# Assessment of therapeutic antibody efficacy without the interference of murine Fc receptors allows for investigation of human antibody-dependent cellular cytotoxicity mediated by NK cells in the FcResolv™ hIL-15 NOG mouse model

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#2836

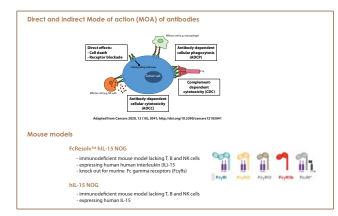
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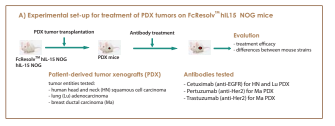
### Background

Targeted antibody therapy is applied to treat various cancer types. In addition to the primary mode of action (MOA), which involves direct binding to the tumor antigen, indirect MOA acting through the constant region (Fc) of the antibody can enhance anti-tumor efficacy. Indirect mechanisms engage the innate immune system, mediated by both the complement system (complement-dependent cytotoxicity (CDC)) and immune cells (antibodydependent cellular phagocytosis (ADCP) and antibody-dependent cellular cytotoxicity (ADCC). These indirect mechanisms can complicate the evaluation and accurate assessment of antibody-induced ADCC by human NK cells in current mouse models. In immune-deficient mouse strains (e.g. NOG), false positives and/or negatives may occur due to interactions with murine Fc receptors. These can either result in anti-tumor responses via activation of the murine innate immune system or can interfere with the human-targeted therapy's primary MOA. To study the response to anti-cancer antibodies without the interference of these murine Fc receptor interactions and to investigate ADCC mediated by human NK cells, a novel mouse model deficient in Fc receptors and expressing human IL-15 (FcReslov<sup>TM</sup>)-IL-15 NOC) was employed for testing antibody therapies.



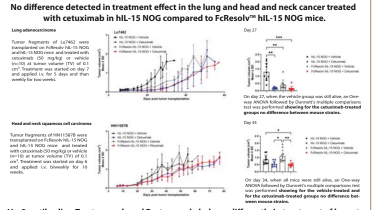
# **Methods**

Patient-derived xenograft (PDX) tumor models were transplanted into hiL-15 NOG and FResolv<sup>™</sup> hiL-15 NOG mice. A human head and neck squamous cell carcinoma and a lung adenocarcinoma PDX model were both treated with cetuximab. Treatment with pertuzumab and trastuzumab was applied in a breast ductal carcinoma PDX model. Based on growth kinetics, the lung cancer PDX model was chosen for further testing of ADCC in the NK cell-humanized FCResolv<sup>™</sup> hiL-15 NOG mouse.

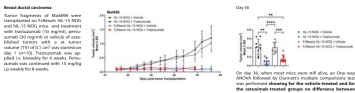


#### Results

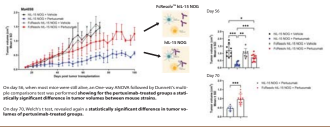
### A) Treatment of PDX tumors on FcResolv<sup>™</sup> hIL-15 NOG mouse model



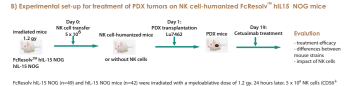
#### Her2-antibodies, Trastuzumab and Pertuzumab, behave differently in treatment of breast carcinoma in hIL-15 NOG and FcResolv™ hIL-15 NOG mice.



#### False-positive efficacy of pertuzumab treatment in hIL-15 NOG compared to FcResolv™ hIL-15 NOG mice.

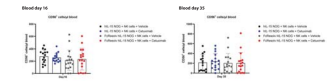


## B) Treatment of PDX tumor on NK cell-humanized FcResolv™ hIL-15 NOG mouse model



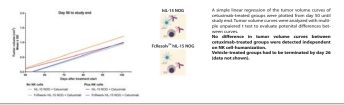
FcResolv hiL15 NOG (n=49 and hL15 NOG mice) rene2] were irradiated with a myeloablative dose of 12 y2 Ahours later, Ya Neols (CD65\* McS-sorted from PMCs of health blood donors) were injected ix. In FcResols hiL15 NOG (n=43) and hL125 NOB (n=63). The Not calls, tumor fragments with L2462 were transplanted on all piled ix. Or day 19, a stratified randomization of tumor volumes was performed and treatment started. Treatment with curvals (SO mg/ka) and all piled ix. Or 6 days and than weekly for two weeks.





Blood was taken on day 16 and day 35 after irradiation to evaluate NK cell engrafitment. Blood was stained for human CD45 and human CD56. Cell count per µL blood are depicted. No difference of NK cell counts in blood was detected between groups tested by One-way ANOVA and Sidak's multiple com parisons test

No difference detected in treatment effect in the lung cancer treated with cetuximab in NK cell-humanized hIL-15 NOG compared to NK cell-humanized FcResolv<sup>™</sup> hIL-15 NOG mice and no NK cell-dependent treatment effect observed in this tumor model.



# Conclusions

There was no difference in percent tumor growth inhibition between the FcResolv<sup>™</sup> hlL-15 NOG and hlL-15 NOG mice with regards to cetuximab treatment in the lung and head and neck cancer of for trastuzumab treatment of breast ductal carcinoma. However, pertuzumab treatment revealed a false positive efficacy, with the false positive effect more pronounced in hlL-15 NOG mice than in FcResolv<sup>™</sup> hlL-15 NOG mice. These results demonstrate that FcResolv<sup>™</sup> hlL-15 NOG mice serve as a suitable mouse model for a more accurate assessment of the therapeutic efficacy of anti-tumor antibodies. Additionally, evaluation of human-mediated ADCC of therapeutic antibodies in NK cell-humanized FcResolv<sup>™</sup> hlL-15 NOG allows detection of effects specifically mediated by human NK cells.



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