#4560 Anti-tumor Activity of a Fully Human anti-CD27 Monoclonal Antibody in a Transgenic Mouse Model

Lawrence J. Thomas, Li-Zhen He, Eric M. Forsberg, Laura Vitale, James M. Boyer, Kristen L. Jones, Naseem Prostak, Jenifer Widger, Andrea Crocker, Jeff Weidlick, Martin J. Glennie¹, Tibor Keler, and Henry C. Marsh Celldex Therapeutics, Inc., Needham, MA and Phillipsburg, NJ, ¹University of Southampton School of Medicine, Southampton, UK

CD27 AS A TARGET FOR IMMUNOTHERAPY

Antibodies that recognize immune cell surface molecules can be used to enhance or target immune responses against tumors. These include antibodies that activate antigen presenting cells (e.g. anti-CD40), antibodies that block immune checkpoints (e.g. anti-CTLA-4, anti-PD-1), and T cell co-stimulatory antibodies (e.g. anti-4-1BB). The costimulatory molecule CD27 is a member of the tumor necrosis factor (TNF) receptor superfamily, and is constitutively expressed on the majority of mature T cells, memory B cells, and a portion of natural killer (NK) cells. The interaction of CD27 with its ligand CD70 plays key roles in the following processes:

- Costimulation through CD27 on T cells causes activation. proliferation, survival, and maturation of effector capacity and memory:
- Costimulation through CD27 on human B cells activates and promotes the generation of plasma cells, proliferation, and the production of immunoglobulin;
- Costimulation through CD27 on NK cells induces cytolytic activity.

In addition to the immune enhancing properties of agonist anti-CD27 mAbs, CD27-targeting antibodies may also provide direct therapeutic effects against tumors with CD27 expression. CD27 expression is well documented on a variety of lymphomas and leukemias and we have previously demonstrated significant antitumor activity in xenograph models using human lymphoblastoid cell lines.

Finally, agonist anti-mouse CD27 mAbs have shown significant anti-tumor efficacy in murine tumor models (French, RR et al. (2007) Blood, 109, 4810-4815; Sakanishi, T et al. (2010) Biochem Biophys Res Commun, 393, 829-835; Roberts, DJ, et al. (2010) J Immunotherapy 33:769-779).

HUMAN ANTI-HUMAN CD27 ANTIBODIES were generated using human Ig Tg-mice.

KD(M

4.02E-10

1.58E-10

3.58E-10

5.56E-11

1 53E-12

1.86E-10

2 02E-10

8 41E-11

CD70-FITC

cell lines, and blocking of sCD70 binding.

Flow cytometric analysis of 1F5 binding to human lymphoblastoid

Biacore - Binding affinity was performed by surface plasmon resonance

analysis using a CD27coated-CM5 sensor chip. Blocking studies were

Human mAb 1F5 is lead development candidate

performed by flow cytometry (below) or ELISA with labeled sCD70

and expressed in CHO cells, as below.

Clone

1G5

1H8

3H12

3H8

269

3410

2C2

Ramos Celli

Anti-hu IgG -

1F5*

Hybridomas expressing human anti-human CD27 mAbs

· Eight selected antibodies have been cloned, sequenced

sCD70 blocking

(Elow and ELISA

Partial

ves

ves

No

No

yes

No

No

Human CD27 Transgenic mice

- · A murine model was established to test the activity of anti-human CD27 HuMAbs.
- · BAC clone, containing the CD27 gene, was used for microiniection of mouse embryos.
- A huCD27-expressing strain was established that showed appropriate expression and regulation of human CD27.
- The huCD27 To mice have been backcrossed to C57BL/6, BALB/c, and C3H mouse strains.
- These mice retain functional mouse CD27.



HuCD27 Tg mice were injected i.v. with 5 mg of ovalburnin on day 0 and 0.25 mg of 1F5 or anti-Mouse CD27 mAb AT-124 on day 0 and 1. On day 7, spleen was harvested and analyzed for CD8+ T cell reactivity to the OVA SIINFEKL peptide (OVA peptide 257-264) by tetramer (A) and IFNy ELISPOT(B).

1F5 Combined with TCR Activation Induces Proliferation and Cvtokine Production from T cells in vitro

1F5 ACTIVATES T CELLS FROM HUMAN CD27 TRANSGENIC MICE



T cells were purified from spleen of hCD27-Tg mice by negative selection with beads. Cells were labeled with CFSE and incubated with antibodies for 3 days. The cross-linking anti-human IoG was passed through an endotoxin removal column before use. The concentration of 1F5 was 0.2 ug/ml . TNFα-ICS showed the same pattern as IFN:

EFFICACY OF 1F5 IN BCL, SYNGENEIC TUMOR MODEL Saline





Groups of 9-10 hCD27 Tg mice (Balb/c background) were challenged with 107 BCL, B-lymphoma cells administered intravenously On Day 0. Animals were then treated with 5 doses of anti-human CD27 mAb 1F5 as indicated. Animals were euthanized upon reaching defined clinical criteria as approved by Celldex IACUC.

SUMMARY AND FUTURE DIRECTIONS

- CD27 is a critical molecule in regulating immunity and tolerance, and is expressed at high level by hematologic malignancies thus representing a new opportunity as an immunotherapy target.
- . We have developed and characterized a fully human monoclonal antibody specific for CD27:
- mAb 1F5 has potent anti-tumor activity in a syngeneic tumor challenge model of the BCL₁ B Cell lymphoma, and
- mAb 1F5 was well tolerated at 1-10 mg/kg i.v. without signs of inflammation or significant lymphocyte depletion.
- Taken together the data support the therapeutic potential of mAb 1F5 (CDX-1127) in cancer therapy.
- A phase 1 clinical trial in hematologic and solid cancers is planned to initiate in 2011.

The authors would like to thank A. Altabef, K. Borrelli Christopher, L. Gergel, T. O'Neill, C. Pilsmaker and R. Weaver for contributions to this work



1F5 BINDING TO MACAQUE CD27

8 '

0.001

Cynomolgus macagues were established as a relevant model for testing 1F5.

- Similar binding to recombinant CD27 from human and macaque Similar distribution of 1F5 binding
- to peripheral blood cells. Similar tissue cross-reactivity by

immunohistochemistry (not shown).

0.01 Various concentrations of purified monkey CD27 (green line) or humar CD27 (red line) were captured to ELISA plates with anti-Flag antibody followed by incubation with 1F5 mAb. A goat anti-human IgG Fc-HRF antibody and substrate Super Blue TMB were used for detection

Monkey CD

0.1 1 10

1F5 concentration, µg/ml

| Analysis | CD4+ T cells | | CD8+T cells | | B cells (CD20+) | | NK cells | |
|----------------------|--------------|---------|-------------|--------|-----------------|----------|----------|----------|
| | human | monkey | human | monkey | human | monkey | human | monkey |
| % CD27+ ^b | 84 ± 5 | 81±1 | 70 ± 12 | 90 ± 1 | 37 ± 4 | 15 ± 1 | 11 ± 4 | 88 ± 6 |
| MFI ^c | 1517± 123 | 416± 14 | 1415±153 | 519±11 | 893±101 | 491± 113 | 667±28 | 1050± 42 |

PBMCs were isolated from whole blood of 3 human and 3 cynomolaus macaques. The cells were stained with 1F5 mAb together with markers to delineate the major T cell and B cell populations that express CD27. The table summarizes the mean ± standard error of results for human and monkey cells with respect to the percent of cells expressing CD27 and the intensity of expression (MFI).

IF5 IS WELL TOLERATED AND NON-DEPLETING IN PILOT NON-HUMAN PRIMATE STUDY

- 1. 3 or 10 mg/kg of anti-human CD27 mAb 1F5.



Total lymphocytes (based on side and forward scatter size) memory B cells (CD20+ and CD95 bright) and monocytes (based on side and forward scatter size) were stained with anti-human IoG antibody (bold line) and compared to unstained controls (shaded histogram).

1F5 does not Significantly Deplete Circulating Lymphocytes after a Single 1 - 10 mg/kg Dose



Lymphocytes were stained with subset markers and the % positive cells plotted vs time for each animal treated at the different doses = 1 ma/ka : • = 3 ma/ka: ▲ = 10 ma/ka

Copyright Celldex Therapeutics Inc.

- 3 cynomolgus monkeys were treated with one i.v. dose of
- - No clinical symptoms.
 - · No elevation in body temperatures and no detectable
 - · Flow was performed on blood drawn at indicated times.
 - 1F5 on Circulating Lymphocytes after a Single 1 mg/kg Dose





- levels of TNF-α, IL-6, or IL-1β.



1F5 Enhances Antigen Specific T cell Response in vivo