



DESCRIPTION

α MFc-NC-MMAF is anti-mouse IgG Fc specific antibody conjugated to monomethyl auristatin F (MMAF) with a non-cleavable linker. The antibody portion is a polyclonal antibody which is specific to the Fc region of mouse IgGs. Monomethyl auristatin F (MMAF) is a cytotoxic small molecule which inhibits cell division by blocking the polymerization of tubulin. The non-cleavable linker connecting MMAF to the antibody is stable in extracellular fluid, but can be cleaved by unspecified mechanisms upon entering cells.

APPLICATIONS

Antibody-drug conjugates (ADCs), which have become a new targeted therapy against cancer, consist of an antibody linked to a cytotoxic drug. The ADCs bind selectively to the target cancer cells via the monoclonal antibody portion. Internalization of the ADCs releases the drug to do its damage. Prior to testing the function of ADCs in cell-based assays, each monoclonal antibody is typically conjugated directly with a cytotoxic drug. This step is time consuming and expensive, requiring milligram quantities of purified antibody, separate conjugation of each antibody, and further isolation of the ADC from the unconjugated drug. Using secondary antibody-drug conjugates (2^oADC) in a cell-based cytotoxic assay is a quick and economical alternative to pre-screening monoclonal antibodies as ADC candidates against tumor cells. Instead of conjugating the monoclonal antibody with a cytotoxic drug, the naked monoclonal antibody is added directly to the cells in the presence of the 2^oADC. Internalization of the monoclonal antibody/2^oADC complex can achieve a similar effect of targeted drug release within the cells as the monoclonal antibody-drug conjugate. Furthermore, the 2^oADC can also be applied to screen protein ligands for receptor-mediated cell targeting.

α MFc-NC-MMAF is a 2^oADC for pre-screening antibodies with a mouse IgG Fc moiety or recombinant mouse IgG Fc fusion proteins to determine their cytotoxicity as MMAF bioconjugates. **α MFc-NC-MMAF** alone displays no obvious toxicity against multiple cell lines at 2 μ g/ml (13 nM) or lower concentration. When applied in combination with tumor specific monoclonal antibodies or recombinant Fc fusion proteins, **α MFc-NC-MMAF** can help determine the cytotoxic potential for these antibodies or proteins against target cell lines.

EXAMPLE DATA

It has been demonstrated that Herceptin-DM1 conjugates (T-DM1) displayed potent killing activity against Her2-overexpressing tumor cells but not normal Her2 expression or Her2 negative cells¹. Here cytotoxicity of a high affinity mouse anti-human Her2 (α HER2) mAb was tested in four breast cancer tumor cell lines expressing different amount of Her2 marker. SKBR3 and HCC1954 are Her2 overexpressing cell lines, MCF7 has normal Her2 expression, and MDA-MB468 is Her2 negative. *In vitro* the growth of SKBR3 is slightly stimulated by the α HER2 mAb, while HCC1954, MCF7 and MDA-MB468 are not affected by the unconjugated α HER2 mAb (data not shown). The unconjugated anti-mouse IgG Fc antibody (α MFc) does not change the apparent effect of the α HER2 mAb against these cell lines (Fig A). In contrast, in the presence of 6.6 nM **α MFc-NC-MMAF**, the α HER2 mAb displays potent cytotoxicity against Her2-overexpressing SKBR3 and HCC1954 cells, while shows no killing of MCF7 or MDA-MB468 cells (Fig B).

Fig A. Cytotoxic Profile of Mouse α HER2 mAb in the Presence of Unconjugated α MFc Secondary IgG

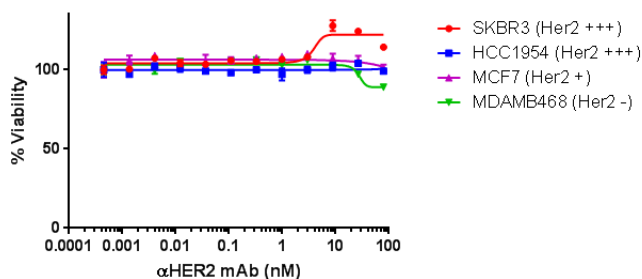
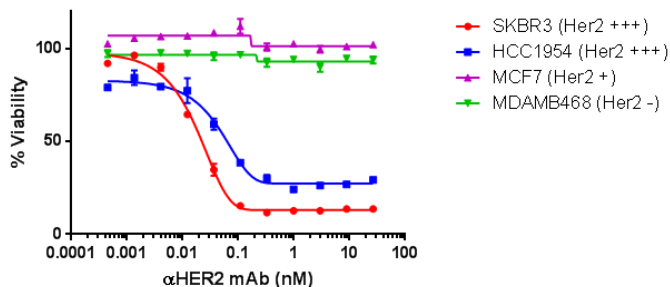


Fig B. Cytotoxic Profile of α HER2 mAb in the Presence of 6.6 nM α MFc-CL-MMAF



STORAGE

Store in -20°C or -70°C manual defrosted freezer. Avoid repeated freeze-thaw cycle.

REFERENCE

1. Lewis Phillips GD, et al. Targeting Her2-positive breast cancer with trastuzumab-DM1, an antibody-cytotoxic drug conjugate. (2008), Cancer Res, **68(22)**, page 9280-90.