Abstract. Photodynamic therapy (PDT) is a minimally invasive cancer therapy that depends on the buildup of a photosensitizing drug within targeted tissue. The photosensitizer is subsequently activated by light of a specific wavelength, resulting in destruction of the targeted tissue by free radicals or singlet oxygen. Successful treatment requires delivery of critical amounts of drug into the cancerous tissue. This frequently demands high doses of the drug in the circulatory system that could lead to side effects in normal daylight due to accumulation of photodrug in normal tissue. To reduce drug load we explored the possibility of targeting cancer with antibody conjugated with photosensitizer. As a model we used LnCAP human prostate cancer cells targeted by antibody (against prostate-specific membrane antigen) which was conjugated with hematoporphyrin (HP). Our results show clearly that mAb/HP conjugates can deliver HP to the tumor cells which would result in considerably less HP in the circulation and, therefore, lower the delivery of HP to normal tissue, resulting in fewer side effects.

Materials and methods

CYT-351 antibody against prostate-specific membrane antigen was purchased from Cytogen Princeton, NJ, USA.

Conjugation of CYT-351 with hematoporphyrin. Direct conjugation of hematoporphyrin (HP) and CYT-351 was done as described by Mew et al (5,6). All reactions were carried out in room temperature and in low light, unless stated differently. Briefly, 60 µM of HP were dissolved with 3 mM of anhydrous 1-ethyl-3-(dimethyl aminopropyl) carbodiimide (EDC) from Pierce (Rockford, IL, USA) and in 6 ml of anhydrous dimethyl sulfoxide (DMSO). Then mixture was stirred for 12 h, and stored at room temperature not more than 10 days. EDC is a zero-length cross linking instrument coupling carboxyl groups to primary amines (Fig. 1). An aliquot of 175 µl of HP-EDC was added to 3.5 ml of antibody (1 mg/ml) and mixed for 1 min in light protected vessel. Conjugated antibody was separated from substrates on PD-10 column from Amersham Biosciences Corp. (NJ, USA). PD-10 column was equilibrated with 25 ml of PBS and 2.5 ml of reacting mixture was applied on top of column, the flow-through was discarded. Next 3.5 ml buffer was added and flow-through containing conjugated antibody was collected.

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**Conjugation of hematoporphyrin and CYT-351 OX by biotin avidin bridge.** We used EZ-Link Hydrazide-LC-Biotin of hydrazide-activated biotin that enables biotinylation of glycoproteins having oxidizable sugars or aldehydes (Pierce) following the manufacturer’s instructions. Weak oxidation of antibodies with sodium periodate (NaIO₄) forms reactive aldehydes on the carbohydrate moieties of the Fc chain to be modified by hydrazides (Fig. 2). This method is beneficial for labeling antibodies since biotinylation takes place only at the sites of glycosylation, which are primarily in the Fc region, far from the antigen binding site (11-13). Briefly, 1 ml of cold sodium periodate solution (20 mM sodium periodate in 0.1 M sodium acetate buffer, pH 5.5) was added to 1 ml of cold CYT-351 antibody (1 mg/ml) mixed and incubated for 30 min at 4°C. Excess of periodate was removed through a desalting column that previously has been equilibrated with coupling buffer (0.1 M sodium phosphate, 0.15 M NaCl, pH 7.2). Next 1 part of biotin hydrazide solution (50 mM) was added to 9 parts of oxidized and buffer-exchanged antibody and mix for 2 h. The biotinylated CYT-351 was separated from non-reacted material by gel filtration (desalting column). Conjugation of hematoporphyrin with avidin (NeutrAvidin Protein (Pierce) was done as described for CYT-351. Biotinylated antibody and HP conjugated avidin where combined 1:1 ratio and mixed for 1 h. Next mixture buffer was adjusted to the composition of the binding buffer (20 mM sodium phosphate, pH 7.0) by buffer exchange using PD-10 desalting column. Conjugated antibody was loaded onto HiTrap Protein G column, Pharmacia Biotech Inc. (Piscataway, NJ, USA). Column was washed with 10 ml of binding buffer and antibody was eluted with 5 ml of elution buffer (0.1 M glycine-HCl, pH 2.7). The purified fractions were immediately buffer-exchanged using PBS and frozen until used.

**Concentration measurement.** Protein and HP concentration were measured spectroscopically at 280 nm for mAb and at 375 nm for HP.

**Gel electrophoresis.** Electrophoresis of HP conjugated CYT-351 antibody was performed at room temperature using 10% SDS-polyacrylamide gels under non-reducing conditions. Gels were stained with Coomassie brilliant blue, Invitrogen (Grand Island, NY). Gels were illuminated with visible and UV light and photographed.

**Prostate cancer cells.** LnCAP human prostate cancer cells were purchased from American Type Culture Collection (ATCC), (Manassas, VA, USA). The cells were propagated in RPMI-1640 medium, Sigma Chemical Co. (St. Louis, MO), and supplemented with 10% fetal bovine serum, Hyclone Laboratories (Logan, UT, USA), penicillin and streptomycin.

**PDT treatment of prostate cancer cells; LnCAP.** Cells were incubated with HP (48, 24, 12, 6, 3 µg/ml) or CYT-351/HP and CYT351 OX/HP (10, 5, 2.5, 1.2, 0.6 µg/ml of HP conjugated to antibody) for 180 min in 24-wells cell culture dish. Next media was removed and washed three times with PBS. Finally 0.5 ml of PBS was added and cells were irradiated to deliver 0.21 J/cm of red light (ruby red filter from Kodak - >610 nm).
Total emitted power was verified before and after PDT photoradiation using an integrating sphere-based power meter (Labsphere, Hanover, NH, USA). After light treatment, cells were trypsinized and resuspended in 5 ml of media and seeded in P100 dish for clonal assay.

Figure 2. Conjugation of hematoporphyrin (HP, shown in red) conjugated to avidin (ribbon model, green color) and antibody (ribbon model colored as rainbow) using Hydrazide-LC-Biotin linker (EDC, shown in blue).

Figure 3. Coomassie stained CYT-351/HP conjugate (upper picture), the same antibody conjugate illuminated by UV light (lower picture).

Figure 4. LnCAP human prostate cancer cells treated with: CYT-351 Ox/HP (green squares); CYT-351/HP (blue circles); HP (red triangles). Concentration of HP conjugated to antibodies was equal to: 10, 5, 2.5, 1.2, 0.6 µg/ml and non-conjugated HP was equal to: 48, 24, 12, 6, 3 µg/ml.
Results and Discussion

Prostate specific membrane antigen is a cell-surface glycoprotein predominantly found in prostate tissue and overexpressed on prostate tumor cells. Expression levels increase as disease progress, and highest levels are observed in metastatic disease. Furthermore, PSMA has also been detected in the neovascularure of a variety of non-prostatic solid malignancies (14). Therefore, PSMA as a biomarker and target for the delivery of therapeutic agents attracted considerable attention in cancer therapy. We chose antibody CYT-351 against PSMA for targeted delivery of PDT agent.

For targeted PDT to be successful, critical amounts of photosensitizer must be conjugated to the MAb without altering its activity. Few methods of conjugation can be used to conjugate the antibody and drug. One of the most frequently used conjugation methods involves the use of a zero-length crosslinker (Fig. 1), EDC for the formation of amide bonds between carboxyl groups and primary amines of antibody (predominantly lysine). However, in this coupling method, there is a risk that the antigen binding sites at the Fab region of antibody are blocked by the nonselective formation of HP/antibody bonds (1,2,15-17). Thus, we chose the second, method of conjugation into the carbohydrate moieties of the Fc chain of the antibody, far from the active site. We use Hydrazide-LC-Biotin linker with spacer of 23.7 Å (Fig. 2). Disadvantage of this method over EDC conjugation is the lengthy and multistep procedure (12,13). Both methods yielded CYT-351/HP and CYT-351 Ox/HP conjugates. As seen in Fig. 3, stained PAGE gel showed distinct bands of HP conjugated CYT-351 antibody. Furthermore, gel illuminated with UV light showed red colored bands of antibody characteristic to HP fluorescence; nonconjugated antibody does not produce any fluorescence.

As shown in Fig. 4 significant cancer cell killing was observed for cells incubated with CYT-351/HP conjugated and non-thermal light treatment starting with concentration of 1.25 µg/ml of HP conjugated to antibodies. Both forms of antibody/HP conjugates had similar cancer cell killing potency indicating that conjugation did not affect the binding site of the antibody but delivered therapeutic amount of HP into cancer cells. Similar cancer cell ablation effect can be observed for unconjugated HP but at concentration of ~24 µg/ml. Above indicates improved targeting delivery and accumulation of HP in cancer cells by CYT-351 antibodies over HP alone. Targeting the tumor cells with antibodies/HP conjugates would result in considerably less HP in the circulation and consequently lower the general skin sensitivity of patients to light. That was the most important contraindication observed in human patient trials (18-22). Additional benefits of PDT with CYT-351 conjugates include damage to newly formed cancer angiogenic vessels which can lead to increased vascular permeability that could greatly improve PDT’s therapeutic margin (23).

CYT-351 was developed in the 1980s and is successfully used in therapy, diagnosis by conjugating CYT-351 with radioactive elements (24-27). Recently, an interesting and similar to our approach to target PSMA with bioconjugate of photosensitizer and inhibitor of PSMA was presented by Liu et al where PSMA targeting small-molecule of this conjugate served as a delivery vehicle for PDT of prostate cancer. They found that PSMA+ cells treated with PS-inhibitor/photosensitizer conjugate and subsequent light treatment underwent apoptosis while PMSA+ cells did not (28). This work confirms our findings that PSMA can be used as a target in for PDT immunotherapy.

Our results clearly demonstrate the potential use of CYT-351/HP conjugates for killing of selectively targeted cells using photo-activation the HP. However, the possible phototherapy might require additional work most likely including optimization of dose, ratio of mAb to photosensitizer or even humanization of antibodies to reduce immune response.

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References


