GB004 Exhibits Protective Effects Directly on Epithelial Cells Using Ex Vivo Organoid and Monolayer Cultures

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INTRODUCTION

- Inflammatory bowel disease (IBD) is characterized by a breach in intestinal barrier integrity, allowing influx of luminal antigens and setting up a vicious cycle of inflammation and epithelial injury.
- Despite efficacy of IBD treatment with anti-tumor necrosis factor agents and anti-integrin agents, a large fraction of patients do not respond adequately to currently available therapies or biologics and do not achieve long-term remission.
- GB004 is an oral, gut-targeted, small molecule that stabilizes hypoxia inducible factor (HIF-1α), a key transcription factor involved in the adaptive and protective cellular responses at the intersection of hypoxia and inflammation (Figure 1).
- Preclinical efficacy of GB004 has been demonstrated in mouse models of colitis and correlated with HIF-1a stabilization in colonic epithelial cells, induction of HIF-1α target genes, downmodulation of inflammatory cytokines, and improvement in histologic parameters of barrier function.
- A phase 2 study evaluating two doses of GB004 as a tablet formulation in mild to moderate UC is ongoing (NCT04550383).

OBJECTIVE

- To assess the effects of HIF stabilizer GB004 on gene expression, tight junctions, and barrier integrity using intestinal epithelial cells and organoids.

METHODS

- Human RepliCult differentiated monolayers assays were performed at Atlas Biosystems (Chapel Hill, NC) by proliferating and differentiating human-derived intestinal epithelial cells on a 2D monolayer platform. These monolayers were assessed with GB004 treatment under normal healthy conditions or with cytokine-stimulated conditions (25 ng/mL TNF-α) to induce barrier damage. Barrier integrity was assessed through measuring Transepithelial Endothelial Electric Resistance (TEER) and a barrier integrity assay using FITC-dextran. HIF-1α target genes were assessed in cell lysates and tight junction formation and adhesion molecules were investigated by immunofluorescence staining. Three independent studies were performed to generate data presented in Figure 3, Figure 4, and Figures 5-7, respectively.
- Unless otherwise noted, data are presented as mean ± SD. Statistical analysis was carried out using GraphPad Prism and one-way ANOVA (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).

RESULTS

- Figure 2. GB004 induces HIF-1α dependent gene expression in murine organoids.
- Figure 3. GB004 stabilizes HIF-1α in human differentiated monolayers at 2 hours (A) and 4 hours (B) after treatment depending on dose (dotted line indicates the maximum HIF-1α stabilization measurement with the positive control lysate). Normal human monolayers have complete barrier integrity (C) as determined by TEER.
- Figure 4. TNF-α or IFN-γ stimulation of human differentiated monolayers induces barrier integrity resulting in reduced TEER measurements (A) and increased barrier permeability (B).
- Figure 5. GB004 treatment of TNF-α-stimulated human differentiated monolayers prevents the loss in barrier integrity as measured by TEER, while tofacitinib (JAK inhibitor) had no effect on barrier integrity.
- Figure 6. TNF-α treatment of human differentiated monolayer results in epithelial cell death at 48 hours. GB004 treatment prevents TNF-α-induced cell death, whereas tofacitinib (Tofa) had no effect.
- Figure 7. GB004 modulates a total of 266 genes in human differentiated monolayers stimulated with TNF-α that are related to the HIF-1α, PHD and hypoxia pathways as determined by RNAseq. Genes are grouped into tightly correlated modules and key genes are highlighted with green boxes.

CONCLUSIONS

- GB004 demonstrates direct protective effects on mouse organoid and human-derived differentiated monolayer epithelial cultures.
- GB004 induces HIF-dependent genes, specifically genes that drive barrier integrity, which are critical to mucosal repair in IBD.
- GB004 preserves barrier integrity in human monolayer cultures that have been stimulated with cytokines.
- GB004 prevents TNF-α-induced cell death and preserves epithelial cell survival.
- This data complements data generated in mouse models of colitis demonstrating induction of HIF-1α dependent genes, reduced barrier dysfunction and beneficial efficacy.

REFERENCES


DISCLOSURES

KRTM, SM, GOG, BGL, LC, and UCD are employed by Gossamer Bio, Inc.

Prepared for presentation at Digestive Disease Week (DDW) virtual meeting, May 21-23, 2021.