23andMe **Fherapeutics**

Phase 1/2a trial of CD200R1 inhibitor 23ME-00610: exploratory analyses of tissue-based and genetic biomarkers

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BACKGROUND

Genetic Signature

• Using the 23andMe database, pleiotropic causal variants with opposing effect on risks for cancer and immune diseases, referred to as an immuno-oncology signature, were observed in 3 components of the CD200R1 pathway, including CD200R1, its ligand CD200 and the signaling adaptor protein DOK2.¹ (Figure 1)

CD200R1

• CD200R1 is expressed on immune cells and binds to CD200, downregulating proinflammatory cytokines by activated T and myeloid cells and promoting an immunosuppressive microenvironment in cancers, where CD200 is highly expressed.¹ (Figure 2)

23ME-00610

- 23ME-00610 is a first-in-class IgG1 antibody that binds CD200R1 with high affinity ($K_n < 0.1 \text{ nM}$) and inhibits immunosuppressive signaling, leading to restoration of T cell activity and killing of CD200-expressing tumor cells in preclinical studies.¹
- 23ME-00610 is being evaluated in a Phase 1/2a clinical trial in patients with advanced solid cancers (NCT05199272) (Figure 3) and has demonstrated acceptable safety and tolerability and anti-tumor activity in a neuroendocrine tumor patient and a renal cell carcinoma patient.²
- For the first time, the exploratory tissue-based biomarker data and genetic analysis are presented.



METHODS



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cell carcinomas.



A. Representative images archival or fresh pre-treatment neuroendocrine neoplasms (NE) with high (b), moderate (d), and no CD200 expression (f) in the tumor cell membrane. Scale bar = 50 µm

Figure 6: Baseline Tumor Immune Contexture: Clinical Benefit May Be More Likely In Immune Low Though More Immune Permissive Tumor Micro-environments (Macrophages < Lymphocytes)



Cancer Stroma Combined

A. Al powered algorithm was applied to H&E images of archival or pre-treatment fresh biopsies from the neuroendocrine cohort to determine spatially defined immune infiltrates (Lunit SCOPE IO).

Tumors characterized by immune low or excluded phenotype might be more likely to benefit from 23ME-00610 treatment Tumor infiltrating lymphocytes (B), and macrophages (C) were quantified in tumor area, tumor stromal area and combined total tumor and stromal areas. Tumors from patients deriving clinical benefit (PR, SD) during 23ME-00610 treatment had fewer infiltrating lymphocytes and macrophages when compared to tumors that progressed at the first radiological tumor assessment (PD)

More immune permissive tumor micro-environments might be more likely to lead to benefit from 23ME-00610 Macrophage to lymphocyte cell density ratios were also calculated **(D)**, with relatively lower macrophage infiltration in relation to /mphocytes associating with clinical benefit.

Stroma Combined

Cancer



Figure 7. Pharmacodynamics: 23ME-00610 Leads to Changes in the Immune **Composition and Immune Activation**



PSMB10 CCL5 IDO1 HLA-E 0.75 **a** 8-0 SD (5) PD (13) SD (5) PD (13) SD (5) PD (13) SD (5) PD (13)

Additionally, a trend in increased expression of interferon inducible genes that may be indicative of activation of cytotoxic CD8+ T cells and NK cells on 23ME-00610 treatment was observed.

Screening

C3D1_C2D9

Bulk RNAseq of pre- and

on-treatment (prior to Cycle 3) tumor

biopsies of neuroendocrine tumors

suggest a trend of increased gene

expression of T cell / NK cell markers

particularly in patients with stable

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Figure 5. Host Genetics: Polygenic Risk Scores May Be Leveraged to Augment the Association of Tumor CD200 Expression With Clinical Benefit (PR/SD) of 23ME-00610 Treatment





- To assess the potential interaction of genetic host immune and tumor parameters on 23ME-00610 efficacy, all Phase 1/2a enrolled patients with genotyping data (n = 81) had ancestry corrected immune-mediated phenotype PRS calculated. The polygenic risk score (PRS) was trained on the research consented 23andME customer database.
- For the hypothyroidism phenotype, there is no difference in median PRS (Z-score transformed using healthy ancestry matched population means), between patients that had a PR or SD vs. PD.
- When segregated based on "high" CD200 tumor expression (H-score > 40), patients with high CD200 who achieved a PR or SD had a higher median (0.6728, ICR = 5.020) genetic risk for hypothyroidism than patients with high CD200 who had PD (median = -0.3707, ICR = 5.568).



All Ph1/2a patients with available CD200 and hypothyroidism



- The combination of CD200 high expression and hypothyroidism PRS > 0 may have a better predictive association with longer time to progression. Univariate
- (time-to-event) analysis using a log rank test was performed. This data might indicate that the immune host genetics might influence the ability of a patient to raise a sufficient anti-tumor immune response against CD200 expression tumors and benefit from CD200R1 blockade.

CONCLUSIONS

- Higher tumor cell expression of CD200 associates with clinical benefit from 23ME-00610 in some patients, and this tumor based putative biomarker might be further augmented if combined with genetic based host immune set point readouts (e.g. hypothyroidism PRS), warranting further exploration of this and other biomarkers for potential future patient selection.
- Neuroendocrine tumors that had tumor shrinkage or prolonged stable disease tended to be less inflamed at baseline potentially due to CD200 mediated immune suppression/exclusion, though these tumors might be more permissive to immune activation due to lower macrophage to lymphocyte ratio.
- Analysis of pre- and on-treatment tumor samples showed an increase in T and NK cell markers and an increase in interferon inducible genes with 23ME-00610 treatment suggesting pharmacodynamic immune modulation.

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