# Potency-reduced IL12 heterodimeric Fc-fusions exhibit strong anti-tumor activity

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# Introduction

- Interleukin-12 (IL12) is a heterodimeric proinflammatory cytokine produced by activated antigen-presenting cells that induces differentiation of Th1 cells and increased proliferation and cytotoxicity of T and NK cells.
- Stimulation of these cells by IL12 leads to production of high levels of IFNγ. These immunestimulating aspects of IL12 are promising for cancer treatment and may help to convert immunologically suppressed "cold" tumors into inflamed "hot" tumors.
- Preclinical studies in mice revealed that IL12 can have a dramatic effect on shrinking syngeneic tumors, however clinical studies in humans have resulted in severe toxicity and a small therapeutic window, limiting response rates.
- Prior work at Xencor demonstrated that reduced-potency IL15/IL15Rα-Fc fusion proteins exhibited superior pharmacokinetics, pharmacodynamics, and safety in non-human primates through reduction of receptor-mediated clearance. Applying similar principles to IL12, we created IL12 heterodimeric Fc-fusions (IL12-Fc) with reduced potency in order to improve tolerability, slow receptor-mediated clearance, and prolong half-life.



# **1. Potency-reduced IL12-Fc are engineered for optimal** activity and extended serum half-life



**Figure 2.** Schematic of IL12-Fc heterodimers and analytical characterization. Monovalent IL12 p35/p40 is attached to Xencor's well-validated heterodimeric Fc domain. The IL12 has been engineered for decreased binding to its receptors in order to reduce potency and the Fc domain is modified to eliminate FcyR interactions. The Fc domain may also be modified with Xtend<sup>™</sup> Fc technology to promote longer half-life. IL12-Fc heterodimers can be produced in high yields and are purified using standard methods (protein A and IEX chromatography).

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## 2. IL12-Fc with up to 10,000-fold reduction of in vitro potency were engineered in order to improve therapeutic index

#### In vitro pSTAT4 activity of WT IL12-Fc on human PBMCs



Figure 3. In vitro activity of WT IL12-Fc was assessed on activated human PBMCs by measuring intracellular pSTAT4 by flow cytometry.

Anti-tumor efficacy at day 33

### **Engineering potency-reduced IL12-Fc**



**Figure 4.** Left: A library exploring amino acid substitutions at putative IL12-receptor-interface positions (in red) was created. Right: The library was screened for reductions of in vitro potency by pSTAT4 and optimal variants were selected for further evaluation.

### 3. Potency-reduced IL12-Fc show strong anti-tumor activity and PD response as single-agent and in combination with anti-PD1



# 4. Potency-reduced IL12-Fc have antibody-like PK in mice



Figure 7. The pharmacokinetics (PK) of potency-reduced IL12-Fc-v1 were evaluated in C57BL6/J mice. N = 4 mice were injected IV with 2 mg/kg IL12-Fc-v1 on Day 0 and drug concentration in serum was measured over time. The estimated half-life  $(t_{1/2})$  is approximately 10 days and similar to that of monoclonal antibodies, indicating that IL12-Fc-v1 has a long half-life and favorable stability in the absence of TMDD.

### Tumor volume - Spider plots



### In vitro pSTAT4 and MLR activity of lead IL12-Fc variants



on activated human PBMCs by measuring intracellular pSTAT4 by flow cytometry (left) and IFN $\gamma$ production in a mixed-lymphocyte reaction (MLR) (right).

### In vivo pharmacodynamics (PD)

# Summary

IL12 heterodimeric Fc-fusions were engineered with a potency-reduced IL12 in order to improve tolerability, slow receptor-mediated clearance, and prolong half-life in vivo.

Potency-reduced IL12-Fc demonstrate significant anti-tumor activity concurrent with activation and proliferation of CD8<sup>+</sup>T cells, increased PD1 expression, and increased serum IFN $\gamma$  in mice.

These results support further testing of potency-reduced IL12-Fc as a potential novel cytokine therapy in cancer patients.