Background

There are few attractive options for treatment of checkpoint inhibitor (CPI)-refractory or -relapsed metastatic melanoma (MeL). Cytotoxic chemotherapy is ineffective. In a recent, multicenter, randomized trial involving ipilimumab provided a response rate of 5.7% and mPFS > 11 mos. in previously untreated patients (pts); however, this combination resulted in high rates of severe adverse effects including immune-related adverse effects.1

The only treatment options for advanced triple-negative (ER/PR/HER2-) breast cancer (TNBC) are cytotoxic chemotherapy regimens which are poorly effective and highly toxic. CPIs, including pembrolizumab, have been studied in pts with advanced TNBC with modest effectiveness. In the KEYNOTE-012 trial, the overall response rate was 27% evaluable cases.2

Less toxic approaches to therapy of CPI-refractory or -relapsed advanced MeL are needed as are improved treatment options for refractory and relapsed TNBC.

Rationale

Imprime PGG is a yeast-derived soluble beta-glucan that acts as a pathogen-associated molecular pattern to stimulate innate immune cells and coordinates a multi-faceted immune response in the tumor microenvironment.3

• Generates immunomodulatory cytokines and chemokines
• Repolarizes M2 to M1 phenotype of macrophages
• Enhances dendritic cell activation

Imprime PGG:4

• Effectively pairs with anti-PD-1 and PD-L1 antibodies to induce immunity against systemic tumors in mouse models4
• Is complexed by naturally occurring IgG anti-beta-glucan antibodies (ABA) which are required for its activity: serum ABA level is a potential predictive biomarker for Imprime PGG activity2
• Has been well-tolerated in clinical trials in healthy volunteers and cancer pts1,5

Pembrolizumab (KEYTRUDA®) is a humanized mAb programmed death receptor (PD-1) approved for advanced MeL advanced NSCLC, recurrent or metastatic head and neck cancer, and classical Hodgkin Lymphoma.

Imprime PGE is a multicenter, open-label, Phase 2 clinical trial of Imprime PGG (PGG beta-glucan) in combination with pembrolizumab in pts with advanced MeL failing front-line treatment with CPIs or TNBC failing front-line chemotherapy for metastatic disease. This trial is registered with Clinicaltrials.gov: NCT02981303 and is enrolling investigational sites located in the United States.

Hypothesis

Imprime PGG in combination with pembrolizumab will restore (for MeL) or enhance (for TNBC) sensitivity to checkpoint inhibitors (CPI) by appropriate and effective stimulation of the pts’ innate and adaptive immune systems in those pts who have failed first-line therapy.

Study Design

Seventy-one (71) pts will be enrolled into the study, 29 pts with advanced MeL and 42 pts with TNBC.

The study design is based on hypothesis testing of the ORR for pembrolizumab + Imprime PGG.

• MeL: null hypothesis of ORR ≤ 5% vs. alternative hypothesis of ORR ≥ 23%
• TNBC: null hypothesis of ORR ≤ 15% vs. alternative hypothesis of ORR ≥ 40%

The study incorporates Simon’s optimal 2-stage design with sample size fixed at 12 pts for each indication in Stage 1, while Stage 2 will enroll an additional 17 MeL pts and 30 TNBC pts.

Type I error of 0.05 and 90% power for both MeL and TNBC

• At Stage 1, in the event there are a total of 0 (or < 33%) of pts with Grade 3/4 adverse events in Cycle 1, Stage 1 will be repeated at a reduced dose of 2 mg/kg with an additional cohort of 12 pts in each arm
• pts was 16.5% 5

• ≤ 4 (or ≤ 5%) pts with Grade 3/4 adverse events in Cycle 1 within either tumor type
• ≤ 1 at least 1 response in MeL pts and 2 responses in TNBC pts
• Either arm may proceed independently
• Rejection of the null hypothesis will require documenting at least 4 objective responses in MeL and 13 objective responses in TNBC at the conclusion of Stage 2.

Study Design Cont.

Study Schema

N = 73 patients MeL ≥ 29 TNBC ≥ 42

Stage 0 = 24 pts (12 MeL + 12 TNBC)

Stage 1 = 24 pts (12 MeL + 12 TNBC)

Imprime PGG 4 mg/m² COV Renublox 200 mg COV

Repeat Stage 1

with Imprime PGG 3 mg/m² COV

Stage 2 = 47 pts (17 MeL + 30 TNBC)

A或多 repeat Stage 1 depending on treatment response

STOP

*Arms may proceed independently into Stage 2 OR, overall response

Key Study Endpoints

Primary Efficacy Endpoint

• Overall Response Rate (ORR) in pts with advanced MeL or metastatic TNBC

Secondary Efficacy Endpoint

• Time to response (TTR)
• Complete response rate (CR)
• Duration of overall response (DOR)
• Progression-Free Survival (PFS) and PFS rate at 6 months and 1 year
• Overall survival (OS) and OS rate at 1 year

Safety Endpoints

Incidence of treatment-emergent adverse events and laboratory abnormalities

Pharmacokinetic Endpoints

• Profile pharmacokinetic (PK) data

Exploratory Immune Correlative Objectives

• Determine ORR and PFS based on RECIST
• Correlate the following with OR and treatment outcome:
  • Levels of baseline serum anti-β-glucan antibody (ABA)
  • Changes in immune cell activation markers in tumor biopsy samples and peripheral blood immune cells
  • Changes in the tumor immune microenvironment including tumor-infiltrating lymphocytes and tumor-infiltrating myeloid cells
  • PD-L1 expression in tumor biopsy samples (in tumor and myeloid cells)

Key Eligibility Criteria

Adults who have:

• MeL: histologically or cytologically confirmed diagnosis of unresectable Stage III or metastatic (Stage IV) MeL not amenable to local therapy and irrespective of PD-L1 status
• TNBC: histologically or cytologically confirmed diagnosis of metastatic (Stage IV) TNBC irrespective of PD-L1 status

• MeL: TPS3105
• TNBC: TPS3105

Key Eligibility Criteria Cont.

• Documented objective radiographic or clinical disease progression after:
  • MeL: PD-L1/PD-L1 +/- anti-CTLA-4 inhibitor therapy
  • TNBC: 1 or more prior lines of chemotherapy for metastatic disease*

• At least one radiologically measurable lesion per RECIST v1.1 imaged by CT scan or MRI within 28 days of study treatment
• Screening peripheral blood levels of IgG anti-β-glucan antibody (ABA) of ≥ 20 mcg/mL
• Criterions to consider providing fresh tissue for biomarker analysis and/or assessment

Excluded:

• Pts who had a prior anti-cancer monoclonal antibody (except immune CPI in the case of MeL pts) within 30 days prior to start of study treatment, and did not respond to CTCAE Grade 1 or better from adverse events related to prior therapies
• TNBC pts who received prior therapy with an anti-PD-1, anti-CTLA-4, or anti-PD-L2 agent
• Eastern Cooperative Oncology Group (ECOG) performance status of 0 or 1
• Adequate organ function
• Documented informed consent
• Life Expectancy of ≥3 months
• Inclusion criteria language pending approval of a protocol amendment: current protocol language reads, “after at least 1 line of chemotherapy for metastatic disease”

General Statistical Methods

Interim Analysis

• At the end of Stage 1 for each tumor type
• Early stopping for futility, not efficacy

Efficacy Analysis

• ORR and CRR: point estimate along with 95% CI
• PFS, OS, DOR and TTR: descriptive summary using Kaplan-Meier method

Safety Analysis

• Summaries of adverse events, hematologic and clinical chemistry values, ECOG performance status

Exploratory Analysis

• Correlation between biomarker data with ORR and other clinical endpoints

Study Overview

A Trial Steering Committee will provide study oversight.

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Corresponding Author

ruhni@biothera.com

References


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Cancer Immunotherapy Clinical Trials

Imprime PGG for the treatment of advanced melanoma and breast cancer