Phase 2 trial of CA-170, a novel oral small molecule dual inhibitor of immune checkpoints VISTA and PD-1, in patients with advanced solid tumor and Hodgkin lymphoma


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Introduction

V-domain Ig suppressor of T-cell activation (VISTA) and Programmed-death 1 (PD-1) are independent immune checkpoints implicated in various malignancies. Preclinical studies have demonstrated that dual blockade of these pathways is synergistic. CA-170 is a first-in-class oral small molecule that directly targets both VISTA and PD-1/PD-L1 pathways and has shown anti-tumor activity in multiple preclinical models. A Phase 1 dose escalation study (Clinicaltrials.gov NCT02812875) has shown acceptable safety of CA-170 with dose escalated up to 1200 mg BID.

Methods

The Phase II study is a multi-tumor (Head & Neck Squamous Cell Cancer [HNSCC], Non-Small Cell Lung Cancer [NSCLC], MSI-H positive solid tumors and Classical Hodgkin Lymphoma [HL]) trial investigating two dosages (400 mg versus 800 mg) of CA-170. [Clinical Trials Registry of India (CTRI) registration no. - CTRI/2017/12/011026]

Key eligibility criteria include: age ≥ 18 years, ECOG ≤1, adequate organ function, no previous exposure to immuno-oncology agents, and 1-3 prior lines of systemic therapy. Primary endpoint is Response Rates by RECIST 1.1 for solid tumors and by IWG Revised Response Criteria for Classical Hodgkin Lymphoma. Additional endpoints include immune-related Response Criteria (irRC) for solid tumors, Clinical benefit Rate (CBR) - defined as Stable Disease or better, as well as safety and Pharmaeconomics. Analysis for Hodgkin Lymphoma was also done by Lugano criteria. CA-170 was given once daily till progression of the disease or intolerable toxicities. Patients with solid tumors were followed with contrast enhanced CT scans and those with Hodgkin lymphoma with PET/CT.

Results

Enrolment started in February 2018, and as of Oct 15, 2018, a total of 62 patients have been enrolled. The tumor type distribution and baseline demographics of patients is provided in Tables 1 and 2.

Table 1: Tumor distribution

<table>
<thead>
<tr>
<th>Tumor Type</th>
<th>No. of Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>HNSCC</td>
<td>20</td>
</tr>
<tr>
<td>NSCLC</td>
<td>24</td>
</tr>
<tr>
<td>MSI-H positive solid tumors</td>
<td>5</td>
</tr>
<tr>
<td>Classical Hodgkin Lymphoma</td>
<td>13</td>
</tr>
</tbody>
</table>

Baseline demographics were similar in the 400 and 800 mg groups

Efficacy Evaluation

Efficacy evaluation is done in “Evaluative Population” - defined as patients who have had at least one follow up scan during study or were withdrawn due to clinical progression before the initial follow up scan. As of Oct. 15, 2018, 37 patients constitute the “Evaluable Population”. Table 3 and Figure 1 provide efficacy evaluation from these patients.

Table 3: Clinical Benefit Rate (CBR) by irRC in solid tumors and by Lugano criterion for Hodgkin lymphoma

<table>
<thead>
<tr>
<th>Tumor type</th>
<th>HNSCC (N=13)</th>
<th>NSCLC (N=10)</th>
<th>MSI-H positive solid tumors (N=5)</th>
<th>Hodgkin Lymphoma (N=9)</th>
<th>Total (N=37)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CBR at 400 mg; n/N (%)</td>
<td>4/7 (57.1)</td>
<td>6/7 (85.7)</td>
<td>0/1 (0)</td>
<td>4/4 (100)</td>
<td>14/19 (73.7)</td>
</tr>
<tr>
<td>CBR at 800 mg; n/N (%)</td>
<td>2/6 (33.3)</td>
<td>1/3 (33.3)</td>
<td>2/4 (50)</td>
<td>3/5 (60)</td>
<td>8/18 (44.4)</td>
</tr>
<tr>
<td>CBR (Total); n/N (%)</td>
<td>6/13 (46.2)</td>
<td>7/10 (70)</td>
<td>2/5 (40)</td>
<td>7/9 (77.8)</td>
<td>22/37 (59.5)</td>
</tr>
</tbody>
</table>

Overall CBR is 59.5% - with a strong trend towards superior benefit at lower dosage. The tendency of improved CBR is seen across all tumor types, except MSI-H where the sample size is smallest (5), and only one (1) patient was randomized to the lower dose (400 mg). The rates of CBR with 400 mg dosage is similar to the rates achieved with PD-1/PD-L1 antibodies. Addition, clear efficacy signals seen among four patients also showed higher activity at 400 mg – out of three (3) patients with Hodgkin lymphoma achieving PR, two (2) received 400 mg: the HNSCC patient having a 48% reduction was also at 400 mg (Figures 1, 2 and 3).

Figure 1: Percentage change from baseline in SPD by irRC criterion in solid tumors and in SUVmax by Lugano criterion in Hodgkin lymphoma.

Safety

Safety is evaluated among all 62 patients who received CA-170. Overall, CA170 has been well tolerated, with Immune Related Adverse Events (irAEs) seen in eight patients – five (5) with increase in TSH or worsening hypothyroidism (2 at 400 mg and 3 at 800 mg), two (2) with skin rash (both at 400 mg) and one (1) with Grade 3 neutropenia and Grade 3 anemia (at 400 mg). The patient (with related Grade 3 neutropenia / Grade 3 anemia) developed lower grade cytophenias within Cycle 1. The counts continued to worsen, necessitating CA-170 interruption in Cycle 3. The hematological values improved after drug interruption, and the counts worsened again when CA-170 was resumed (Figure 5).

Figure 5: Changes in Absolute Neutrophil Count (ANC) and Hemoglobin in a HNSCC Patient (400 mg).

Conclusions

• The Clinical Benefit Rate (CBR) with CA-170 is in a similar range as PD-1/PD-L1 antibodies with Objective Response Rates being lower.
• Better efficacy results are observed at the lower dosage (400 mg), consistent with pre-clinical findings showing bell-shaped curve of immune activation with CA-170 (Figure 4).
• Higher proportion of patients at 400 mg developed Immune Related Adverse Events (irAEs).
• Immune related hematological events, reported either as fatal or of prolonged duration with antibodies[8,9,10], have been reversible and of shorter duration.
• The CA-170 dose and dosing regimen are being optimized in relevant tumor types to achieve higher responses, in view of short half life[1] and PD data showing bell shaped immune activation.

References


Figure 4: Functional antagonism of PD-L1/L2 and VISTA with CA-170

Figure 5: Changes in Absolute Neutrophil Count (ANC) and Hemoglobin in a HNSCC Patient (400 mg).