier, J., Sami, H., and Ogris, M. (2016). Up-Scaled Synthesis and Characterization of Nonviral Gene Delivery Particles for Transient In Vitro and In Vivo Transgene Expression. Hum. Gene Ther. Methods 27, 87–97.
20. Magnusson, T., Haase, R., Schlie, M., Wagner, E., and Ogris, M. (2011). Sustained, high transgene expression in liver with plasmid vectors using optimized promoter-enhancer combinations. J. Gene Med. 13, 382–391.
21. Subramanian, S., Shin, J.W., and Discher, D.E. (2020). Macrophages show higher levels of engulfment after disruption of cis interactions between CD47 and the checkpoint receptor SIRPa. J. Cell Sci. 133, jcs237800.
37. Mathung, H.L., Babes, L., Zhao, X.W., van Houdt, M., Treffers, L.W., van Rees, D.J., Franke, K., Schornagel, K., Verkuijlen, P., Janssen, H., et al. (2018). Neutrophils Kill Antibody-Opsonized Cancer Cells by Trogoptosis. Cell Rep. 23, 3946–3959.e6.

38. Nath, P.R., Pal-Nath, D., Mandal, A., Cam, M.C., Schwartz, A.L., and Roberts, D.D. (2019). Natural Killer Cell Recruitment and Activation Are Regulated by CD47 Expression in the Tumor Microenvironment. Cancer Immunol. Res. 7, 1547–1561.

39. Deuse, T., Hu, X., Agbor-Enoh, S., Jang, M.K., Alawi, M., Saygi, C., Gravina, A., Tediashvili, G., Nguyen, V.Q., Liu, Y., et al. (2021). The SIRPa:CD47 immune checkpoint in NK cells. J. Exp. Med. 218, e20200839.

40. Taylor, R.J., Chan, S.L., Wood, A., Voskens, C.J., Wolf, J.S., Lin, W., Chapoval, A., Schulze, D.H., Tian, G., and Strome, S.E. (2009). FcgammaRIIIa polymorphisms and cetuximab induced cytotoxicity in squamous cell carcinoma of the head and neck. Cancer Immunol. Immunother. 58, 997–1006.

41. Hayek, S., Bekaddour, N., Besson, L., Alves de Sousa, R., Pietrancosta, N., Viel, S., Smith, N., Jacob, Y., Nisole, S., Mandal, R., et al. (2019). Identification of Primary Natural Killer Cell Modulators by Chemical Library Screening with a Luciferase-Based Functional Assay. SLAS Discov. 24, 25–37.

42. Shenouda, M.M., Gillgrass, A., Nham, T., Hogg, R., Lee, A.J., Chew, M.Y., Shafaei, M., Aarts, C., Lee, D.A., Hassell, J., et al. (2017). Ex vivo expanded natural killer cells from breast cancer patients and healthy donors are highly cytotoxic against breast cancer cell lines and patient-derived tumours. Breast Cancer Res. 19, 76.

43. Huang, Y., Lv, S.Q., Liu, P.Y., Ye, Z.L., Yang, H., Li, L.F., Zhu, H.L., Wang, Y., Cui, L.Z., Jiang, D.Q., et al. (2020). A SIRPa-Fc fusion protein enhances the antitumor effect of oncolytic adenovirus against ovarian cancer. Mol. Oncol. 14, 657–668.

44. Advani, R., Flinn, I., Popplewell, L., Foster, A., Bartlett, N.L., Ghosh, N., Kline, J., Roschewski, M., LaCasce, A., Collins, G.P., et al. (2018). CD47 Blockade by Hu5F9-G4 and Rituximab in Non-Hodgkin’s Lymphoma. N. Engl. J. Med. 379, 1711–1721.

45. Klutz, K., Schaffert, D., Willhauck, M.J., Grünwald, G.K., Haase, R., Wunderlich, N., Zach, C., Gildehaus, F.J., Senekowitsch-Schmidtke, R., Göke, B., et al. (2011). Epidermal growth factor receptor-targeted (131)I-therapy of liver cancer following systemic delivery of the sodium iodide symporter gene. Mol. Ther. 19, 676–685.

46. Groza, D., Gehrig, S., Kudela, P., Holcman, M., Pirker, C., Dinhof, C., Schueff, H.H., Stramko, M., Hoerbart, J., Alioglu, F., et al. (2018). Bacterial ghosts as adjuvant to oxaliplatin chemotherapy in colorectal carcinomatosis. OncolImmunology 7, e1244676.

47. Liu, X., Pu, Y., Cron, K., Deng, L., Kline, J., Frazier, W.A., Xu, H., Peng, H., Fu, Y.X., and Xu, M.M. (2015). CD47 blockade triggers T cell-mediated destruction of immunogenic tumors. Nat. Med. 21, 1209–1215.

48. Albers, J., Danzer, C., Rechsteiner, M., Lehmanna, H., Brandt, L.P., Hejhal, T., Catalano, A., Busenhart, P., Gonçalves, A.F., Brandt, S., et al. (2015). A versatile modular vector system for rapid combinatorial mammalian genetics. J. Clin. Invest. 125, 1603–1619.

49. Maier, J., Elmenofi, S., Taschauer, A., Anton, M., Sami, H., and Ogris, M. (2019). Luminescent and fluorescent triple reporter plasmid constructs for Wnt, Hedgehog and Notch pathway. PLoS ONE 14, e0226570.

50. Taschauer, A., Polzer, W., Alieoglu, F., Billerhart, M., Decker, S., Kittelmann, T., Geppl, E., Elmenofi, S., Zehl, M., Urban, E., et al. (2019). Peptide-Targeted Polyplexes for Aerosol-Mediated Gene Delivery to CD49f-Overexpressing Tumor Lesions in Lung. Mol. Ther. Nucleic Acids 18, 774–786.