LOCAL BIOACTIVATION AND EFFICACY OF PALI-2108: A PROMISING PDE4 INHIBITOR PRODRUG FOR ULCERATIVE COLITIS TREATMENT

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Introduction
Phosphodiesterase-4 (PDE4) is a key enzyme in cAMP hydrolysis and its inhibition elevates intracellular cAMP, downregulates inflammatory cytokines, and reduces the expression of cell adhesion molecules, thereby preventing local infiltration and activation of inflammatory cells. Approved PDE4 inhibitors include rufinamide for use in chronic obstructive pulmonary disease (COPD) and apremilast for psoriasis/psoriatic arthritis. Despite the development of subtype specific PDE4 inhibitors, oral administration and systemic distribution has resulted in central nervous system (CNS) toxicity, such as headaches, nausea or vomiting, leading to discontinuation of therapy and limiting potential efficacy. Targeted and better-tolerated oral PDE4 inhibitors hence remain an unmet need in IBD.

We investigated PALI-2108, an orally delivered, intestinally activated PDE4 inhibitor prodrug designed with a D-glucoronic sugar moiety. PALI-2108 minimizes systemic exposure until cleaved by the colonic bacterium enzyme β-glucuronidase, producing active drug PALI-0008, therefore reducing potential CNS toxicity. PALI-2108 proposed mechanism of action is outlined in figure 9.

Methods
Healthy mice were dosed with a single oral dose of PALI-2108 and sacrificed at multiple time points across 24 hours. Proximal colon, distal colon and plasma were collected and PALI-0008 measured (Figure 3). Mice were treated once with 3% oxazolone i.r. to induce colitis, followed by treatment with PALI-2108 BID at 4.2 mg/kg for 3 days. PALI-2108, -0708, and -0008 were measured at 72 hours in plasma, duodenum, ileum and colon (Figure 4). A classic cellular thermal shift assay (CETSA) was developed to assess on-target PDE4 binding within colon tissue homogenates dosed with apremilast, PALI-2108 or Vehicle and detecting changes in thermal stability (Figure 1 and 5). A second acute colitis model of disease induction in mice was utilized with 2% DSS in drinking water administered from D0 to D7. Mice were treated on D1 through D6 with PALI-2108 BID, PALI-2108 1.5 mg/kg, tacrolimus and apremilast were administered OD. Mice were sacrificed on day 7 (Figure 2). Disease activity index (DAI) score was assessed based on a scoring system including body weight loss, stool consistency and the presence of blood in feces in DSS acute colitis mouse model. Treatment with PALI-2108 at 21, 4 and 0.75 mg/kg BID and 1.5 mg/kg daily significantly decreased the mean DAI score compared to DSS control.

Results
Figure 3. PALI-0008 levels in healthy mice dosed PO with PALI-2108. PALI-0008 is favorably distributed to the colon as indicated by a colon/plasma AUC ratio of >200 and the bioavailability of PALI-0008 when delivered as the prodrug PALI-2108 is only 4%.

Figure 4. Concentrations of PALI-2108 (prodrug), PALI-0708 (inactive intermediate) and PALI-0008 (active) where measured at 72h upon dosing PALI-2108 BID at 4.2 mg/kg for 3 days in mice treated once with 3% oxazolone i.r. Study demonstrated colon biodistribution of PALI-2108 to PALI-0008 and negligible plasma levels of all forms.

Figure 5. Mouse colon homogenate with 10 µM of apremilast resulted in a stable assay for PDE4D where the target protein was thermally stabilized by the compound. Target engagement was validated in tissue from animals that were in vivo dosed with apremilast, PALI-2108 or Vehicle controls. For both compounds a thermal stabilization effect on PDE4D was observed confirming target engagement in vivo.

Figure 6. Disease activity index (DAI) score was assessed based on a scoring system including body weight loss, stool consistency and the presence of blood in feces in DSS acute colitis mouse model. Treatment with PALI-2108 at 21, 4 and 0.75 mg/kg BID and 1.5 mg/kg daily significantly decreased the mean DAI score compared to DSS control.

Figure 7. Treatment with PALI-2108 at 21, 4 and 0.75 mg/kg BID and 1.5 mg/kg decreased the mean body weight loss score for DAI compared to DSS Vehicle control group in DSS mouse model.

Figure 8. PO administration of produg PALI-2108 at 43 mg/kg was well tolerated, while PO administration of PALI-0008 resulted in emesis at 1 and 3 mg/kg, suggesting systemic toxicity only with administration of the active compound and not the prodrug.

Figure 9. PALI-2108 mechanism of action. PDE4 is highly expressed in immune and proinflammatory cells. The inhibition of PDE4 prevents transcription factors, such as NF-κB, from inducing the expression of various proinflammatory mediators involved in IBD, such as TNF-α and various interleukins. The PDE4 inhibitor PALI-2108 is orally delivered and colon activated allowing for local activity with low systemic exposure.

Conclusions
PDE4 inhibitor pro-drug, PALI-2108 shows:
• Local bioactivation
• Colon-specific distribution
• Similar target engagement to other PDE4 inhibitors
• Dose-dependent efficacy in a mouse UC model
• No systemic toxicity in dog and large therapeutic window due to local activation

PALI-2108 is a promising novel therapy for UC, with localized bioactivation, expanded therapeutic window, and potent PDE4 inhibitory activity. PALI-2108 is in development for moderate-to-severe UC and is advancing toward regulatory filing for first-in-human studies.

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