Abstract. It was confirmed that CD147 (Emmprin) was expressed on the cell surface of carcinoma cells. For the purpose of studying the efficacy of a CD147-targeting agent on CD147-expressing carcinoma cells, we investigated the effect of a conjugate of glutathione-doxorubicin (GSH-DXR) encapsulated in an anti-CD147 antibody-labeled liposome (aCD147ab-liposome) in terms of specific accumulation and cytotoxicity in CD147-expressing human carcinoma cells. Expression of CD147 was not observed in many normal human tissues. However, slight expression of CD147 in kidney, prostate and breast tissues was observed. By contrast, high-level expression of CD147 in all carcinoma cells such as A431, PC3 and Ishikawa cell lines was confirmed by fluorescent microscopy and Western blot analysis. Specific accumulation of the aCD147ab-liposome in the above-described CD147-expressing cells was observed. GSH-DXR encapsulated in an aCD147ab-liposome expressed specific cytotoxicity against these carcinoma cells. These results suggested that target chemotherapy of GSH-DXR encapsulated in an aCD147ab-liposome on CD147-expressing carcinoma cells was effective.

Introduction

Conjugates of monoclonal antibodies (MAbs) with drugs or toxins have been investigated for many years as a potential approach to delivering these agents more specifically to cancers (1). For example, rituximab, a chimeric monoclonal antibody targeted against the pan-B-cell marker CD20, was the first monoclonal antibody to be approved for therapeutic use (2-4). Treatment with rituximab at standard weekly dosing is effective in more than 50% of patients with relapsed or refractory CD20-positive follicular non-Hodgkin's lymphoma (5). Moreover, drug-MAb conjugates are currently under development for the treatment of various solid tumors (6).

Degradation of the basement membrane by matrix metalloproteinases (MMPs) is one of the most critical steps in the various stages of tumor disease progression, including tumor angiogenesis, tumor growth, as well as local invasion and subsequent distant metastasis (7-9). Extracellular matrix MMP, a family of zinc-dependent proteolytic enzymes, plays a central role in these processes, due to the ability to break down basement membranes and most extracellular matrix (ECM) components. Extracellular matrix MMP inducer (EMMPRIN, also known as CD147 or basigin) is a 55-kDa molecule that is found on the surface of tumor cells and up-regulates the expression of MMPs in surrounding fibroblasts and endothelial cells to enhance the invasiveness of cancer cells (10). Involvement of tumor-derived EMMPRIN in tumor angiogenesis through stimulation of MMPs and VEGF production was demonstrated by Tang et al (11), suggesting that CD147 could represent a key molecule in tumor cell invasion and metastasis. To assess its putative role as a target for anticancer therapies, Riethdorf et al (12), analyzed the overall incidence of CD147 expression in a wide spectrum of normal and tumor tissues. They showed that CD147 expression was detected frequently in the vast majority of human malignancies and also in a subset of benign tumors. Nevertheless, there are some very interesting differences both in the intensity and distribution of CD147 staining among different malignant tumors as well as benign lesions (12). It was predicted that CD147 has the potential to serve as a target for anti-tumor therapy.

A murine monoclonal antibody (MoAb 12C3) that is specific to human ovarian carcinoma was generated by immunizing mice with a human ovarian germinoma cell line (JOHYC-2) in a previous study by us (13). A single-phage clone was purified by immunostaining with MoAb 12C3 among the phages selected from either the human ovarian carcinoma (SKOV3) or colon carcinoma cDNA libraries, termed SKOV3-1 and colon-1, respectively. Based on the selected DNA sequences of the phage, a homology search
using an amino acid sequence with the BLAST program was performed and revealed that SKOV3-1 and Colon-1 matched the 17th-174th and the 17th-149th amino acid residues, respectively, which were on the extracellular region of CD147.

On the other hand, a recent study revealed that a conjugate of doxorubicin (DXR) with glutathione (GSH) (GSH-DXR) potently induced apoptosis in rat hepatoma AH66 cells relative to DXR (15-17). The same study also showed that GSH-DXR inhibited GSH activity and suppressed P1 mRNA, but DXR did not, indicating that inhibition of the enzyme makes an important contribution to manifestation of potent GSH-DXR cytotoxicity against AH66 cells (18,19).

Therefore, it was predicted that the disappearance of GST P1-1 enzymatic activity did not suppress JNK activity in spite of binding to the JNK molecule and potently inducing apoptosis.

In the present study, we investigated the therapeutic efficacy of the aCD147ab-DXR conjugate for the treatment of several cancer cell lines. Moreover, GSH-DXR encapsulated in the aCD147ab-liposome was also examined for its specific cytotoxicity against these carcinoma cells.

Materials and methods

Materials. DXR was obtained from Kyowa Hakko Kogyo (Tokyo, Japan). GSH, MTT, 5-bromo-4-chloro-3-indolyl phosphate and nitro blue tetrazolium (NBT) (Dojindo, Japan). Distearoyl-phosphatidylethanolamine-polyethylene glycol maleimide (DSPE-PEG-MAL), DSPE-PEG-activated carboxylic acid (PE-PEG-NHS) and Coatsome (EL-01-A) were obtained from Nippon MARUNISHI (Tokyo, Japan). A431, Ishikawa and PC3 were cultured with RPMI-1640 medium (containing 10% heat inactivated fetal bovine serum (growth medium)) under conventional conditions.

Preparation of aCD147 monoclonal antibody. Murine anti-human CD147 monoclonal antibody (aCD147ab, MoAb 12C3) that is specific to human ovarian carcinoma was generated by immunizing mice with a human ovarian germinoma cell line (JOHYC-2) in our study (13).

Cell lines. Human epithelial carcinoma cell line, A431, human placental carcinoma cell line, Ishikawa, and human prostate carcinoma cell line, PC3 were cultured with RPMI-1640 containing 10% heat inactivated fetal bovine serum (growth medium) under conventional conditions.

Knockdown of CD147 expression in PC3 cells. Plasmid of shRNA for human CD147 (0.1 μg of plasmid in 10 cm dish) was transfected into PC3 cells using the FuGENE6 transfection reagent (Roche). The transfectants were selected by treatment with puromycin. A permanent CD147-suppressive clone (PC3/CD147-KD) was obtained.

Conjugation of DXR with GSH. GSH-DXR was prepared as described previously (16-19). In brief, the combination of 1 mg of each GSH and 0.5 mg of DXR in 0.5 ml of 0.15 M NaCl containing 0.1% glutaraldehyde was incubated at room temperature for 30 min. After incubation, GSH-DXR was separated from GSH and DXR using Dowex 50 W x8 (H-form, 5 x 15 mm). The obtained GSH-DXR was filter-sterilized by a 0.45-μm syringe filter (Corning Costar, Tokyo, Japan). The concentration of DXR was measured by absorbance at 495 nm. A conjugation of aCD147ab and DXR was carried out in 0.1% glutaraldehyde at room temperature for 30 min. aCD147ab-DXR was purified by gel chromatography on Sephadex G-100.

Preparation of immuno (aCD147ab)-liposome. One milligram of aCD147ab was mixed with 0.3 mg of DSPE-PEG-NHS (activated carboxylic acid-terminal for NH2 group reaction) in 0.1 M potassium phosphate buffer (K-PB) (pH 8.0) at room temperature (RT) overnight. One milligram of recombinant GFP was mixed with 0.3 mg of DSPE-PEG-MAL (maleimide-terminal for SH group reaction) in 0.1 M K-PB (pH 6.5) at RT overnight. After the reaction, excess non-reactive DSPE-PEG-NHS and DSPE-PEG-MAL were quenched by incubation with 50 mM Tris-HCl (pH 8.0) and 2 mM 2-mercaptoethanol, respectively, for 30 min at RT. 2-ME, DSPE-PEG and unconjugated antibody were removed by gel chromatography on Sephacryl S300HR. The resulting DSPE-PEG-aCD147ab and/or DSPE-PEG-GFP were mixed with 10 mg of Coatsome EL-01-A in EtOH. Excess PBS was added to the mixture and centrifuged at 20,000 g for 1 h. The precipitate was suspended in PBS containing 100 μM GSH-DXR and sterilized by passage through a 0.45-μm sterile filter.

Expression of CD147 in human carcinoma cells (A431, Ishikawa and PC3). The cells were incubated with 1 μg/ml of aCD147ab for 1 h. The cells were washed with PBS and were incubated with Alexa546-labeled anti-mouse IgG. The staining CD147 was observed by fluorescent microscopy.

Western blot analysis. CD147 in the cell extract with 1% Triton X-100 was separated by SDS-PAGE (10% acrylamide) and analyzed by Western blot using aCD147ab as the primary antibody and ALPase-labeled anti-mouse IgG as the secondary antibody.

Immunoreaction of aCD147-labeled liposome. The cells were incubated with aCD147ab- and GFP-double labeled liposome for 2 h. The cells were washed with PBS and were incubated with Alexa546-labeled anti-mouse IgG. The fluorescence of GFP and Alexa546 were detected by fluorescent microscopy.

Cytotoxicity of GSH-DXR-encapsulated immuno-liposome. Cytotoxicity of GSH-DXR-encapsulated immuno-liposome against A431, Ishikawa, PC3 and PC3/CD147-KD cells treated for 4 h was measured after 96 h by MTT assay. The cell death rate for aCD147ab-liposome/GSH-DXR (0.1 μM) and mouse IgG-liposome/GSH-DXR (0.1 μM) was expressed as drug-non treated control (100%).
Protein determination. Protein concentration was assayed by a Bio-Rad protein assay kit using BSA as the standard.

Results

Expression of CD147 in A431, Ishikawa, PC3 and PC3/CD147-KD cells with immunofluorescence and Western blot analysis. Representative profiles of immunofluorescence for CD147 are shown in Fig. 1A. Specific staining for CD147 was commonly observed in the cellular membrane of A431, Ishikawa and PC3 cells, whereas little immunofluorescent staining of the cellular membrane of CD147-knockdown cells, PC3/CD147-KD, was observed. Moreover, the expression of CD147 (55-kDa) in A431, Ishikawa and PC3 cells at protein
levels was observed by Western blot analysis using 12C3 as the primary antibody. Nevertheless, expression in PC3/CD147-KD cells was not detected (Fig. 1B). The reason why broad and some bands of CD147 detected in this Western blot analysis was dependent upon different amount of glycosylation ratio of CD147.

Cytotoxicity of aCD147ab-DXR conjugates. We prepared the conjugate with DXR and aCD147ab, and examined the cytotoxicity. As shown in Fig. 2, after continuous exposure to aCD147ab-DXR, the A431, Ishikawa, PC3 and PC3/CD147-KD cells exhibited the same degree of cytotoxicity in comparison to murine IgG-DXR and BSA-DXR used as the negative control.

As the 80% growth-inhibitory concentration value of these drugs was approximately 1 μM of DXR-equivalent with these cell lines, the results of treatment conducted using the drugs at the concentration of 1 μM, with the cells exposed to the drugs for 2, 4 and 6 h after 96 h of incubation in drug-free medium are shown in Fig. 3. Cytotoxicity of aCD147ab-DXR was apparently higher than that of IgG-DXR or BSA-DXR. Moreover, the survival rates after exposure to the drug for 4 h were comparable among aCD147ab-DXR, IgG-DXR and BSA-DXR. The survival rate was 54, 90 and 89%, respectively, for A431 cells, 66, 86 and 91%, respectively, for Ishikawa cells, 62, 90 and 93% (51, 82 and 86%), respectively, for PC3 cells, and 72, 80 and 82%, respectively, for PC3/CD147-KD cells. Accordingly, the efficacy of anti-cancer drugs targeting CD147 was demonstrated.

Accumulation of aCD147ab-DXR. When PC3 and PC3-KD cells were exposed to 1 μM aCD147ab-DXR for 2 h, the drug accumulated in the cell membrane of PC3 cells. As shown in Fig. 4, aCD147ab-DXR was combined and accumulated clearly at the CD147 discovery plasma membrane, and accumulation in PC3/CD147-KD cells was not observed.

Accumulation of aCD147ab-liposome. After treatment of A431 cells with an aCD147ab-/GFP-double-labeled liposome for 2 h, the cells were stained with Alexa546-labeled mouse IgG (secondary antibody for aCD147ab). The liposomes accumulated in the cells (Fig. 5A), but GFP-liposomes did not (data not shown). Moreover, co-treatment of the cells with the immuno-liposome and excess aCD147ab competitively suppressed accumulation of the liposome (Fig. 5B). Therefore, specific accumulation of the aCD147ab-liposome in CD147-expressing cells was observed.

On the other hand, accumulation of the liposome in PC3 cells was observed by GFP-fluorescence. However, a small amount of the aCD147ab-liposome accumulated in PC3/CD147-KD cells (Fig. 5C and D).

Cytotoxicity of aCD147ab-liposome against A431, Ishikawa, PC3 and PC3/CD147-KD. Cytotoxicity of GSH-DXR-encapsulated immuno-liposome against A431, Ishikawa, PC3 and PC3/CD147-KD cells treated for 4 h was measured after 96 h. The cell death rate for the aCD147ab-liposome/GSH-DXR (0.1 μM) and mouse IgG-liposomes/GSH-DXR (0.1 μM)
was 46 and 18%, respectively, for A431, 37 and 20%, respectively, for Ishikawa, 38 and 17%, respectively, for PC3, and 18 and 16%, respectively, for PC3/CD147-KD (Fig. 6). On the other hand, little cytotoxic effect on these cells treated with aCD147ab-liposome without encapsulated-GSH-DXR was observed (less than 2.5%). Moreover, leakage of GSH-DXR from aCD147ab-liposome/GSH-DXR to culture medium for 4 h-incubation was found in less than 2% of liposomal GSH-DXR (data not shown). These findings suggest that cytotoxicity of the aCD147ab-liposome/GSH-DXR was specifically related to the expression of CD147.

Discussion

Immunofluorescent study revealed a high level of CD147 protein expression in various carcinoma cells, but not in normal tissues. When taking into consideration that an anti-cancer drug acts at an early stage, it was predicted that CD147 would have sufficient potential for use in target therapies.

No significant difference in cytotoxicity was observed as a result of treating the cells with aCD147ab-DXR for 96 h compared with IgG-DXR or BSA-DXR. Nevertheless, even if the cells were exposed to aCD147ab-DXR for a short period of time (2-6 h), potent cytotoxicity of aCD147ab-DXR against carcinoma cells was observed compared with that of DXR. Moreover, it was demonstrated that the effect of aCD147ab-DXR on CD147-expressing cells specifically accumulated and exhibited potent cytotoxicity after only 2 h of exposure (Figs. 4-6). The CD147-negative normal cells did not show accumulation, and only weak cytotoxicity was observed against these cells. This anti-cancer drug targeting CD147 was shown to be effective even after short-term exposure. However, it was suggested that the same degree of cytotoxicity against all conjugates for 96 h exposure was caused by non-specific endocytotic uptake of drugs in 2-dimentional culture system.

Various physiological processes might be influenced by anti-EMMPRIN therapy (12), such as proliferation and differentiation of epithelial cells (20-22), fertilization (23), differentiation and activation of immune cells (24-26), integrin-mediated adhesion of myocytes of the left heart ventricle to basement membrane components (27), selective transport processes in endothelial cells to maintain the blood-brain-barrier function (28), erythrocyte maturation (29) or wound
healing (30). Accordingly, the side effects of target therapy using CD147 conjugates should be evaluated carefully.

We demonstrated in the previous study that GSH-DXR exhibited potent cytotoxicity compared to DXR (17-19). Here, the effect of drug-delivery of GSH-DXR on target chemotherapy was examined using the aCD147ab-liposome. The aCD147ab-liposome specifically accumulated in CD147-expressing cells, but not in CD147-deficient PC3/CD147-KD cells. Moreover, the aCD147ab-liposome/GSH-DXR exhibited potent cytotoxicity against CD147-expressing cells. The drug was induced apoptotic cell death against these cells through caspase-3 activation (data not shown). We also demonstrated that GSH-DXR reversed the response of MDR cells in which AH66/DR cells over-expressed Pgp and showed resistance to 10 μM DXR (15-17). Therefore, it was predicted that target chemotherapy using GSH-DXR was effective on both drug-sensitive and -resistant tumor cells.

In this study, we investigated the therapeutic efficacy of an aCD147ab-DXR conjugate for the treatment of several cancer cell lines. Moreover, the specific cytotoxicity of GSH-DXR encapsulated in an aCD147ab-liposome against these carcinoma cells was also examined. Further study will attempt to confirm specific cytotoxicity in a three-dimensional cell culture system using the Radial-flow Bio-reactor.

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