Genetically Encoded Immunophotosensitizer 4D5scFv-miniSOG is a Highly Selective Agent for Targeted Photokilling of Tumor Cells In Vitro

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Abstract
Tumor-targeted delivery of cytotoxins presents considerable advantages over their passive transport. Chemical conjugation of cytotoxic module to antibody is limited due to insufficient reproducibility of synthesis, and recombinant immunotoxins are aimed to overcome this disadvantage. We obtained genetically encoded immunophotosensitizer 4D5scFv-miniSOG and evaluated its phototoxic effect in vitro. A single-chain variable fragment (scFv) of humanized 4D5 antibody was used as a targeting vehicle for selective recognition of the extracellular domain of human epidermal growth factor receptor 2 (HER2/neu) overexpressed in many human carcinomas. As a phototoxic module we used a recently described photoactivated fluorescent flavoprotein miniSOG. We found that recombinant protein 4D5scFv-miniSOG exerts a highly specific photo-induced cytotoxic effect on HER2/neu-positive human breast adenocarcinoma SK-BR-3 cells (IC50 = 160 nM). We demonstrated that the 4D5scFv-miniSOG specifically binds to HER2-positive cells and internalizes via receptor-mediated endocytosis. Co-treatment of breast cancer cells with 4D5scFv-miniSOG and Taxol or junction opener protein JO-1 produced remarkable additive effects.

Key words: tumor-targeted delivery, immunophototoxin, phototoxicity, HER2/neu, single-chain variable fragment.

Introduction
Photodynamic therapy (PDT) is a minimally invasive cancer therapy that includes accumulation of a photosensitizer in the tumor and subsequent illumination with light of a specific wavelength [1, 2].

Being in activated state, a photosensitizer (PS) transfers the energy from light to the molecular oxygen and generates reactive oxygen species (ROS, free radicals or singlet oxygen) that cause destruction of the targeted tissue. As ROS have short lifetime, the reactions of oxidative damage occur in the immediate vicinity of the PS. A short range of action (~10 nm) in combination with precise light direction (for example, by means of fiber optics), ensures a high degree of spatial control during PDT.
Chemically based photosensitizing reagents have been widely used in photodynamic therapy [1]. However, due to accumulation in normal tissue commercially available PS (Photofrin, Foscan and Verteporfin) cause side effects such as skin and eyes photosensitization.

To overcome this obstacle, a new approach for drug delivery, called targeted photodynamic therapy, has been developed [3, 4]. One way for accurate delivery of PS to the tumor cells is chemical conjugation of PS to antibodies that recognize tumor-associated antigens. Following the pioneering work of Mew et al. [5] in the early 1980s the use of photosensitizer immunoconjugates became an advanced approach broadening the utility of photodynamic therapy [6]. Such macromolecular conjugates enable direct delivery of the PS to the target cell types. However, the main disadvantages of this approach are the limited reproducibility of the conjugate synthesis and the loss of activity of antibody or the PS [7, 8].

Another way for selective and effective therapeutic treatment is applying of genetically encoded PS.

We have previously reported a promising genetically encoded PS KillerRed, which can specifically recognize HER2/neu receptor (human epidermal growth factor receptor) and destroy adenocarcinoma cells in vitro due to KillerRed ability to generate ROS upon green light irradiation [9]. In general, HER2/neu, overexpressed on the surface of many cancer cells, is an important target for cancer therapy [10, 11]. The first to demonstrate this in animals were Drebin and co-workers [12], who proved that monoclonal antibody specific to the extracellular domain of HER2/neu protein possesses a significant anti-tumor effect. This research became the keystone for development of an antibody-based therapy of cancer. 4D5 full-size humanized monoclonal antibody is widely used in clinical practice for immunotherapy of HER2/neu-overexpressing tumors [10, 13]. A recombinant 4D5scFv mini-antibody is an example of high-efficiency HER2/neu-targeting vehicle that represents a single chain variable fragment of immunoglobulin (Ig) and exhibits lower cross-reactivity and immunogenicity and faster tissue penetration in comparison with the corresponding full-size antibody [13].

Recently a new highly phototoxic protein mini-iSOG (mini Singlet Oxygen Generator) has been reported [14]. This is a green fluorescent flavoprotein engineered from Arabidopsis phototropin 2, which uses flavin mononucleotide (FMN) as a cofactor. Upon blue-light illumination, miniSOG generates a sufficient quantity of singlet oxygen to induce diaminobenzidine (DAB) photooxidation and also photoinduces cell ablation in C. elegans [15], making it a potentially useful tool for PDT and optogenetics [14-16].

We have taken into account the advantage of the ROS-producing properties of miniSOG and created the recombinant immunoPS consisting of an anti-HER2/neu 4D5 single chain variable fragment (4D5scFv) and miniSOG as a phototoxic module.

In this study we show that genetically encoded immuno-photosensitizer 4D5scFv-miniSOG specifically targets and photo-kills HER2/neu overexpressing cells but spare HER2-negative cells in vitro. Excellent 4D5scFv-miniSOG targeting and photokilling properties in vitro give its opportunity to be a reasonable immunoPS in vivo.

Materials and Methods

Construction of 4D5scFv-miniSOG expression plasmid. miniSOG coding region was amplified from pC1-miniSOG plasmid [17] using specific primers (5’GGTGGCGCGCCCTATGGAAAAGAGCTTTGT, 5’GGCGCGCGCCGCCATCCAGCTGCACACC). The amplification product of 340 bp was digested with AscI (recognition region is underlined in primer structure) and cloned under lac-promoter in one reading frame with the coding sequence of 4D5scFv hybrid gene into pSD-4D5 [24]. E. coli colonies carrying inserts of miniSOG in direct orientation were selected. Generated plasmid pSD-4D5scFv-miniSOG was sequenced and proven to harbor correct coding sequence.

4D5scFv-miniSOG purification. Freshly transformed Escherichia coli SB536 strain [18] were grown in the YTPS medium (1% triptone, 1% yeast extract, 45 mM K2HPO4, 5 mM KH2PO4, 0.1 M NaCl, 2 mM MgCl2) containing 0.1 g/l ampicillin; the lac-promoter was induced with 1 mM IPTG at OD600 of 0.7. Expression was allowed to continue for 6 h at room temperature. Cells were harvested by centrifugation at 6,000g for 10 min at 4°C. Cell pellets were resuspended in lysis buffer (200 mM NaCl, 30 mM NaH2PO4, 2 mM Tris, pH 8.1) and then sonicated on ice in dark. Cellular debris was removed by centrifugation at 15,000g and the supernatant was applied to a Ni2+-NTA column (GE Healthcare) according to the manufacturer’s instruction. The immobilized proteins were eluted with 100 mM imidazole and equilibrated on a PD10 column (GE Healthcare) in storage buffer (20 mM KH2PO4, 20 mM HEPES, 10 mM EDTA, 0.5 M NaCl, 10% glycerol, 0.1% Tween, pH 8.0). The yield of recombinant protein was 1.5 mg purified protein per one liter of growth medium.

JO-1 purification. Plasmid pQE30-K-Ad3-6th-
SK (provided by Andre Lieber) encoding JO-1 protein was expressed in XL1-Blue (Stratagene) and purified on Ni²⁺–NTA column (GE Healthcare) according to the manufacturer’s instruction.

**SDS-PAGE and western blot analysis.** SDS-PAGE analysis was performed under denaturing conditions according to standard protocols using 12% polyacrylamide gels.

Band corresponding to 4D5scFv-miniSOG was excised from Coomassie blue-stained gel, digested with trypsin and subjected to MALDI–TOF mass spectrometry [19].

Immunoblots on Immobilon-P transfer membrane (Millipore) were carried out according to the manufacturer’s instructions using anti-His tag mouse monoclonal antibody alkaline phosphotase conjugate (Abcam) for detection. The blots were visualized with nitroblue tetrázolium/5-bromo-4-chloro-3-indolyl phosphate.

**Cell lines and incubation conditions.** The human breast adenocarcinoma SK-BR-3 (HTB-30™; ATCC) and Chinese hamster ovary CHO cell lines (Russian Cell Culture Collection) were maintained in RPMI-1640 medium (HyClone) supplemented with 10% fetal calf serum (HyClone) and 10% L-glutamine (PanEko). Cells were incubated in a 5% CO₂ atmosphere at 37°C.

**Internalization/plasma membrane localization studies.** SK-BR-3 cells were cultured on chamber-slides (Nalge Nunc International) overnight at 37°C, washed and incubated with 500 nM 4D5scFv-miniSOG at 4°C for 1 h following plasma membrane localization studies. For 4D5scFv-miniSOG internalization, cells were further washed and incubated for additional 30 min at 37°C. CHO cells were used as nontarget cells (negative control) to demonstrate HER2/neu-specific binding of 4D5scFv-miniSOG. The slides were analyzed with the laser scanning microscope Carl Zeiss LSM-710-NLO (Prokhorov General Physics Institute, Russian Academy of Sciences). The 4D5scFv-miniSOG was excited at 458 nm. The MitoTracker Red and LysoTracker Red organelle probe dyes were excited at the 561 nm. Colocalization analysis of photomicrographs was performed using the Carl Zeiss LSM-710-NLO software.

**Cell viability assay (MTT).** The effect of 4D5scFv-miniSOG or miniSOG on cell viability was studied using the microculture tetrazoline test (MTT) [20]. Briefly, SK-BR-3 and CHO cells were seeded in 96-well microtiter plates at 25,000 and 30,000 cells/ml, respectively. After 24 h media was removed, and the cells were treated with 4D5scFv-miniSOG or miniSOG at concentrations ranging from 1 nM to 500 nM. To investigate 4D5scFv-miniSOG cytotoxic action in combination with Taxol (Paclitaxel) or junction opened protein JO-1, cells were treated with 4D5scFv-miniSOG (1 to 500 nM) in the presence of 1 µM Taxol or 5µg/ml JO-1. To avoid unspecific effects the storage buffer was replaced with the RPMI-1640 medium. After 1 h incubation at 37°C the media was replaced by a fresh portion and the cells were exposed to white light with 1W/cm² from a lamp Dolan-Jenner Fiber-Lite 180 Illuminator (1W/cm² of white light gives 55mW/cm² of blue light within the efficient absorption range of 4D5scFv-miniSOG) 10 min. After 48 h incubation, growth medium was replaced by RPMI-1640 containing 0.5 mg/mL 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide. Cells were incubated at 37°C for 2 h, then the medium was withdrawn and 100 µl of DMSO (Sigma) was added to each well. Cell viability was quantified spectrophotometrically at 540 nm using a microplate reader (Awareness Technology, INC. USA). The cell viability was calculated as the percentage of proliferating cells relative to the number of untreated cells. All experiments were performed in triplicate.

**Statistical analysis.** All results are expressed as means ± standard deviations (SD). The statistical differences between groups of data were analyzed by one-way ANOVA. P<0.05 was considered significant.
Absorbance and fluorescence spectroscopy. Absorbance spectra (400–600 nm) of purified proteins (10 µM) were recorded on a Hitachi U-3400 spectrophotometer at 25°C. Emission spectra were recorded on a laser scanning microscope Carl Zeiss LSM-710-NLO in λ-mode regime at the excitation wavelength of 458 nm.

**K_D-Determination by surface plasmon resonance.** Dissociation constant (K_D) was determined by surface plasmon resonance with a BIAcore instrument (GE Healthcare Life Sciences). CM-5 sensor chips were coated with 1500 or 5000 resonance units of p185^HER-2-ECD antigen. 4D5scFv-miniSOG in concentration of 5 µM was injected in a final volume of 30 µl on the chips. Dissociation rate constant was calculated from the sensorgram using the BIAevaluation 3.0 software (GE Healthcare Life Sciences).

**Results**

**Design and characteristics of 4D5scFv-miniSOG**

It has been previously shown that natural N-terminus of 4D5scFv is important for proper antibody function [21]. Thus, the nucleotide sequence of the hybrid gene encoding anti-HER2/neu 4D5scFv fragment was placed at the 5'-terminus of the miniSOG coding sequence. The target and phototoxic domains were connected via flexible linker that prevented spatial interference between the two domains (Fig. 1A).

The resulting genetic construct (Fig. 1B) was expressed in *E. coli* SB536 strain [18] as the C-terminus His5-tag protein and purified using Ni²⁺-NTA affinity chromatography (Fig. 2A).

As was determined by SDS-PAGE electrophoresis, the produced 4D5scFv-miniSOG had an expected molecular mass (~ 44 kDa, Mw_{theor} 42 kDa) and was homogenous. The identity of the purified protein was verified by western blot analysis (Fig. 2B) and MALDI-TOF mass-spectrometry.

Absorption and emission spectra of 4D5scFv-miniSOG and free miniSOG are demonstrated in Fig. 2C. Excitation of the hybrid protein 4D5scFv-miniSOG leads to green emission with the two peaks at 500 and 526 nm (Fig. 2C) that is in good agreement with the fluorescence spectra for free miniSOG [14].

![Fig 1. 3D model and gene construct of the genetically encoded immunoPS. A, Molecular model (ribbon representation) of the 4D5scFv (PDB entry 1FCV) [22] and miniSOG. The 3D structure of miniSOG was made using SWISS-MODEL structure homology-modeling server [23, 24]. Cofactor FMN is shown in the miniSOG chromophore pocket. The image of 3D immunoPS structure was made using DS ViewerPro 5.0 software. B, Gene construct for expression of the 4D5scFv-miniSOG. OmpA, signal peptide for direct secretion of the recombinant protein to the periplasmic space of *E. coli* (white); VL, light chain of 4D5scFv (magenta); L, designed peptide linker (21 a.a., brown) that links the carboxyl terminus of the VL sequence to the amino terminus of the VH sequence; VH, heavy chain (blue); H, hinge-like linker (16 a.a., red) derived from the murine IgG3 hinge region; miniSOG, coding region of miniSOG (green); His, C-terminal 5-His-tag (yellow).](http://www.thno.org)
Evaluation of the HER2/neu binding

Binding of 4D5scFv-miniSOG to a purified HER2/neu receptor was evaluated by Plasmon surface resonance (Fig. 2D). The $K_D$ value of the 4D5scFv-miniSOG to HER2/neu receptor was determined by BIAdore to be 10±2 nM, which corresponds well to that of 4D5scFv (9±2 nM) [25].

Breast adenocarcinoma SK-BR-3 cells, which express ~1x10^6 to 2x10^6 HER2/neu receptors per cell [26], were chosen as a model cell line. As was determined by confocal microscopy, 4D5scFv-miniSOG accumulated effectively on the cellular membrane surface after 1 h incubation at 4°C (Fig. 3A). After removing all non-specifically bounded 4D5scFv-miniSOG by washing with PBS, the cells were left for 30 min at 37°C for internalization. Fluorescent signal inside the SK-BR-3 cells on the micrograph (Fig. 3B) clearly indicates the ability of 4D5scFv-miniSOG to penetrate into the cells by receptor-mediated endocytosis. Fluorescent signal was undetectable when free miniSOG was added to SK-BR-3 cells (Supplementary Material: Fig. S1A). No binding activity was detected in the case of CHO cells (negative control) that do not overexpress HER2/neu (Supplementary Material: Fig. S1B).

**Competition experiments**

4D5scFv-miniSOG binding specific properties was also investigated in competition experiments with 4D5scFv mini-antibody. Figure 4 clearly demonstrates that SK-BR-3 binding of 4D5scFv-miniSOG was inhibited when competed with 4D5scFv. Integrated fluorescence intensity signal for SK-BR-3 cells treated with 500 nM 4D5scFv-miniSOG was in 2.5-fold greater than for SK-BR-3 cells treated with 500 nM 4D5scFv-miniSOG in the presence of 500 nM 4D5scFv (Fig. 4C). The SK-BR-3 cell amount was equal in both cases.

**Intracellular localization**

For a better understanding of the immunoPS-mediated PDT, we characterized the intracellular localization of the immunoPS and quantified...
the colocalization patterns with lysosomes and mitochondria.

Micrographs of the SKBR-3 cells double stained with 4D5scFv-miniSOG and LysoTracker Red or MitoTrackerRed are presented in Figure 5. 4D5scFv-miniSOG was shown to localize mainly in the cytoplasm. Only weak fluorescence intensity was found in the nucleus. Yellow areas correspond to the colocalization fraction of 4D5scFv-miniSOG and organelle specific dyes (Figures 5Aiii and 5Biii). Thus, lysosomes localized mainly at the same site of the cytoplasm where 4D5scFv-miniSOG accumulation occurred (Figure 5Aiii). But yellow areas were slightly detectable in cells double-stained with 4D5scFv-miniSOG and the MitoTrackerRed (Figure 5Biii).

In order to increase the significance of the co-staining experiments, fluorescence images were analyzed quantitatively by the Carl Zeiss LSM-710-NLO software, and results are depicted in scatterplots (Figure 5Aiv, Biv).

Colocalization coefficients were calculated as relative number of colocalizing pixels in channel 1 or 2, respectively, as compared to the total number of pixels above threshold. In cells double stained with 4D5scFv-miniSOG and LysoTracker, the colocalization fraction of 4D5scFv-miniSOG found in the lysosomes was 21% (Ch2-T1), whereas 72% (Ch1-T1) of the lysosome fluorescence coincided with the 4D5scFv-miniSOG fluorescence (Figure 5Aiv). Due to receptor-mediated nature of 4D5scFv-miniSOG endocytosis it was of our expectation to observe its localization in lysosomes. In the case of cells double stained with 4D5scFv-miniSOG and MitoTracker, there was also some fluorescence coincidence in mitochondrions vs 4D5scFv-miniSOG and vice versa (colocalisation coefficients were 28% (Ch2-T1) and 23% (Ch1-T2) respectively) (Figure 5Biv).

Thus, the majority of the immunoPS is located in cytoplasm. As we previously proved an absence of unspecific entry of 4D5scFv-miniSOG through the cell membrane it may indicate its endosomal localization or the photo-induced release of 4D5scFv-miniSOG from lysosomes under laser scanning manipulation.

Fig 3. Confocal fluorescence images of SK-BR-3 cells after 1 h incubation with 4D5scFv-miniSOG at 4°C (A) and after additional 30 min incubation at 37°C (B). Left row, a bright-field image; middle row, fluorescence image in green channel; right row, fluorescence image in green and blue channels (nuclei stains with Hoechst 33258).
Fig 4. 4D5scFv-miniSOG competitive uptake experiments. HER2/neu-overexpressing SK-BR-3 cells were incubated for 1 h at 37°C with 500 nM 4D5scFv-miniSOG (A) or 500 nM 4D5scFv-miniSOG plus 500 nM 4D5scFv (competitive mini-antibody) (B). The integral signal of fluorescence in both cases were measured using confocal microscopy and presented as a histogram (C): red field – fluorescence signal corresponding to 4D5scFv-miniSOG-treated cells, black field – fluorescence signal corresponding to 4D5scFv-miniSOG plus 4D5scFv-treated cells.

Fig 5. 4D5scFv-miniSOG localization in different organelles in SK-BR-3 cells. 4D5scFv-miniSOG vs LysoTracker (A), 4D5scFv-miniSOG vs MitoTracker (B). Representative 4D5scFv-miniSOG images of the cells are shown in green (i), images of each organelle-specific dye are shown in red (ii), and merged fluorescent images are displayed in iii. Scatterplots of 4D5scFv-miniSOG and organelle-specific dye signal frequency and intensity are shown in iv. The cell on fig. Biii indicated in red frame was chosen for scatterplot measurement.

Specific photo-induced cytotoxicity of 4D5scFv-miniSOG

In order to estimate the specific photo-induced cytotoxic effect of 4D5scFv-miniSOG on the viability of SK-BR-3 cells, a microculture tetrazolium test (MTT) was used. Decreasing concentrations of immunoPS were added to SK-BR-3 cells, than incubated for 1 h at 37°C, followed by exposure to light from a lamp (Dolan-Jenner Fiber-Lite 180 Illuminator) with 1 W/cm² of bright white light (55 mW/cm² of blue light within the efficient absorption range of 4D5scFv-miniSOG) for 10 min, the percentage of viable cells was determined compared to the untreated cells. Using regression analysis, IC₅₀ was estimated as 160 nM. We observed that the maximal cytotoxic effect (21% of viable cells) for SK-BR-3 cells was achieved at 500 nM of 4D5scFv-miniSOG (Fig. 6A). It should be noted that this cytotoxic effect of 4D5scFv-miniSOG was observed under 55 mW/cm² blue light intensity, but not under typical indoor illumination (~0.2 mW/cm²) (Fig. 6A). ImmunoPS did not affect viability of the CHO cells in the dark or under white light even if treated with maximum concentration of immunoPS (Fig. 6B).

Free 4D5scFv, free miniSOG and free FMN did not affect cell viability (Fig. 6C), indicating high specificity of the 4D5scFv-miniSOG phototoxic effect.

We also investigated the cytotoxic action of 4D5scFv-miniSOG in combination with mitotic inhibitor Taxol (Paclitaxel) or junction opener protein JO-1 [27].
Taxol is a well-known cytostatic drug, and in our experiment it was used to estimate the cytostatic action of 4D5scFv-miniSOG in non-proliferating cell culture. Checking several toxic concentrations of Taxol we have found that 1 μM of anticancer drug causes 40% cell death (Supplementary Material: Fig. S2), which is in good agreement with literature data [28]. This concentration of Taxol was further used in toxicity experiments. SK-BR-3 cells were treated with various concentrations of 4D5scFv-miniSOG and 1 μM of Taxol, and the percentage of viable cells relative to the immunoPS/drug-free control was determined. We have found that 1 μM of Taxol significantly increased cytotoxicity of 4D5scFv-miniSOG, so that co-treatment produced 100% cell death (Fig. 6A). On the other hand, the same concentration of Taxol showed no increase in 4D5scFv-miniSOG citotoxicity on CHO cells (Fig. 6B).

Another agent that we used in our experiments was junction opener protein JO-1. JO-1 is a self-dimerizing recombinant protein derived from the adenovirus serotype 3 fiber [27]. The pretreatment of epithelial cells with JO-1 resulted in increased access to receptors that are localized in or masked by epithelial junctions [27]. JO-1 was shown recently to increase intratumoral penetration of the anti-HER2/neu mAb Herceptin [29]. We tested whether JO-1 would improve SK-BR-3 killing by immunoPS. Cells in 100% confluency were co-incubated with JO-1 and 4D5scFv-miniSOG for 2 h and then irradiated with white light. Addition of 5 μg/ml JO-1 decreased IC50 of 4D5scFv-miniSOG from 160 to 5 nM (Fig. 6A).

**Fig 6. In vitro cytotoxicity analysis of 4D5scFv-miniSOG.** A. The impact of 4D5scFv-miniSOG decreasing concentrations on the SK-BR-3 cell viability under different conditions: white light and dark, taxol or JO-1 cotreatment. B. Relative viability of CHO after the treatment with 4D5scFv-miniSOG under white light or in the dark and after the co-treatment with 4D5scFv-miniSOG and Taxol under white light. C. Effects of free 4D5scFv, free miniSOG and free FMN on the SK-BR-3 cell viability. Bars indicate SD for 3 independent experiments. P<0.05.

**Discussion**

In this study we have clearly demonstrated that genetically encoded PS 4D5scFv-miniSOG can be successfully used for tumor cell destruction *in vitro.*

Our experiments indicate that photo-damage of tumor cells by targeting the 4D5scFv-miniSOG to HER2/neu-overexpressing cells is highly specific only for these cells and occurs only upon light irradiation. Being in fuse with miniSOG, 4D5scFv reserves binding activity to HER2/neu (K_D =10±2 nM) and at the same time miniSOG retains its ability to generate ROS, which is proved by its cytotoxic effect on SK-BR-3 cells *in vitro.* Our data (Figs. 4 and 6) clearly show that the phototoxic effects of 4D5scFv-miniSOG is mediated via specific HER2/neu targeting and is not due to the nonspecific effects.

*In vitro* cell killing using 4D5scFv-miniSOG is much effective than in the case of 4D5scFv-KillerRed [9]. Being compared to anti-HER2/neu scFv-photosensitizer immunoconjugates, 4D5scFv-miniSOG showed a greater than 8-fold and 30-fold improvement of IC50 over anti-HER2/neu-pyropheophorbide-a and anti-HER2/neu-verteporfin respectively [30]. Most of photosensitizers used in photodynamic therapy are hydrophobic and lipo-
philic and tend to aggregate in aqueous solutions. So it is not easy to conjugate them to antibodies. Besides this many photosensitizer immunoconjugates can contain some amounts of free photosensitizer impurities. Also conjugation can sterically hinder or disrupt antibody antigen binding sites [31]. Being genetically-encoded, 4D5scFv miniSOG has fixed photosensitizer/antibody ratio, a defined junction of two functional modules, absence PS impurity, as well as stable and simple production of the active recombinant protein in bacteria.

There is one more approach utilizing minISOG as a genetically encoded photosensitizer for cancer cells has been recently published [17]. Ryumina and co-workers received HeLa Kyoto cell lines stably expressing minISOG and established its phototoxic effect in vitro. But they did not observe photoinduced tumor reduction in vivo. Authors believe that such unexpected in vivo result can be due to unsaturation of minISOG by its cofactor FMN. In this case our approach based on the production of minISOG in bacteria has one crucial advantage: bacterial expression allows to obtain minISOG fully saturated with FMN which is extremely necessary for strong damage of cancer cells. Moreover we produce HER2/neu-targeting minISOG that broadens the utility of PS in vivo.

Generally, photosensitizers can generate type I (photoinduced electron transfer) or type II (energy transfer from exited photosensitizer to oxygen) photochemical reactions. It has been previously shown that due to bounded FMN, excited minISOG generates singlet oxygen with a high yield (0.47) [14] and thus to be a type II photosensitizer. In contrast to this claim there are two recent publications stated that minISOG indeed is a very poor singlet oxygen sensitizer, and FMN being enclosed in minISOG produces a reactive species other than 1O2 [32, 33]. So minISOG may be considered as type I photosensitizer. Taking into account all the statements [32, 33] and our in vitro results we can conclude that minISOG does initiate photochemical reactions defining its high cytotoxic properties. This result can be due to both type I and type II photochemical reactions.

Combining PDT with other therapies is known to increase effectiveness of anti-tumor treatments [1, 34]. We showed that co-treatment of cells with immunoPS and mitotic inhibitor Taxol leads to 100% SK-BR-3 cells death, while complete cell killing was not achieved if 4D5scFv miniSOG was applied alone (Fig. 6A). This strategy gives opportunity to use drugs in lower concentrations and decreases possibility of side effects.

Another way for the reduction of immunoPS doses is to improve an access to its target receptor. It has been shown previously, that adenovirus junction opener protein JO-1 is capable of transient opening of intercellular junctions, leading to increase of intratumoral penetration of the anti-HER2/neu monoclonal antibody trastuzumab (Herceptin) [29]. We have found that JO-1 enhances immunoPS cytotoxicity, significantly improving the outcome of the breast cancer treatment with 4D5scFv miniSOG (Fig. 6A).

It is obvious that the efficacy of the diseased tissue destruction during photodynamic therapy depends on the depth of the light penetration. Blue light has only limited propagation through the tissues. As 4D5scFv miniSOG is excited by blue light, its application in vivo might require light delivery system coupled with fiber-optic technology allowed the interstitial light delivery [35]. And we hope that well-coordinated interdisciplinary efforts would cope with this challenge. Correct approach might enable in vivo implementation of 4D5scFv miniSOG cytotoxic potential and could make photodynamic properties of our immunoPS comparable to other commercially available red PSs.

Supplementary Material

Figure S1. Confocal fluorescence images of SK-BR-3 (A) and CHO (B) after 1 h incubation with free minISOG (500 nM) at 37°C. Left row, a bright-field image; right row, green fluorescence image.

Figure S2. Relative cell viability of SK-BR-3 cells after treatment with various Taxol concentrations. http://www.thno.org/v03p0831s1.pdf

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Author contributions

K.M., G.P., A.R., O.S., S.L., R.P. and S.D. designed research; K.M., G.P., A.R. and O.S. performed research; K.M., G.P. and A.R. analyzed data; G.P.
K.M. and S.D. wrote the paper.

Competing Interests

The authors have declared that no competing interest exists.

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