<table>
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<th><strong>Official Protocol Title:</strong></th>
<th>A Phase Ib Multi-Cohort Study of MK-3475 in Subjects with Advanced Solid Tumors</th>
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<td><strong>NCT number:</strong></td>
<td>NCT01848834</td>
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<td><strong>Document Date:</strong></td>
<td>11-Dec-2017</td>
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Protocol-specific Sponsor Contact information can be found in the Investigator Trial File Binder.

TITLE:

A Phase Ib Multi-Cohort Study of MK-3475 in Subjects with Advanced Solid Tumors

IND NUMBER: 110,080

EudraCT NUMBER: 2012-005771-14
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### SUMMARY OF CHANGES

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<th>Section Title(s)</th>
<th>Description of Change(s)</th>
<th>Rationale</th>
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<tbody>
<tr>
<td>5.2.1.2</td>
<td>Dose Modification (Escalation/Titration/Other)</td>
<td>The dose modification guidelines are expanded to cover supportive care, monitoring, and follow-up. Myocarditis is added.</td>
<td>To provide current, comprehensive guidelines for management of immune-related adverse events.</td>
</tr>
<tr>
<td>6.1</td>
<td>Study Flow Chart for Cohorts A, B, C and D (10 mg/kg Q2W dosing)</td>
<td>Survival status activities are shown taking place throughout the trial. The frequency of telephone contacts during the survival follow-up phase is changed from every 12 weeks to approximately every 12 weeks.</td>
<td>To allow flexibility of survival status activities and ensure that current, complete survival data are available at the time of database locks.</td>
</tr>
<tr>
<td>6.2</td>
<td>Study Flow Chart for Cohort B2 (200 mg Q3W dosing)</td>
<td>Survival status activities are shown taking place throughout the trial. The frequency of telephone contacts during the survival follow-up phase is changed from every 12 weeks to approximately every 12 weeks.</td>
<td>To allow flexibility of survival status activities and ensure that current, complete survival data are available at the time of database locks.</td>
</tr>
<tr>
<td>6.3</td>
<td>Second Course Phase (Retreatment ONLY) for Cohorts A, B, C and D</td>
<td>ORST (Oncologic Response-Solid Tumor) is defined.</td>
<td>To provide definition of this term.</td>
</tr>
<tr>
<td>6.4</td>
<td>Second Course Phase (Retreatment ONLY) for Cohort B2</td>
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<td>Section Title(s)</td>
<td>Description of Change (s)</td>
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<tr>
<td>7.1.5.4.1</td>
<td>Survival Follow-up</td>
<td>The frequency of telephone contacts during the survival follow-up phase is changed from every 12 weeks to approximately every 12 weeks.</td>
<td>To allow flexibility of survival status activities</td>
</tr>
<tr>
<td>7.1.5.5</td>
<td>Survival Status</td>
<td>This section is added to state that the Sponsor may request updated survival data during the course of the trial.</td>
<td>To allow flexibility of survival status activities and ensure that current, complete survival data are available at the time of database locks</td>
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**ADDITIONAL CHANGE(S) FOR THIS AMENDMENT:**

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<th>Rationale</th>
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<tr>
<td>1.0</td>
<td>Trial Summary</td>
<td>Once subjects have achieved the trial objective or the trial has ended, subjects are discontinued from this trial and will be enrolled in an extension trial to continue protocol-defined assessments and treatment.</td>
<td>This trial has been identified to roll over into an extension trial.</td>
</tr>
<tr>
<td>2.2</td>
<td>Trial Diagram</td>
<td>The pembrolizumab extension trial after survival follow-up is added.</td>
<td></td>
</tr>
<tr>
<td>5.10</td>
<td>Beginning and End of the Trial</td>
<td>Upon trial completion, subjects are discontinued and enrolled in a pembrolizumab extension trial.</td>
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<tr>
<td>Section Number(s)</td>
<td>Section Title(s)</td>
<td>Description of Change(s)</td>
<td>Rationale</td>
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</tr>
<tr>
<td>6.1</td>
<td>Study Flow Chart for Cohorts A, B, C and D (10 mg/kg Q2W dosing)</td>
<td>Text was added to allow flexibility around the imaging schedule for subjects in follow-up.</td>
<td>To maintain subject retention and the collection of survival status for every subject enrolled</td>
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<tr>
<td>6.2</td>
<td>Study Flow Chart for Cohort B2 (200 mg Q3W dosing)</td>
<td></td>
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<tr>
<td>6.3</td>
<td>Second Course Phase (Retreatment ONLY) for Cohorts A, B, C and D</td>
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<td></td>
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<tr>
<td>6.4</td>
<td>Second Course Phase (Retreatment ONLY) for Cohort B2</td>
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<tr>
<td>7.1.2.5.1</td>
<td>Assessment of Disease</td>
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<td>7.1.5.4</td>
<td>Follow-up Visits</td>
<td></td>
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<tr>
<td>5.6.1</td>
<td>Supportive Care</td>
<td>Reference to the ECI (events of clinical interest) guidance document is removed.</td>
<td>This guidance document has been retired.</td>
</tr>
<tr>
<td>7.1.2.1</td>
<td>Adverse Event (AE) Monitoring</td>
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1.0 TRIAL SUMMARY

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<th>Abbreviated Title</th>
<th>Phase Ib Multicohort Study of MK-3475 in Subjects with Advanced Solid Tumors</th>
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<tr>
<td>Trial Phase</td>
<td>Ib</td>
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<tr>
<td>Clinical Indication</td>
<td>The treatment of subjects with triple negative breast cancer, head/neck cancer, urothelial tract cancer, or gastric cancer.</td>
</tr>
<tr>
<td>Trial Type</td>
<td>Interventional</td>
</tr>
<tr>
<td>Type of control</td>
<td>No treatment control</td>
</tr>
<tr>
<td>Route of administration</td>
<td>Intravenous</td>
</tr>
<tr>
<td>Trial Blinding</td>
<td>Unblinded Open-label</td>
</tr>
</tbody>
</table>
| Treatment Groups  | MK-3475 10 mg/kg every 2 weeks (Cohort A, B, C and D)  
MK-3475 200 mg every 3 weeks (Cohort B2) |
| Number of trial subjects | Approximately 114 subjects originally planned for enrollment into Cohorts A, B, C, and D, plus 110 additional H/N expansion (Cohort B2) subjects will be enrolled. |
| Estimated duration of trial | The sponsor estimates that the trial will require approximately 42 months from the time the first subject signs the informed consent until the last subject’s last visit. |
| Duration of Participation | Each subject will participate in the trial from the time the subject signs the Informed Consent Form (ICF) through the final protocol-specified contact. After a screening phase of 28 days, eligible subjects will receive treatment on Day 1 of each 2-week dosing cycle (Cohorts A, B, C, and D) or 3-week dosing cycle (Cohort B2). Treatment with MK-3475 will continue until documented confirmed disease progression, unacceptable adverse event(s), intercurrent illness that prevents further administration of treatment, investigator’s decision to withdraw the subject, subject withdraws consent, pregnancy of the subject, noncompliance with trial treatment or procedure requirements, subject receives 24 months of study medication, or administrative reasons. Subjects who attain a complete response may consider stopping trial treatment if they meet criteria for holding therapy. Subjects who stop trial treatment after receiving 24 months of study medication for reasons other than disease progression or intolerability or who attain a complete response and stop trial treatment may be eligible for up to one year of retreatment after experiencing disease progression. The decision to retreat will be at the discretion of the investigator only if they meet the criteria for retreatment and the trial is ongoing. After the end of treatment, each subject will be followed for 30 days for adverse event monitoring (serious adverse events will be collected for 90 days after the end of treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier). Subjects who discontinue for reasons other than disease progression will have post-treatment follow-up for disease status until disease progression, initiating a non-study cancer treatment, withdrawing consent, or becoming lost to follow-up. All subjects will be followed by telephone for overall survival until death, withdrawal of consent, or the end of the study. Once subjects have achieved the trial objective or the trial has ended, subjects are discontinued from this trial and will be enrolled in an extension trial to continue protocol-defined assessments and treatment. |
2.0 TRIAL DESIGN

2.1 Trial Design

This is a multicenter, nonrandomized, multi-cohort trial of MK-3475 in subjects with advanced solid tumors. Subjects will be enrolled into Cohort A for triple negative breast cancer (TNBC), Cohort B for the initial head and neck (H/N) cancer cohort and Cohort B2 for the H/N cancer expansion cohort, Cohort C for urothelial tract cancer, or Cohort D for gastric cancer. Only subjects with PD-L1 positive tumors will be enrolled in Cohorts A, B, C and D. Subjects in Cohort B2 may be enrolled regardless of PD-L1 status. Approximately 114 subjects were planned to be enrolled in Cohorts A, B, C and D of this trial to examine the safety and efficacy of MK-3475 in these populations. Approximately 110 subjects will be enrolled into Cohort B2 of the study to further explore the safety and efficacy in the head and neck cancer population at a different dose and schedule of MK-3475 and including both PD-L1 positive and negative subjects. Subjects enrolled in Cohorts A, B, C and D will receive 10 mg/kg of MK-3475 administered every 2 weeks. Subjects enrolled in Cohort B2 will receive 200 mg of MK-3475 administered every 3 weeks. Subjects will be evaluated every 8 weeks (56 days ± 7 days) with radiographic imaging to assess response to treatment. RECIST 1.1 response rate as assessed by independent central radiology review will be used as the primary efficacy endpoint for Cohorts A, B, C and D. RECIST 1.1 response rate as assessed by independent central radiology review will be used as the primary efficacy endpoint for Cohort B2. RECIST 1.1 will be adapted as described in Section 4.2.3.1 to accommodate for the tumor response patterns seen with MK-3475 treatment (e.g., tumor flare), and this adapted RECIST will be used by the sites for treatment decisions in all cohorts. Adverse events will be monitored throughout the trial and graded in severity according to the guidelines outlined in the NCI Common Terminology Criteria for Adverse Events (CTCAE) version 4.0. Treatment with MK-3475 will continue until documented disease progression, unacceptable adverse event(s), intercurrent illness that prevents further administration of treatment, investigator’s decision to withdraw the subject, subject withdraws consent, pregnancy of the subject, noncompliance with trial treatment or procedure requirements, completion of 24 months of treatment with MK-3475, or administrative reasons. Subjects who attain an investigator-determined confirmed complete response (CR) may consider stopping trial treatment after receiving at least 24 weeks of treatment. Subjects who discontinue after at least 24 months of therapy for reasons other than disease progression or intolerability or who discontinue after attaining a CR may be eligible for up to one year of retreatment after they have experienced radiographic disease progression. The decision to retreat will be at the discretion of the investigator only if no cancer treatment was administered since the last dose of MK-3475, the subject still meets the safety parameters listed in the Inclusion/Exclusion criteria and the trial remains open (refer to Section 7.1.5.2.1 for further details). After the end of treatment, each subject will be followed for 30 days for adverse event monitoring (serious adverse events will be collected for 90 days after the end of treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier). Subjects who discontinue treatment for reasons other than disease progression will have post-treatment follow-up for disease status until disease progression, initiating a non-study cancer treatment, withdrawing consent,
or becoming lost to follow-up. All subjects will be followed by telephone contact for overall survival until death, withdrawal of consent or the end of the study, whichever comes first.

The primary objectives of the trial are to determine the safety, tolerability, and anti-tumor activity of MK-3475 in subjects with advanced solid tumors receiving either the 10 mg/kg Q2W dose (Cohorts A, B, C and D) or the 200 mg Q3W dose (Cohort B2). Secondary objectives include progression-free survival (PFS), overall survival (OS) and response duration in subjects with advanced solid tumors. In addition, the anti-tumor activity of MK-3475 in subjects with PD-L1 positive advanced human papillomavirus (HPV) positive head/neck cancer will be evaluated as a secondary objective. The relationship between candidate efficacy/resistance biomarkers (including PD-L1 expression in the tumor and its microenvironment) and anti-tumor activity of MK-3475 will also be investigated as a secondary objective. The pharmacokinetic (PK) properties of MK-3475 in the advanced solid tumor population will be investigated as an exploratory objective.

Participation in this trial will be dependent upon supplying tissue from an archival tissue sample or newly obtained biopsy of a tumor lesion not previously irradiated (subjects in the H/N cohort may submit tissue from a previously-irradiated lesion). This specimen will be evaluated at a central laboratory for expression status of PD-L1 by immunohistochemistry (IHC). Only subjects with PD-L1 positive tumors will be enrolled into Cohorts A, B, C and D of the trial. Both PD-L1 positive and negative subjects will be enrolled into Cohort B2, and the clinical activity in both subsets will be evaluated.

This study will be conducted in conformance with Good Clinical Practices.

Specific procedures to be performed during the trial, as well as their prescribed times and associated visit windows, are outlined in the Trial Flow Chart - Section 6.0. Details of each procedure are provided in Section 7.0 – Trial Procedures.
2.2 Trial Diagram

The trial design is depicted in Figure 1 below.

![Trial Design Diagram]

*An interim analysis for each cohort may be performed depending on the rate of enrollment or other factors determined during the course of the trial. This interim analysis would only be performed when ≥ 10 patients in the respective cohort have had at least two post-baseline scans.
*A total of 34 subjects with head/neck cancer will be enrolled in Cohort B of the study.
#The gastric cancer cohort will be stratified to enroll 16 patients in Asia Pacific (AP) and 16 patients ex AP. No interim analysis will be performed in this cohort. 
PD = Progressive Disease
SFU = Survival Follow-up

Figure 1 Trial Design

3.0 OBJECTIVE(S) & HYPOTHESIS(ES)

3.1 Primary Objective(s) & Hypothesis(es)

(1) **Objective:** To determine the safety and tolerability of the 10 mg/kg Q2W dose of MK-3475 in subjects with PD-L1 positive advanced solid tumors enrolled into Cohorts A, B, C, and D.

**Hypothesis:** Intravenous administration of the single agent 10 mg/kg Q2W dose of MK-3475 is sufficiently well-tolerated to permit continued clinical investigation.
(2) **Objective:** To evaluate anti-tumor activity of the 10 mg/kg Q2W dose of MK-3475 in subjects with PD-L1 positive advanced solid tumors enrolled into Cohorts A, B, C, and D based on RECIST 1.1 assessed by independent central radiology review.

a. **Hypothesis:** Intravenous administration of single agent MK-3475 10 mg/kg Q2W to subjects with PD-L1 positive triple negative advanced breast cancer (Cohort A) will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1.

b. **Hypothesis:** Intravenous administration of single agent MK-3475 10 mg/kg Q2W to subjects with PD-L1 positive HPV negative advanced head and neck cancer (Cohort B) will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1.

c. **Hypothesis:** Intravenous administration of single agent MK-3475 10 mg/kg Q2W to subjects with PD-L1 positive advanced urothelial tract cancer (Cohort C) will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1.

d. **Hypothesis:** Intravenous administration of single agent MK-3475 10 mg/kg Q2W to subjects with PD-L1 positive advanced gastric cancer (Cohort D) will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1.

(3) **Objective:** To determine the safety and tolerability of the 200 mg Q3W dose of MK-3475 in subjects with advanced head and neck cancer enrolled into Cohort B2.

**Hypothesis:** Intravenous administration of single agent MK-3475 200 mg Q3W in subjects with advanced head and neck cancer is sufficiently well-tolerated to permit continued clinical investigation.

(4) **Objective:** To evaluate anti-tumor activity of MK-3475 200 mg Q3W (Cohort B2) in subjects with advanced head and neck cancer enrolled into Cohort B2 based on RECIST 1.1 assessed by independent central radiology review.

**Hypothesis:** Intravenous administration of the single agent 200 mg Q3W dose of MK-3475 to subjects with advanced head and neck cancer (Cohort B2) will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1 assessed by independent central radiology review.

3.2 **Secondary Objective(s) & Hypothesis(es)**

(1) **Objective:** To evaluate the anti-tumor activity of MK-3475 in subjects with PD-L1 positive advanced HPV positive head/neck cancer in Cohort B as determined by independent central radiology review.

**Hypothesis:** Intravenous administration of single agent MK-3475 to subjects with PD-L1 positive HPV positive advanced head and neck cancer enrolled into Cohort B will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1 as assessed by independent central radiology review.
 Objective: To evaluate the anti-tumor activity of MK-3475 in subjects with PD-L1 positive advanced gastric cancer in the Asia Pacific region as assessed by independent central radiology review.

Hypothesis: Intravenous administration of single agent MK-3475 to subjects with PD-L1 positive advanced gastric cancer in the Asia Pacific region will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1 as assessed by independent central radiology review.

Objective: To evaluate the anti-tumor activity of MK-3475 in subjects with advanced head and neck cancer previously treated with cetuximab and platinum enrolled in Cohort B or Cohort B2 based on RECIST 1.1 as assessed by independent central radiology review.

Hypothesis: Intravenous administration of single agent MK-3475 in subjects with advanced head and neck cancer previously treated with cetuximab and platinum enrolled in Cohort B or Cohort B2 will result in a clinically meaningful overall response rate (ORR) based on RECIST 1.1 as assessed by independent central radiology review.

Objective: To evaluate the anti-tumor activity of MK-3475 in subjects with PD-L1 positive advanced solid tumors based on RECIST 1.1 as determined by the Investigator (Cohorts A, B, C and D).

Objective: To evaluate the anti-tumor activity of MK-3475 at 200 mg Q3W in subjects with advanced head and neck cancer (Cohort B2) based on RECIST 1.1 as assessed by the Investigator.

Objective: To investigate the correlation between PD-L1 expression and anti-tumor activity of pembrolizumab in subjects with advanced head and neck cancer enrolled into Cohort B2.

Objective: To investigate the relationship between candidate efficacy/resistance biomarkers and anti-tumor activity of pembrolizumab utilizing pre- and post-treatment tumor biopsies and blood sampling.

Objective: To evaluate the progression-free survival (PFS) in subjects with advanced solid tumors receiving MK-3475.

Objective: To evaluate the overall survival (OS) in subjects with advanced solid tumors receiving MK-3475.

Objective: To evaluate the response duration in subjects with advanced solid tumors receiving MK-3475.
3.3 Exploratory Objective

(1) **Objective:** To explore the PK profile of MK-3475 in the advanced solid tumor population.

4.0 BACKGROUND & RATIONALE

4.1 Background

4.1.1 Pharmaceutical and Therapeutic Background

The importance of intact immune surveillance in controlling outgrowth of neoplastic transformation has been known for decades [1]. Accumulating evidence shows a correlation between tumor-infiltrating lymphocytes (TILs) in cancer tissue and favorable prognosis in various malignancies [2; 3; 4; 5; 6]. In particular, the presence of CD8+ T-cells and the ratio of CD8+ effector T-cells / FoxP3+ regulatory T-cells seems to correlate with improved prognosis and long-term survival in many solid tumors.

The PD-1 receptor-ligand interaction is a major pathway hijacked by tumors to suppress immune control. The normal function of PD-1, expressed on the cell surface of activated T-cells under healthy conditions, is to down-modulate unwanted or excessive immune responses, including autoimmunity reactions. PD-1 (encoded by the gene Pdcd1) is an Ig superfamily member related to CD28 and CTLA-4 which has been shown to negatively regulate antigen receptor signaling upon engagement of its ligands (PD-L1 and/or PD-L2) [7; 8]. The structure of murine PD-1 has been resolved [9]. PD-1 and family members are type I transmembrane glycoproteins containing an Ig Variable-type (V-type) domain responsible for ligand binding and a cytoplasmic tail which is responsible for the binding of signaling molecules. The cytoplasmic tail of PD-1 contains 2 tyrosine-based signaling motifs, an immunoreceptor tyrosine-based inhibition motif (ITIM) and an immunoreceptor tyrosine-based switch motif (ITSM). Following T-cell stimulation, PD-1 recruits the tyrosine phosphatases SHP-1 and SHP-2 to the ITSM motif within its cytoplasmic tail, leading to the dephosphorylation of effector molecules such as CD3ζ, PKCθ and ZAP70 which are involved in the CD3 T-cell signaling cascade [7; 10; 11; 12]. The mechanism by which PD-1 down modulates T-cell responses is similar to, but distinct from that of CTLA-4 as both molecules regulate an overlapping set of signaling proteins [13; 14]. PD-1 was shown to be expressed on activated lymphocytes including peripheral CD4+ and CD8+ T-cells, B-cells, Tregs and Natural Killer cells [15; 16]. Expression has also been shown during thymic development on CD4-Cd8- (double negative) T-cells as well as subsets of macrophages and dendritic cells [17]. The ligands for PD-1 (PD-L1 and PD-L2) are constitutively expressed or can be induced in a variety of cell types, including non-hematopoietic tissues as well as in various tumors [18; 19; 20; 13]. Both ligands are type I transmembrane receptors containing both IgV- and IgC-like domains in the extracellular region and contain short cytoplasmic regions with no known signaling motifs. Binding of either PD-1 ligand to PD-1 inhibits T-cell activation triggered through the T-cell receptor. PD-L1 is expressed at low levels on various non-hematopoietic tissues, most notably on vascular endothelium, whereas PD-L2 protein is only detectably expressed on antigen-presenting cells found in lymphoid tissue or...
chronic inflammatory environments. PD-L2 is thought to control immune T-cell activation in lymphoid organs, whereas PD-L1 serves to dampen unwarranted T-cell function in peripheral tissues [13]. Although healthy organs express little (if any) PD-L1, a variety of cancers were demonstrated to express abundant levels of this T-cell inhibitor. PD-1 has been suggested to regulate tumor-specific T-cell expansion in subjects with melanoma (MEL) [21]. This suggests that the PD-1/PD-L1 pathway plays a critical role in tumor immune evasion and should be considered as an attractive target for therapeutic intervention.

MK-3475 (previously known as SCH 900475) is a potent and highly selective humanized monoclonal antibody (mAb) of the IgG4/kappa isotype designed to directly block the interaction between PD-1 and its ligands, PD-L1 and PD-L2.

4.1.2 Pre-clinical and Clinical Trials

Therapeutic studies in mouse models have shown that administration of antibodies blocking PD-1/PD-L1 interaction enhances infiltration of tumor-specific CD8+ T-cells and leads ultimately to tumor rejection, either as a mono-therapy or in combination with other treatment modalities. Anti-mouse PD-1 or anti-mouse PD-L1 antibodies have demonstrated anti-tumor responses as a mono-therapy in models of squamous cell carcinoma, pancreatic carcinoma, MEL and colorectal carcinoma. Blockade of the PD-1 pathway effectively promoted CD8+ T-cell infiltration into the tumor and the presence of IFN-γ, granzyme B and perforin, indicating that the mechanism of action involved local infiltration and activation of effector T-cell function in vivo [22; 23; 24; 25; 26; 27]. Experiments have confirmed the in vivo efficacy of PD-1 blockade as a mono-therapy as well as in combination with chemotherapy in syngeneic mouse tumor models (see the Investigator’s Brochure [IB]).

4.1.3 Ongoing Clinical Trials

Ongoing clinical trials are being conducted in advanced melanoma, non-small cell lung cancer, a number of advanced solid tumor indications and hematologic malignancies. For study details please refer to the Investigator’s Brochure.

4.2 Rationale

4.2.1 Rationale for the Trial and Selected Subject Population

Details regarding specific benefits and risks for subjects participating in this clinical trial may be found in the accompanying IB and Informed Consent documents.

This is a multicenter, nonrandomized, multi-cohort trial of MK-3475 in subjects with advanced solid tumors. Subjects with PD-L1 positive tumors will be enrolled into Cohort A for TNBC, Cohort B for H/N cancer, Cohort C for urothelial tract cancer or Cohort D for gastric cancer. Subjects with advanced head and neck cancer will be enrolled into Cohort B2 irrespective of PD-L1 expression (including subjects whose tumors do not express PD-L1).
Participation in this trial will be dependent upon supplying tissue from an archival tissue sample or newly obtained core or excisional biopsy of a tumor lesion not previously irradiated (subjects in the H/N cohort may provide tissue from a previously irradiated lesion) to evaluate for PD-L1 expression by IHC. For Cohorts A, B, C or D, if an archival specimen is PD-L1 negative but a newly obtained biopsy is positive, the subject would be eligible. The specimen will be evaluated at a central laboratory for expression status of PD-L1. Only subjects with PD-L1 positive tumors will be enrolled into Cohorts A, B, C and D of the trial. The hypothesis that PD-L1 may predict potential response to anti-PD-1 therapy is based on the results from Topalian et al [30] who examined PD-L1 expression in the archival specimens of 42 of the 296 subjects treated with the PD-1 inhibitor nivolumab. Of those 17 subjects whose tumor cells did not stain positive for PD-L1 using a 5% threshold of tumor cell surface expression, no objective response by RECIST 1.1 was observed. But among the 25 subjects whose tumor cells were considered positive for PD-L1, 9 responded (36%).

Further, data from MK-3475 in patients with non-small cell lung cancer (NSCLC) showed that pretreatment tumor PD-L1 expression was a statistically significant predictor of response. In patients with evaluable tumor PD-L1 expression, confirmed responses by RECIST v1.1 (and irRC) occurred in patients with tumors strongly positive for PD-L1 [42]. Therefore, PD-L1 may be a predictive biomarker of anti-PD-1 activity, and will be tested in this study as an enrollment criterion for select cohorts A, B, C, and D.

A subset of head and neck cancer is caused by infection with HPV. It is hypothesized that virally induced tumors may have enhanced immunogenicity and PD-L1 expression [31]. Therefore, additional subjects that are HPV positive will be enrolled into the initial H/N cohort (Cohort B). Cetuximab and platinum are commonly used therapies for head and neck patients in the recurrent and metastatic setting. Understanding the response rate of pembrolizumab in patients who have received these standard treatments is of interest. A subset of patients with prior cetuximab and platinum treatment will be analyzed to further characterize clinical response in head and neck cancer patients exposed to cetuximab and platinum therapy.

### 4.2.2 Rationale for Dose Selection/Regimen/Modification

An open-label Phase I trial (Protocol 001) is being conducted to evaluate the safety and clinical activity of single agent MK-3475. The dose escalation portion of this trial evaluated three dose levels, 1 mg/kg, 3 mg/kg, and 10 mg/kg, administered every 2 weeks (Q2W) in subjects with advanced solid tumors. All three dose levels were well tolerated and no dose-limiting toxicities were observed. This first in human study of MK-3475 showed evidence of target engagement and objective evidence of tumor size reduction at all dose levels (1 mg/kg, 3 mg/kg and 10 mg/kg Q2W). No MTD has been identified to date. 10.0 mg/kg Q2W, the highest dose tested in PN001, will be the dose and schedule utilized in Cohorts A, B, C and D of this protocol to test for initial tumor activity. Recent data from other clinical studies within the MK-3475 program has shown that a lower dose of MK-3475 and a less frequent schedule may be sufficient for target engagement and clinical activity [43]. To further investigate this hypothesis subjects in Cohort B2 will be dosed at 200 mg Q3W.
PK data analysis of MK-3475 administered Q2W and Q3W showed slow systemic clearance, limited volume of distribution, and a long half-life (refer to IB). Pharmacodynamic data (IL-2 release assay) suggested that peripheral target engagement is durable (>21 days). This early PK and pharmacodynamic data provides scientific rationale for testing a Q2W and Q3W dosing schedule.

A population pharmacokinetic analysis has been performed using serum concentration time data from 476 patients. Within the resulting population PK model, clearance and volume parameters of MK-3475 were found to be dependent on body weight. The relationship between clearance and body weight, with an allometric exponent of 0.59, is within the range observed for other antibodies and would support both body weight normalized dosing or a fixed dose across all body weights. MK-3475 has been found to have a wide therapeutic range based on the melanoma indication. The differences in exposure for a 200 mg fixed dose regimen relative to a 2 mg/kg Q3W body weight based regimen are anticipated to remain well within the established exposure margins of 0.5 – 5.0 for MK-3475 in the melanoma indication. The exposure margins are based on the notion of similar efficacy and safety in melanoma at 10 mg/kg Q3W vs. the proposed dose regimen of 2 mg/kg Q3W (i.e. 5-fold higher dose and exposure). The population PK evaluation revealed that there was no significant impact of tumor burden on exposure. In addition, exposure was similar between the NSCLC and melanoma indications. Therefore, there are no anticipated changes in exposure between different indication settings.

4.2.3 Rationale for Endpoints

4.2.3.1 Efficacy Endpoints

The primary efficacy objective of this study is to evaluate the anti-tumor activity of MK-3475 in subjects with advanced solid tumors. Response rates per RECIST 1.1 and volumetric analysis will be evaluated.

RECIST 1.1[32] as assessed by the independent central radiology vendor will be used as the primary response rate efficacy endpoint for Cohorts A, B, C and D. RECIST 1.1 [32] as assessed by independent central radiology review will be used as the primary response rate efficacy endpoint for Cohort B2.

RECIST 1.1 will also be used by the local site to determine eligibility and make treatment decisions. For this purpose RECIST 1.1 will be adapted to account for the unique tumor response profile seen with treatment of MK-3475.

Immunotherapeutic agents such as MK-3475 may produce antitumor effects by potentiating endogenous cancer-specific immune responses which may be functionally anergic. The response patterns seen with such an approach may extend beyond the typical time course of responses seen with cytotoxic agents, and can manifest a clinical response after an initial increase in tumor burden or even the appearance of new lesions. Standard RECIST criteria may not provide a complete response assessment of immunotherapeutic agents such as MK-3475. Therefore, RECIST 1.1 will be used with the following adaptation:
If radiologic imaging shows PD, tumor assessment should be repeated ≥ 4 weeks later in order to confirm PD with the option of continuing treatment per below while awaiting radiologic confirmation of progression. If repeat imaging shows a reduction in the tumor burden compared to the initial scan demonstrating PD, treatment may be continued as per treatment calendar. If repeat imaging confirms progressive disease, subjects will be discontinued from study therapy (exception noted in Section 7.1.2.5.1). In determining whether or not the tumor burden has increased or decreased, Investigators should consider all target lesions as well as non-target lesions (please refer to the Site Imaging Manual).

In subjects who have initial evidence of radiological PD, it is at the discretion of the treating physician whether to continue a subject on study treatment until repeat imaging is obtained a minimum of 4 weeks later. This decision should be based on the clinical judgment of the subject’s overall clinical condition, including performance status, clinical symptoms, and laboratory data. Subjects may receive treatment while waiting for confirmation of PD if they are clinically stable as defined by the following criteria:

- Absence of signs and symptoms indicating disease progression
- No decline in ECOG performance status
- Absence of rapid progression of disease
- Absence of progressive tumor at critical anatomical sites (e.g., cord compression) requiring urgent alternative medical intervention

When feasible, subjects should not be discontinued until progression is confirmed. This allowance to continue treatment despite initial radiologic progression takes into account the observation that some subjects can have a transient tumor flare in the first few months after the start of immunotherapy, but with subsequent disease response. Subjects that are deemed clinically unstable are not required to have repeat imaging for confirmation of progressive disease. Retrospective independent central radiology review of all imaging time points will be performed for this study using RECIST 1.1 and volumetric analysis. Additional information is included in the Site Imaging Manual.

### 4.2.3.2 Safety Endpoints

The primary safety objective of this study is to characterize the safety and tolerability of MK-3475 in subjects with advanced solid tumors. The primary safety analysis will be based on subjects who experienced toxicities as defined by CTCAE criteria. Safety will be assessed by quantifying the toxicities and grades experienced by subjects who have received MK-3475, including serious adverse events (SAEs) and events of clinical interest (ECIs).

Safety will be assessed by reported adverse experiences using CTCAE, Version 4.0. The attribution to drug, time-of-onset, duration of the event, its resolution, and any concomitant medications administered will be recorded. AEs will be analyzed including but not limited to all AEs, SAEs, fatal AEs, and laboratory changes. Furthermore, specific immune-related adverse events (irAEs) will be collected and designated as immune-related events of clinical interest (ECIs) as described in Section 7.2.3.2.
4.2.3.3 Biomarker Research

Additional biomarker research to identify factors important for MK-3475 therapy may also be pursued. For example, pre- and post-dose tumor and blood samples from this study may undergo proteomic, genomic and transcriptional analyses. Additional research may evaluate factors important for predicting responsiveness or resistance to MK-3475 therapy and other immunologic targets.

Assays may include but are not be limited to:

**Multiparametric (Two-Color) IHC**

Spatial association of PD-1+ tumor infiltrating lymphocytes (TILs) and PD-L1+ cells (tumor and myeloid cells) suggests “induction” of PD-L1. Interferon-gamma production by antigen-specific PD-1+ CD8+ T cells is hypothesized to drive local intratumoral upregulation of PD-L1 on adjacent tumor and myeloid cells, leading to a “stalled CTL” response which may be predictive of response to MK-3475 therapy. By assessing both of the required elements, i.e. PD-L1 positive cells and PD-1+ T cells, a two-color IHC assay may be a better predictor of response than PD-L1 positivity alone.

**Transcriptional Analyses**

Messenger RNA (mRNA) expression profiling in archival material will be completed to assess expression of approximately 400 genes and attempt to define a gene set critical for clinical response to MK-3475. The hypothesis to be tested is that MK-3475 responders will exhibit a “stalled Cytotoxic T Lymphocyte (CTL)” response within the tumor reflected in the physical proximity between PD-1 and PD-L1 expression and the presence of an aborted (e.g., weak but discernible) interferon-gamma transcriptional program will be detectable by profiling analyses. Global profiling will also be pursued.

Expression of individual genes related to the immune system may also be evaluated such as immune signatures and critical cytokines (e.g., IL-10).

**Gene Sequencing**

New data are emerging that suggest we can define certain tumor types as being ‘hypermutated’. There is a potential that this hypermutated state may correlate with response to MK-3475 therapy, and/or that the converse, ‘hypomutated’ state may correlate with non-response.

4.2.3.4 Future Biomedical Research

Merck will conduct Future Biomedical Research on blood and tumor tissue specimens collected during this clinical trial. This research may include genetic analyses (DNA), gene expression profiling (RNA), proteomics, metabolomics (serum, plasma) and/or the measurement of other analytes.
Such research is for biomarker testing to address emergent questions not described elsewhere in the protocol (as part of the main trial) and will only be conducted on specimens from appropriately consented subjects. The objective of collecting specimens for Future Biomedical Research is to explore and identify biomarkers that inform the scientific understanding of diseases and/or their therapeutic treatments. For instance, exploratory pharmacogenetic (PGt) studies may be performed if significant Pharmacokinetic/Pharmacodynamic (PK/PD) relationships are observed or adverse events are identified. Genomic markers of disease may also be investigated. Such retrospective pharmacogenetic studies will be conducted with appropriate biostatistical design and analysis and compared to PK/PD results or clinical outcomes. Any significant PGt relationships to outcome would require validation in future clinical trials. The overarching goal is to use such information to develop safer, more effective drugs, and/or to ensure that subjects receive the correct dose of the correct drug at the correct time. The details of this Future Biomedical Research sub-trial are presented in Section 12.2 - Collection and Management of Specimens for Future Biomedical Research. Additional informational material for institutional review boards/ethics committees (IRBs/ERCs) and investigational site staff is provided in Section 12.3.

5.0 METHODOLOGY

5.1 Entry Criteria

5.1.1 Diagnosis/Condition for Entry into the Trial

Male/Female subjects with triple negative breast cancer (Cohort A), head/neck cancer (Cohort B and Cohort B2), urothelial tract cancer (Cohort C), or gastric cancer (Cohort D) of at least 18 years of age will be enrolled in this trial.

5.1.2 Subject Inclusion Criteria

In order to be eligible for participation in this trial, the subject must:

1. Be willing and able to provide written informed consent/assent for the trial. The subject may also provide consent/assent for Future Biomedical Research. However, the subject may participate in the main trial without participating in Future Biomedical Research.

2. Be ≥ 18 years of age on day of signing informed consent.
3. Have histologically or cytologically-confirmed diagnosis of cancer that is recurrent, metastatic, or persistent and meet the following corresponding requirements for the cohort of the study they will enroll into. There is no limit to the number of prior treatment regimens.

   a. **Cohort A**: Have diagnosis of triple negative breast cancer (Estrogen, Progesterone, and HER2 negative carcinoma of the breast)

   b. **Cohort B and B2**: Have a diagnosis of squamous cell carcinoma of the head and neck

      Note: A subset of subjects enrolled to Cohort B known to have HPV positive head and neck squamous cell cancer will be assessed.

      Note: Cohort B2 will enroll both HPV positive and HPV negative subjects

   c. **Cohort C**: Have a diagnosis of urothelial tract cancer of the renal pelvis, ureter, bladder, or urethra. Both transitional cell and non-transitional cell histologies are allowed.

   d. **Cohort D**: Have a diagnosis of adenocarcinoma of the stomach or gastro-esophageal junction

4. Have provided tissue for biomarker analysis from an archival tissue sample or newly obtained core or excisional biopsy of a tumor lesion not previously irradiated. (Subjects with H/N cancer may provide tissue from a previously irradiated lesion.)

5. Cohorts A, B, C and D only: Have a PD-L1 positive tumor as determined by IHC at a central laboratory from either an archived formalin fixed paraffin embedded (FFPE) tumor sample or a newly obtained biopsy.

6. Have measurable disease based on RECIST 1.1. Tumor lesions situated in a previously irradiated area are considered measurable if progression has been demonstrated in such lesions.

7. Have a performance status of 0 or 1 on the ECOG Performance Scale.

8. Demonstrate adequate organ function as defined in Table 1; all screening labs should be performed within 10 days of treatment initiation.
Table 1 Adequate Organ Function Laboratory Values

<table>
<thead>
<tr>
<th>System</th>
<th>Laboratory Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hematological</strong></td>
<td></td>
</tr>
<tr>
<td>Absolute neutrophil count (ANC)</td>
<td>≥1,500 /mcL</td>
</tr>
<tr>
<td>Platelets</td>
<td>≥100,000 / mcL</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>≥9 g/dL or ≥5.6 mmol/L</td>
</tr>
<tr>
<td><strong>Renal</strong></td>
<td></td>
</tr>
<tr>
<td>Creatinine <strong>OR</strong></td>
<td>≤1.5 X upper limit of normal (ULN) <strong>OR</strong></td>
</tr>
<tr>
<td>Measured or calculated(^a) creatinine clearance (GFR can also be used in place of creatinine or CrCl)</td>
<td>≥60 mL/min for subject with creatinine levels &gt; 1.5 X institutional ULN</td>
</tr>
<tr>
<td><strong>Hepatic</strong></td>
<td></td>
</tr>
<tr>
<td>Total bilirubin</td>
<td>≤1.5 X ULN <strong>OR</strong></td>
</tr>
<tr>
<td>Direct bilirubin ≤ ULN for subjects with total bilirubin levels &gt; 1.5 ULN</td>
<td></td>
</tr>
<tr>
<td>AST (SGOT) and ALT (SGPT)</td>
<td>≤2.5 X ULN <strong>OR</strong></td>
</tr>
<tr>
<td>≤ 5 X ULN for subjects with liver metastases</td>
<td></td>
</tr>
<tr>
<td><strong>Coagulation</strong></td>
<td></td>
</tr>
<tr>
<td>International Normalized Ratio (INR) or Prothrombin Time (PT)</td>
<td>≤1.5 X ULN unless subject is receiving anticoagulant therapy as long as PT or PTT is within therapeutic range of intended use of anticoagulants</td>
</tr>
<tr>
<td>Activated Partial Thromboplastin Time (aPTT)</td>
<td>≤1.5 X ULN unless subject is receiving anticoagulant therapy as long as PT or PTT is within therapeutic range of intended use of anticoagulants</td>
</tr>
</tbody>
</table>

\(^a\) Creatinine clearance should be calculated per institutional standard.

9. Female subject of childbearing potential should have a negative urine or serum pregnancy test within 72 hours prior to receiving the first dose of study medication. If the urine test is positive or cannot be confirmed as negative, a serum pregnancy test will be required.

10. Female subjects of childbearing potential should be willing to use 2 methods of birth control or be surgically sterile, or abstain from heterosexual activity for the course of the study through 120 days after the last dose of study medication (Reference Section 5.7.2). Subjects of childbearing potential are those who have not been surgically sterilized or have not been free from menses for > 1 year.

11. Male subjects should agree to use an adequate method of contraception starting with the first dose of study therapy through 120 days after the last dose of study therapy.

5.1.3 Subject Exclusion Criteria

The subject must be excluded from participating in the trial if the subject:

1. Is currently participating in or has participated in a study of an investigational agent or using an investigational device within 4 weeks of the first dose of treatment.
2. Has a diagnosis of immunosuppression or is receiving systemic steroid therapy or any other form of immunosuppressive therapy within 7 days prior to the first dose of trial treatment.

3. Has had a prior anti-cancer monoclonal antibody within 4 weeks prior to study Day 1 or who has not recovered (i.e., ≤ Grade 1 or at baseline) from adverse events due to agents administered more than 4 weeks earlier.

4. Has had prior chemotherapy, targeted small molecule therapy, or radiation therapy within 2 weeks prior to study Day 1 or who has not recovered (i.e., ≤ Grade 1 or at baseline) from adverse events due to a previously administered agent.
   - Note: Subjects with ≤ Grade 2 neuropathy are an exception to this criterion and may qualify for the study.
   - Note: If subject received major surgery, they must have recovered adequately from the toxicity and/or complications from the intervention prior to starting therapy.

5. Has a known additional malignancy that is progressing or requires active treatment. Exceptions include basal cell carcinoma of the skin, squamous cell carcinoma of the skin, or in situ cervical cancer that has undergone potentially curative therapy.

6. Has known active central nervous system (CNS) metastases and/or carcinomatous meningitis. Subjects with previously treated brain metastases may participate provided they are stable (without evidence of progression by imaging for at least four weeks prior to the first dose of trial treatment and any neurologic symptoms have returned to baseline), have no evidence of new or enlarging brain metastases, and are not using steroids for at least 7 days prior to trial treatment.

7. Has an active autoimmune disease requiring systemic treatment within the past 3 months or a documented history of clinically severe autoimmune disease, or a syndrome that requires systemic steroids or immunosuppressive agents. Subjects with vitiligo, diabetes mellitus type I, or resolved childhood asthma/atopy would be an exception to this rule. Subjects that require intermittent use of bronchodilators or local steroid injections would not be excluded from the study. Subjects with hypothyroidism stable on hormone replacement or Sjogren’s syndrome will not be excluded from the study.


9. Has an active infection requiring systemic therapy.

10. Has a history or current evidence of any condition, therapy, or laboratory abnormality that might confound the results of the trial, interfere with the subject’s participation for the full duration of the trial, or is not in the best interest of the subject to participate, in the opinion of the treating investigator.
11. Has known psychiatric or substance abuse disorders that would interfere with cooperation with the requirements of the trial.

12. Is pregnant or breastfeeding, or expecting to conceive or father children within the projected duration of the trial, starting with the pre-screening or screening visit through 120 days after the last dose of trial treatment.

13. Has received prior therapy with an anti-PD-1, anti-PD-L1, anti-PD-L2, anti-CD137, or anti-Cytotoxic T-lymphocyte-associated antigen-4 (CTLA-4) antibody (including ipilimumab or any other antibody or drug specifically targeting T-cell co-stimulation or checkpoint pathways).


15. Has known active Hepatitis B (e.g., HBsAg reactive) or Hepatitis C (e.g., HCV RNA [qualitative] is detected).

16. Has received a live vaccine within 30 days of planned start of study therapy.

   Note: Seasonal influenza vaccines for injection are generally inactivated flu vaccines and are allowed; however intranasal influenza vaccines (e.g., Flu-Mist®) are live attenuated vaccines, and are not allowed.

17. Is or has an immediate family member (e.g., spouse, parent/legal guardian, sibling or child) who is investigational site or sponsor staff directly involved with this trial, unless prospective IRB approval (by chair or designee) is given allowing exception to this criterion for a specific subject.

### 5.2 Trial Treatments

The treatment to be used in this trial is outlined below in **Table 2**.

**Table 2  Trial Treatment**

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Drug</th>
<th>Dose/Potency</th>
<th>Dose Frequency</th>
<th>Route of Administration</th>
<th>Regimen/Treatment Period</th>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>A, B, C, D</td>
<td>MK-3475</td>
<td>10 mg/kg</td>
<td>Q2W</td>
<td>IV infusion</td>
<td>Day 1 of each cycle</td>
<td>Experimental</td>
</tr>
<tr>
<td>B2</td>
<td>MK-3475</td>
<td>200 mg</td>
<td>Q3W</td>
<td>IV infusion</td>
<td>Day 1 of each cycle</td>
<td>Experimental</td>
</tr>
</tbody>
</table>

Trial treatment should begin on the day of randomization or as close as possible to the date on which treatment is allocated/assigned.
5.2.1 Dose Selection/Modification

5.2.1.1 Dose Selection

The rationale for selection of doses to be used in this trial is provided in Section 4.0 – Background and Rationale.

For Cohorts A, B, C, and D, the dose amount required to prepare the MK-3475 infusion solution will be based on the subject’s weight in kilograms (kg). Details on the dose calculation, preparation and administration are provided in the Pharmacy Manual.

For Cohort B2, subjects will be given 200 mg Q3W.

5.2.1.2 Dose Modification

Adverse events (both non-serious and serious) associated with pembrolizumab exposure may represent an immunologic etiology. These adverse events may occur shortly after the first dose or several months after the last dose of treatment. Pembrolizumab must be withheld for drug-related toxicities and severe or life-threatening AEs as per Table 3 below. See Section 5.6 and the Events of Clinical Interest Guidance Document for supportive care guidelines, including use of corticosteroids.
Table 3 Dose Modification and Toxicity Management Guidelines for Immune-related AEs Associated with Pembrolizumab

<table>
<thead>
<tr>
<th>Immune-related AEs</th>
<th>Toxicity grade or conditions (CTCAEv4.0)</th>
<th>Action taken to pembrolizumab</th>
<th>irAE management with corticosteroid and/or other therapies</th>
<th>Monitor and follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pneumonitis</td>
<td>Grade 2</td>
<td>Withhold</td>
<td>• Administer corticosteroids (initial dose of 1-2 mg/kg prednisone or equivalent) followed by taper</td>
<td>Monitor subjects for signs and symptoms of pneumonitis</td>
</tr>
<tr>
<td></td>
<td>Grade 3 or 4, or recurrent Grade 2</td>
<td>Permanently discontinue</td>
<td>• Evaluate subjects with suspected pneumonitis with radiographic imaging and initiate corticosteroid treatment</td>
<td></td>
</tr>
<tr>
<td>Diarrhea / Colitis</td>
<td>Grade 2 or 3</td>
<td>Withhold</td>
<td>• Add prophylactic antibiotics for opportunistic infections</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Grade 4</td>
<td>Permanently discontinue</td>
<td>• Monitor subjects for signs and symptoms of enterocolitis (ie, diarrhea, abdominal pain, blood or mucus in stool with or without fever) and of bowel perforation (ie, peritoneal signs and ileus).</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Subjects with ≥ Grade 2 diarrhea suspecting colitis should consider GI consultation and performing endoscopy to rule out colitis.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Subjects with diarrhea/colitis should be advised to drink liberal quantities of clear fluids. If sufficient oral fluid intake is not feasible, fluid and electrolytes should be substituted via IV infusion.</td>
<td></td>
</tr>
<tr>
<td>Immune-related AEs</td>
<td>Toxicity grade or conditions (CTCAEv4.0)</td>
<td>Action taken to pembrolizumab</td>
<td>irAE management with corticosteroid and/or other therapies</td>
<td>Monitor and follow-up</td>
</tr>
<tr>
<td>-------------------</td>
<td>---------------------------------------</td>
<td>---------------------------------</td>
<td>----------------------------------------------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>AST / ALT elevation or Increased bilirubin</td>
<td>Grade 2</td>
<td>Withhold</td>
<td>• Administer corticosteroids (initial dose of 0.5-1 mg/kg prednisone or equivalent) followed by taper</td>
<td>• Monitor with liver function tests (consider weekly or more frequently until liver enzyme value returned to baseline or is stable)</td>
</tr>
<tr>
<td></td>
<td>Grade 3 or 4</td>
<td>Permanently discontinue</td>
<td>• Administer corticosteroids (initial dose of 1-2 mg/kg prednisone or equivalent) followed by taper</td>
<td></td>
</tr>
<tr>
<td>Type 1 diabetes mellitus (T1DM) or Hyperglycemia</td>
<td>Newly onset T1DM or Grade 3 or 4 hyperglycemia associated with evidence of β-cell failure</td>
<td>Withhold</td>
<td>• Initiate insulin replacement therapy for subjects with T1DM • Administer anti-hyperglycemic in subjects with hyperglycemia</td>
<td>• Monitor subjects for hyperglycemia or other signs and symptoms of diabetes.</td>
</tr>
<tr>
<td>Hypophysitis</td>
<td>Grade 2</td>
<td>Withhold</td>
<td>• Administer corticosteroids and initiate hormonal replacements as clinically indicated.</td>
<td>• Monitor for signs and symptoms of hypophysitis (including hypopituitarism and adrenal insufficiency)</td>
</tr>
<tr>
<td></td>
<td>Grade 3 or 4</td>
<td>Withhold or permanently discontinue</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hyperthyroidism</td>
<td>Grade 2</td>
<td>Continue</td>
<td>• Treat with non-selective beta-blockers (eg, propranolol) or thionamides as appropriate</td>
<td>• Monitor for signs and symptoms of thyroid disorders.</td>
</tr>
<tr>
<td></td>
<td>Grade 3 or 4</td>
<td>Withhold or permanently discontinue</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypothyroidism</td>
<td>Grade 2-4</td>
<td>Continue</td>
<td>• Initiate thyroid replacement hormones (eg, levothyroxine or liothyronine) per standard of care</td>
<td>• Monitor for signs and symptoms of thyroid disorders.</td>
</tr>
<tr>
<td>Nephritis and Renal dysfunction</td>
<td>Grade 2</td>
<td>Withhold</td>
<td>• Administer corticosteroids (prednisone 1-2 mg/kg or equivalent) followed by taper.</td>
<td>• Monitor changes of renal function</td>
</tr>
<tr>
<td></td>
<td>Grade 3 or 4</td>
<td>Permanently discontinue</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immune-related AEs</td>
<td>Toxicity grade or conditions (CTCAEv4.0)</td>
<td>Action taken to pembrolizumab</td>
<td>irAE management with corticosteroid and/or other therapies</td>
<td>Monitor and follow-up</td>
</tr>
<tr>
<td>-------------------</td>
<td>--------------------------------------</td>
<td>-------------------------------</td>
<td>-------------------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>Myocarditis</td>
<td>Grade 1 or 2</td>
<td>Withhold</td>
<td>• Based on severity of AE administer corticosteroids</td>
<td>• Ensure adequate evaluation to confirm etiology and/or exclude other causes</td>
</tr>
<tr>
<td></td>
<td>Grade 3 or 4</td>
<td>Permanently discontinue</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All other immune-related AEs</td>
<td>Intolerable/ persistent Grade 2</td>
<td>Withhold</td>
<td>• Based on type and severity of AE administer corticosteroids</td>
<td>• Ensure adequate evaluation to confirm etiology and/or exclude other causes</td>
</tr>
<tr>
<td></td>
<td>Grade 3</td>
<td>Withhold or discontinue based on the type of event. Events that require discontinuation include and not limited to: Guillain-Barre Syndrome, encephalitis</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Grade 4 or recurrent Grade 3</td>
<td>Permanently discontinue</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. Withhold or permanently discontinue pembrolizumab is at the discretion of the investigator or treating physician.

**NOTE:**
For subjects with Grade 3 or 4 immune-related endocrinopathy where withhold of pembrolizumab is required, pembrolizumab may be resumed when AE resolves to ≤ Grade 2 and is controlled with hormonal replacement therapy or achieved metabolic control (in case of T1DM).
In case toxicity does not resolve to Grade 0-1 within 12 weeks after last infusion, trial treatment should be discontinued after consultation with the Sponsor. With investigator and Sponsor agreement, subjects with a laboratory adverse event still at Grade 2 after 12 weeks may continue treatment in the trial only if asymptomatic and controlled. For information on the management of adverse events, see Section 5.6.1.

5.2.2 Timing of Dose Administration

Trial treatment should be administered on Day 1 of each cycle after all procedures/assessments have been completed as detailed on the Trial Flow Chart (Section 6.0). Trial treatment may be administered up to 3 days before or after the scheduled Day 1 of each cycle due to administrative reasons.

All trial treatments will be administered on an outpatient basis.

Dosing interruptions are permitted in the case of medical / surgical events or logistical reasons not related to study therapy (e.g., elective surgery, unrelated medical events, subject vacation, and/or holidays). Subjects should be placed back on study therapy within 3 weeks of the scheduled interruption, unless otherwise discussed with the Sponsor. The reason for interruption should be documented in the patient's study record.

MK-3475 will be administered as a 30 minute IV infusion every 2 or 3 weeks depending on the Cohort the subjects have been enrolled into. Sites should make every effort to target infusion timing to be as close to 30 minutes as possible. However, given the variability of infusion pumps from site to site, a window of -5 minutes and +10 minutes is permitted (i.e., infusion time is 30 minutes: -5 min/+10 min).

The Pharmacy Manual contains specific instructions for MK-3475 dose calculation, reconstitution, preparation of the infusion fluid, and administration.

Treatment after initial evidence of radiologic disease progression

Immunotherapeutic agents such as MK-3475 may produce antitumor effects by potentiating endogenous cancer-specific immune responses. The response patterns seen with such an approach may extend beyond the typical time course of responses seen with cytotoxic agents, and can manifest as a clinical response after an initial increase in tumor burden or even the appearance of new lesions.

If radiologic imaging shows PD, tumor assessment should be repeated ≥ 4 weeks later in order to confirm PD with the option of continuing treatment per below while awaiting radiologic confirmation of progression. If repeat imaging shows a reduction in the tumor burden compared to the initial scan demonstrating PD, treatment may be continued as per treatment calendar. If repeat imaging confirms progressive disease, subjects will be discontinued from study therapy (exception noted in Section 7.1.2.5.1). In determining whether or not the tumor burden has increased or decreased, Investigators should consider all target lesions as well as non-target lesions (please refer to the Site Imaging Manual).
When feasible, subjects should not be discontinued until progression is confirmed; however, the decision to continue study treatment after the 1\textsuperscript{st} evidence of disease progression is at the Investigator’s discretion based on the clinical status of the subject as described in Table 4 below. Subjects may receive study treatment while waiting for confirmation of PD if they are clinically stable as defined by the following criteria:

- Absence of signs and symptoms (including worsening of laboratory values) indicating disease progression
- No decline in ECOG performance status
- Absence of rapid progression of disease
- Absence of progressive tumor at critical anatomical sites (e.g., cord compression) requiring urgent alternative medical intervention

Table 4 Imaging and Treatment After 1st Radiologic Evidence of PD

<table>
<thead>
<tr>
<th>Clinical Status</th>
<th>Imaging</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinically Stable</td>
<td>Repeat imaging at ≥ 4 weeks to confirm PD</td>
<td>May continue study treatment at the Investigator’s discretion while awaiting confirmatory scan</td>
</tr>
<tr>
<td>Clinically Unstable</td>
<td>Repeat imaging at ≥ 4 weeks to confirm PD if possible</td>
<td>Discontinue treatment</td>
</tr>
</tbody>
</table>

| Repeat scan confirms PD | No additional imaging required | Discontinue treatment (exception noted in Section 7.1.2.5.1) | No additional imaging required | N/A |

| Repeat scan shows SD, PR or CR | Continue regularly scheduled imaging assessments every 8 weeks | Continue study treatment at the Investigator’s discretion | Continue regularly scheduled imaging assessments every 8 weeks | May restart study treatment if condition has improved and/or clinically stable per Investigator’s discretion |

5.2.3 Trial Blinding/Masking

This is an open-label trial; therefore, the Sponsor, investigator and subject will know the treatment administered.

PD-L1 IHC biomarker testing is used to determine study eligibility for Cohorts A, B, C, and D and therefore the subject, investigator and SPONSOR will know the PD-L1 IHC result. Access to PD-L1 subject-level biomarker results for subjects enrolled in Cohort B2 will be
limited to an unblinded SPONSOR statistician, and unblinded SPONSOR statistical programmer who will be responsible for data review to ensure validity of results and summary reporting of clinical response by biomarker status, but who will have no other responsibilities associated with the study.

5.3 Randomization or Treatment Allocation

Subjects participating in this trial will be allocated to trial treatment by non-random assignment.

5.4 Stratification

No stratification based on age, sex or other characteristics will be used in this trial.

5.5 Concomitant Medications/Vaccinations (allowed & prohibited)

Medications or vaccinations specifically prohibited in the exclusion criteria are not allowed during the ongoing trial. Listed below are some specific restrictions for concomitant therapy or vaccination during the course of the trial. If there is a clinical indication for one of these or other medications or vaccinations specifically prohibited during the trial, discontinuation from trial therapy or vaccination may be required. The investigator should discuss any questions regarding this with the local Clinical Monitor. The final decision on any supportive therapy or vaccination rests with the investigator and/or the subject's primary physician. However, the decision to continue the subject on trial therapy or vaccination schedule requires the mutual agreement of the investigator, the Sponsor and the subject.

5.5.1 Acceptable Concomitant Medications

All treatments that the investigator considers necessary for a subject’s welfare may be administered at the discretion of the investigator in keeping with the community standards of medical care. All concomitant medication will be recorded on the case report form (CRF) including all prescription, over-the-counter (OTC), herbal supplements, and IV medications and fluids. If changes occur during the trial period, documentation of drug dosage, frequency, route, and date may also be included on the CRF.

All concomitant medications received within 28 days before the first dose of trial treatment and 30 days after the last dose of trial treatment should be recorded. Concomitant medications administered after 30 days after the last dose of trial treatment should be recorded for SAEs and ECIs as defined in Section 7.2.

5.5.2 Prohibited Concomitant Medications

Subjects are prohibited from receiving the following therapies during the Screening and Treatment Phase (including retreatment for post-complete response relapse) of this trial:

- Antineoplastic systemic chemotherapy or biological therapy
- Immunotherapy not specified in this protocol
- Chemotherapy not specified in this protocol
- Investigational agents other than MK-3475
- Radiation therapy
  - Note: Radiation therapy to a symptomatic solitary lesion or to the brain may be allowed after consultation with Sponsor.
- Live vaccines within 30 days prior to the first dose of trial treatment and while participating in the trial. Examples of live vaccines include, but are not limited to, the following: measles, mumps, rubella, chicken pox, yellow fever, rabies, BCG, and typhoid (oral) vaccine. Seasonal influenza vaccines for injection are generally inactivated flu vaccines and are allowed; however intranasal influenza vaccines (e.g., Flu-Mist®) are live attenuated vaccines, and are not allowed.
- Glucocorticoids for any purpose other than to modulate acute symptoms from an adverse event. The use of physiologic doses for subjects requiring ongoing corticosteroids, may be approved after consultation with the Sponsor.

Subjects who, in the assessment by the investigator, require the use of any of the aforementioned treatments for clinical management should be removed from the trial. Subjects may receive other medications that the investigator deems to be medically necessary.

The Exclusion Criteria describes other medications which are prohibited in this trial.

There are no prohibited therapies during the Post-Treatment Follow-up Phase.

### 5.6 Rescue Medications & Supportive Care

#### 5.6.1 Supportive Care Guidelines

Subjects should receive appropriate supportive care measures as deemed necessary by the treating investigator. Suggested supportive care measures for the management of adverse events with potential immunologic etiology are outlined below. Where appropriate, these guidelines include the use of oral or intravenous treatment with corticosteroids as well as additional anti-inflammatory agents if symptoms do not improve with administration of corticosteroids. Note that several courses of steroid tapering may be necessary as symptoms may worsen when the steroid dose is decreased. For each disorder, attempts should be made to rule out other causes such as metastatic disease or bacterial or viral infection, which might require additional supportive care. The treatment guidelines are intended to be applied when the investigator determines the events to be related to pembrolizumab.

Note: if after the evaluation the event is determined not to be related, the investigator is instructed to follow the ECI reporting guidance. Refer to Section 5.2.1 for dose modification.

It may be necessary to perform conditional procedures such as bronchoscopy, endoscopy, or skin photography as part of evaluation of the event.
- **Pneumonitis:**
  
  o For **Grade 2 events**, treat with systemic corticosteroids. When symptoms improve to Grade 1 or less, steroid taper should be started and continued over no less than 4 weeks.
  
  o For **Grade 3-4 events**, immediately treat with intravenous steroids. Administer additional anti-inflammatory measures, as needed.
  
  o Add prophylactic antibiotics for opportunistic infections in the case of prolonged steroid administration.

- **Diarrhea/Colitis:**

  Subjects should be carefully monitored for signs and symptoms of enterocolitis (such as diarrhea, abdominal pain, blood or mucus in stool, with or without fever) and of bowel perforation (such as peritoneal signs and ileus).

  o All subjects who experience diarrhea/colitis should be advised to drink liberal quantities of clear fluids. If sufficient oral fluid intake is not feasible, fluid and electrolytes should be substituted via IV infusion. For Grade 2 or higher diarrhea, consider GI consultation and endoscopy to confirm or rule out colitis.
  
  o For **Grade 2 diarrhea/colitis** that persists greater than 3 days, administer oral corticosteroids.
  
  o For **Grade 3 or 4 diarrhea/colitis** that persists > 1 week, treat with intravenous steroids followed by high dose oral steroids.
  
  o When symptoms improve to Grade 1 or less, steroid taper should be started and continued over no less than 4 weeks.

- **Type 1 diabetes mellitus (if new onset, including diabetic ketoacidosis [DKA]) or ≥ Grade 3 Hyperglycemia, if associated with ketosis (ketonuria) or metabolic acidosis (DKA)**

  o For **T1DM** or **Grade 3-4 Hyperglycemia**
    - Insulin replacement therapy is recommended for Type I diabetes mellitus and for Grade 3-4 hyperglycemia associated with metabolic acidosis or ketonuria.
    - Evaluate patients with serum glucose and a metabolic panel, urine ketones, glycosylated hemoglobin, and C-peptide.
- **Hypophysitis:**
  - For **Grade 2** events, treat with corticosteroids. When symptoms improve to Grade 1 or less, steroid taper should be started and continued over no less than 4 weeks. Replacement of appropriate hormones may be required as the steroid dose is tapered.
  - For **Grade 3-4** events, treat with an initial dose of IV corticosteroids followed by oral corticosteroids. When symptoms improve to Grade 1 or less, steroid taper should be started and continued over no less than 4 weeks. Replacement of appropriate hormones may be required as the steroid dose is tapered.

- **Hyperthyroidism or Hypothyroidism:**

  Thyroid disorders can occur at any time during treatment. Monitor patients for changes in thyroid function (at the start of treatment, periodically during treatment, and as indicated based on clinical evaluation) and for clinical signs and symptoms of thyroid disorders.

  - **Grade 2** hyperthyroidism events (and **Grade 2-4** hypothyroidism):
    - In hyperthyroidism, non-selective beta-blockers (e.g. propranolol) are suggested as initial therapy.
    - In hypothyroidism, thyroid hormone replacement therapy, with levothyroxine or liothyronine, is indicated per standard of care.
  - **Grade 3-4** hyperthyroidism
    - Treat with an initial dose of IV corticosteroid followed by oral corticosteroids. When symptoms improve to Grade 1 or less, steroid taper should be started and continued over no less than 4 weeks. Replacement of appropriate hormones may be required as the steroid dose is tapered.

- **Hepatic:**
  - For **Grade 2** events, monitor liver function tests more frequently until returned to baseline values (consider weekly).
    - Treat with IV or oral corticosteroids
  - For **Grade 3-4** events, treat with intravenous corticosteroids for 24 to 48 hours.
  - When symptoms improve to Grade 1 or less, a steroid taper should be started and continued over no less than 4 weeks.
- **Renal Failure or Nephritis:**
  - For **Grade 2** events, treat with corticosteroids.
  - For **Grade 3-4** events, treat with systemic corticosteroids.
  - When symptoms improve to Grade 1 or less, steroid taper should be started and continued over no less than 4 weeks.

- **Management of Infusion Reactions:** Signs and symptoms usually develop during or shortly after drug infusion and generally resolve completely within 24 hours of completion of infusion.

**Table 5** below shows treatment guidelines for subjects who experience an infusion reaction associated with administration of pembrolizumab (MK-3475).

**Table 5** Infusion Reaction Treatment Guidelines

<table>
<thead>
<tr>
<th>NCI CTCAE Grade</th>
<th>Treatment</th>
<th>Premedication at subsequent dosing</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Grade 1</strong></td>
<td>Increase monitoring of vital signs as medically indicated until the subject is deemed medically stable in the opinion of the investigator.</td>
<td>None</td>
</tr>
<tr>
<td>Mild reaction; infusion interruption not indicated; intervention not indicated</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| **Grade 2** | Stop Infusion and monitor symptoms. Additional appropriate medical therapy may include but is not limited to: | Subject may be premedicated 1.5h (± 30 minutes) prior to infusion of pembrolizumab (MK-3475) with: |
| Requires infusion interruption but responds promptly to symptomatic treatment (e.g., antihistamines, NSAIDS, narcotics, IV fluids); prophylactic medications indicated for < =24 hrs | IV fluids Antihistamines NSAIDS Acetaminophen Narcotics Increase monitoring of vital signs as medically indicated until the subject is deemed medically stable in the opinion of the investigator. If symptoms resolve within one hour of stopping drug infusion, the infusion may be restarted at 50% of the original infusion rate (e.g., from 100 mL/hr to 50 mL/hr). Otherwise dosing will be held until symptoms resolve and the subject should be premedicated for the next scheduled dose. Subjects who develop Grade 2 toxicity despite adequate premedication should be permanently discontinued from further trial treatment administration. | | Diphenhydramine 50 mg po (or equivalent dose of antihistamine). Acetaminophen 500-1000 mg po (or equivalent dose of antipyretic). |
### 5.7 Diet/Activity/Other Considerations

#### 5.7.1 Diet

Subjects should maintain a normal diet unless modifications are required to manage an AE such as diarrhea, nausea or vomiting.

#### 5.7.2 Contraception

MK-3475 may have adverse effects on a fetus in utero. Furthermore, it is not known if MK-3475 has transient adverse effects on the composition of sperm. Non-pregnant, non-breast-feeding women may be enrolled if they are willing to use 2 methods of birth control or are considered highly unlikely to conceive. Highly unlikely to conceive is defined as 1) surgically sterilized, or 2) postmenopausal (a woman who is ≥45 years of age and has not had menses for greater than 1 year will be considered postmenopausal), or 3) not heterosexually active for the duration of the study. The two birth control methods can either be two barrier methods or a barrier method plus a hormonal method to prevent pregnancy. Subjects should start using birth control from study Visit 1 throughout the study period up to 120 days after the last dose of study therapy.

The following are considered adequate barrier methods of contraception: diaphragm, condom (by the partner), copper intrauterine device, sponge, or spermicide. Appropriate hormonal contraceptives will include any registered and marketed contraceptive agent that contains an estrogen and/or a progestational agent (including oral, subcutaneous, intrauterine, or intramuscular agents).
Subjects should be informed that taking the study medication may involve unknown risks to
the fetus (unborn baby) if pregnancy were to occur during the study. In order to participate
in the study they must adhere to the contraception requirement (described above) for the
duration of the study and during the follow-up period defined in section 7.2.2-Reporting of
Pregnancy and Lactation to the Sponsor. If there is any question that a subject will not
reliably comply with the requirements for contraception, that subject should not be entered
into the study.

5.7.3 Use in Pregnancy

If a subject inadvertently becomes pregnant while on treatment with MK-3475, the subject
will immediately be removed from the study. The site will contact the subject at least
monthly and document the subject’s status until the pregnancy has been completed or
terminated. The outcome of the pregnancy will be reported to the Sponsor without delay and
within 24 hours if the outcome is a serious adverse experience (e.g., death, abortion,
congenital anomaly, or other disabling or life-threatening complication to the mother or
newborn). The study investigator will make every effort to obtain permission to follow the
outcome of the pregnancy and report the condition of the fetus or newborn to the Sponsor.
If a male subject impregnates his female partner the study personnel at the site must be
informed immediately and the pregnancy reported to the Sponsor and followed as described
above and in Section 7.2.2.

5.7.4 Use in Nursing Women

It is unknown whether MK-3475 is excreted in human milk. Since many drugs are excreted
in human milk, and because of the potential for serious adverse reactions in the nursing
infant, subjects who are breast-feeding are not eligible for enrollment.

5.8 Subject Withdrawal/Discontinuation Criteria

Subjects may withdraw consent at any time for any reason or be dropped from the trial at the
discretion of the investigator should any untoward effect occur. In addition, a subject may be
withdrawn by the investigator or the Sponsor if enrollment into the trial is inappropriate, the
trial plan is violated, or for administrative and/or other safety reasons. Specific details
regarding discontinuation or withdrawal procedures; including specific details regarding
withdrawal from Future Biomedical Research, are provided in Section 7.1.4 – Other
Procedures.

A subject must be discontinued from the trial for any of the following reasons:

- The subject or legal representative (such as a parent or legal guardian) withdraws
  consent.
A subject must be discontinued from treatment (but may continue to be monitored in the post-treatment follow-up portion of the trial) for any of the following reasons:

- The subject or legal representative (such as a parent or legal guardian) withdraws consent for treatment
- Confirmed radiographic disease progression
  
  Note: For unconfirmed radiographic disease progression, please see Section 5.2.2.
  
  Note: A subject may be granted an exception to continue on treatment with confirmed radiographic progression if clinically stable or clinically improved, please see Section 7.1.2.5.1.
- Unacceptable adverse experiences as described in Section 5.2.1.2
- Intercurrent illness that prevents further administration of treatment
- Investigator’s decision to withdraw the subject
- The subject has a confirmed positive serum pregnancy test
- Noncompliance with trial treatment or procedure requirements
- The subject is lost to follow-up
- Completed 24 months of treatment with MK-3475
  
  Note: 24 months of study medication is calculated from the date of first dose. Subjects who stop MK-3475 after 24 months may be eligible for up to one year of additional study treatment if they progress after stopping study treatment provided they meet the requirements detailed in Section 7.1.5.2.1.
- Administrative reasons

The End of Treatment and Follow-up visit procedures are listed in Section 6 (Protocol Flow Chart) and Section 7.1.5 (Visit Requirements). After the end of treatment, each subject will be followed for 30 days for adverse event monitoring (serious adverse events will be collected for 90 days after the end of treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier, as described in Section 7.2.3.1). Subjects who discontinue for reasons other than progressive disease will have post-treatment follow-up for disease status until disease progression, initiating a non-study cancer treatment, withdrawing consent or becoming lost to follow-up. After documented disease progression each subject will be followed by telephone for overall survival until death, withdrawal of consent, or the end of the study, whichever occurs first.
5.8.1 Discontinuation of Study Therapy after CR

Discontinuation of treatment may be considered for subjects who have attained a confirmed CR that have been treated for at least 24 weeks with MK-3475 and had at least two treatments with MK-3475 beyond the date when the initial CR was declared. Subjects who then experience radiographic disease progression may be eligible for up to one year of additional treatment with MK-3475 at the discretion of the investigator if no cancer treatment was administered since the last dose of MK-3475, the subject meets the safety parameters listed in the Inclusion/Exclusion criteria, and the trial is open. Subjects will resume therapy at the same dose and schedule at the time of initial discontinuation. Additional details are provided in Section 7.1.5.2.1.

5.9 Subject Replacement Strategy

Additional subjects may be enrolled in a given cohort to ensure that the required number of evaluable subjects in each cohort is achieved. A subject that discontinues the trial for progressive disease or a drug-related AE will not be replaced and will be counted in the evaluable population of subjects for the respective cohort. Further details are provided in Section 8.1.3.

5.10 Beginning and End of the Trial

The study begins when the first subject signs the informed consent (either pre-screening consent or main study consent). The end of the study may be designated as the time point when all subjects have discontinued the study or are a minimum of 6 months post initial study medication administration or all subjects still on study have received their 24-week scan (i.e. completed their third 8-week efficacy assessment). If, by the end of the study, there remains at least 1 subject still on study treatment for at least 6 months, the subject(s) may enter additional treatment cycles. At this point a database lock of the trial may occur to allow the analysis of the study data. Any remaining subjects may continue to receive study medication and be seen by the investigator per usual standard of care for this subject population. In addition, the investigator will be expected to monitor for and report any serious adverse events, events of clinical interest, and pregnancies, as detailed in Section 7.2.3 (Serious Adverse Experiences). The subject is considered on study until such time that he/she meets any of the discontinuation criteria and written notification is given to the Sponsor. Upon study completion, subjects are discontinued and enrolled in a pembrolizumab extension trial.

5.11 Clinical Criteria for Early Trial Termination

Early trial termination will be the result of the criteria specified below:

1. Quality or quantity of data recording is inaccurate or incomplete

2. Poor adherence to protocol and regulatory requirements
3. Incidence or severity of adverse drug reaction in this or other studies indicates a potential health hazard to subjects

4. Plans to modify or discontinue the development of the study drug

In the event of Sponsor decision to no longer supply study drug, ample notification will be provided so that appropriate adjustments to subject treatment can be made.
### 6.0 TRIAL FLOW CHART

#### 6.1 Study Flow Chart for Cohorts A, B, C and D (10 mg/kg Q2W dosing)

<table>
<thead>
<tr>
<th>Trial Period:</th>
<th>Screening Phase</th>
<th>Treatment Cycles</th>
<th>End of Treatment</th>
<th>Post-Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pre-screening</td>
<td>Main Study</td>
<td>To be repeated beyond 8 cycles</td>
<td>Discon</td>
</tr>
<tr>
<td></td>
<td>(Visit 1)</td>
<td>Screening (Visit 2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 2 3 4 5 6 7 8</td>
<td>± 3 ± 3 ± 3 ± 3 ± 3 ± 3 ± 3</td>
<td>At time of Discon</td>
<td>30 days post discon</td>
</tr>
<tr>
<td>Scheduling Window (Days):</td>
<td>-28 to -1</td>
<td></td>
<td></td>
<td></td>
</tr>
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</table>

#### Administrative Procedures

- Pre-screening Consent
- Informed Consent
- Informed Consent for Future Biomedical Research
- Inclusion/Exclusion Criteria
- Subject Identification Card
- Demographics and Medical History
- Prior and Concomitant Medication Review
- Trial Treatment Administration
- Post-study anticancer therapy status
- Survival Status

#### Clinical Procedures/Assessments

- Review Adverse Events
- Full Physical Examination
- Directed Physical Examination
- Vital Signs and Weight
- ECOG Performance Status

#### Laboratory Procedures/Assessments: analysis performed by local laboratory

- Pregnancy Test – Urine or Serum
- PT/INR and aPTT
- CBC with Differential
- Comprehensive Chemistry Panel
- Urinalysis
- T3, FT4 and TSH

---
<table>
  <thead>
    <tr>
      <th>Trial Period:</th>
      <th>Screening Phase</th>
      <th>Treatment Cycles<sup>a</sup></th>
      <th>End of Treatment</th>
      <th>Post-Treatment</th>
    </tr>
  </thead>
  <tbody>
    <tr>
      <td>Treatment Cycle/Title:</td>
      <td>Pre-screening (Visit 1)<sup>b</sup> Main Study Screening (Visit 2)<sup>c</sup></td>
      <td>1 2 3 4 5 6 7 8</td>
      <td><span>To be repeated beyond 8 cycles</span></td>
      <td>Post-Treatment Safety Follow-up</td>
    </tr>
    <tr>
      <td>Scheduling Window (Days)<sup>d</sup>:</td>
      <td>-28 to -1 ± 3 ± 3 ± 3 ± 3 ± 3 ± 3 ± 3 ± 3</td>
      <td><span>At time of Discon</span></td>
      <td>30 days post discon</td>
      <td>Every 8 weeks post discon</td>
    </tr>
  </tbody>
</table>

**Laboratory Procedures/Assessments:** analysis performed by central laboratory

- **Anti-MK-3475 Antibodies:**
  - X<sup>t</sup> X<sup>t</sup> X<sup>t</sup> X<sup>t</sup> X<sup>t</sup> X<sup>t</sup>
- **Pharmacokinetics:**
  - X<sup>t,u</sup> X<sup>t</sup> X<sup>t</sup> X<sup>t</sup> X<sup>t</sup> X<sup>t</sup>
- **Blood for Future Biomedical Research:**
  - X

**Efficacy Measurements**

<table>
<thead>
<tr>
<th>Tumor Imaging&lt;sup&gt;x&lt;/sup&gt;</th>
<th>X</th>
<th>X</th>
<th>X</th>
<th>X&lt;sup&gt;y&lt;/sup&gt;</th>
<th>X&lt;sup&gt;d&lt;/sup&gt;</th>
</tr>
</thead>
</table>

**Tumor Biopsies/Archival Tissue Collection/Correlative Studies Blood**

<table>
<thead>
<tr>
<th>Archival Tissue Collection&lt;sup&gt;z&lt;/sup&gt;</th>
<th>X&lt;sup&gt;aa&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cohort B (H/N) and Cohort D (gastric cancer) Tumor Tissue Collection&lt;sup&gt;z&lt;/sup&gt;</td>
<td>X&lt;sup&gt;aa,bb&lt;/sup&gt; X&lt;sup&gt;bb&lt;/sup&gt; X&lt;sup&gt;cc&lt;/sup&gt;</td>
</tr>
<tr>
<td>Cohort A (TNBC) and Cohort C (urothelial tract cancer) Tumor Tissue Collection&lt;sup&gt;z&lt;/sup&gt;</td>
<td>X&lt;sup&gt;aa,dd&lt;/sup&gt; X&lt;sup&gt;dd&lt;/sup&gt; X&lt;sup&gt;cc&lt;/sup&gt;</td>
</tr>
<tr>
<td>Correlative Studies Blood Collection</td>
<td>X&lt;sup&gt;ee&lt;/sup&gt; X&lt;sup&gt;ee&lt;/sup&gt; X&lt;sup&gt;ee&lt;/sup&gt;</td>
</tr>
</tbody>
</table>
a. In general, assessments/procedures are to be performed on Day 1 and prior to the first dose of treatment for each cycle unless otherwise specified. Treatment cycles are 2 weeks. Imaging should always be performed every 8 weeks (± 7 days) regardless of any treatment delays.
b. At the pre-screening visit, subjects will sign the pre-screening consent and submit an archival sample for PD-L1 characterization.
c. Subjects who submit an archival tumor sample at the prescreening visit and are found to be PD-L1 positive will continue to the screening portion of the study. Subjects who do not have an archival sample will go directly to the screening phase of the study.
d. In subjects who discontinue study therapy without documented disease progression, every effort should be made to continue monitoring their disease status by radiologic imaging every 8 weeks (± 7 days) until (1) the start of new anti-cancer treatment, (2) documented disease progression, (3) death, or (4) the end of the study, whichever occurs first. If imaging assessment beyond the protocol-required imaging time frame is necessary, the reason for the extended imaging schedule should be documented in the subject’s chart and recorded in the Oncologic Response-Solid Tumor (ORST) eCRF for each imaging assessment completed within the extended time frame. The maximum imaging time frame allowed is no greater than twice the imaging time frame noted (ie, for every 8 weeks’ imaging, the maximum imaging schedule allowance is every 16 weeks). Every effort should be made to target the protocol-required imaging schedule as closely as possible; however, in the interest of the subject’s continued participation in the follow-up phase of the study, the extended imaging time frame may be considered at the discretion of the investigator.
e. After the start of new anti-cancer treatment or documented disease progression, the subject should be contacted by telephone approximately every 12 weeks to assess for
f. In general, the window for each visit is ± 3 days unless otherwise noted.
g. Pre-screening informed consent must be obtained prior to sending an archival sample to the lab for characterization. Subjects that do not have archival tissue available to send must sign the main study consent prior to undergoing a newly obtained biopsy.
h. Written consent must be obtained prior to performing any protocol specific procedure. Results of a test performed prior to the subject signing consent as part of routine clinical management are acceptable in lieu of a screening test if performed within the specified time frame (e.g., within 28 days prior to the first dose of trial treatment). Screening number will be assigned when the study informed consent is signed.
i. Signing the informed consent for future biomedical research (FBR) samples is optional. Detailed instructions for the collection and management of specimens for FBR are provided in the Procedures Manual and Section 12.2.
j. Prior medications – Record all medications taken within 28 days of screening visit. Concomitant medications – Enter new medications started during the trial through the
k. AEs and laboratory safety measurements will be graded per NCI CTCAE version 4.0. All AEs, whether gradable by CTCAE or not, will also be evaluated for seriousness.
l. Record all AEs occurring within 30 days after the last dose of trial treatment. Report all SAEs (related and unrelated to trial treatment) and ECIs occurring up until 90 days after the last dose of trial treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier. Afterwards, report only SAEs and ECIs that are related to trial treatment.
m. To be repeated every 4 cycles after cycle 5.

o. Vital signs to include temperature, pulse, respiratory rate, weight and blood pressure. Height will be measured at visit 2 only.
p. Coagulation factors (PT/INR and aPTT) should be tested as part of the screening procedures for all subjects. Any subject receiving anticoagulant therapy should have coagulation factors monitored closely throughout the trial.
q. Laboratory tests for screening are to be performed within 10 days prior to the first dose of trial treatment. See Section 7.1.3 for details regarding laboratory tests.
r. After Cycle 1, lab samples can be collected up to 72 hours prior to the scheduled time point. See Section 7.1.3 for details regarding laboratory tests.
s. Unresolved abnormal labs that are drug related AEs should be followed until resolution. Labs do not need to be repeated after the end of treatment if labs are within normal range.
| t. | Pre-dose trough and post-dose peak PK samples will be collected at Cycles 1 and 2. Pre-dose trough samples only will be collected every 4 cycles starting with Cycle 5 and through Cycle 37, 30 days after discontinuation of study drug, and 3 months and 6 months after discontinuation of study drug (or until the subject starts new anti-neoplastic therapy). All trough samples should be drawn within 24 hours before infusion of MK-3475. All peak samples should be drawn within 30 minutes after the end of the infusion. Anti-MK-3475 antibodies should be drawn with all pre-dose trough PK samples, the 30 day discontinuation draw and 3 months and 6 months after discontinuation of study drug (or until the subject starts new anti-neoplastic therapy). Procedures for sample collection are described in the Procedures Manual. |
| u. | An additional single PK sample should be drawn between 24 to 96 hours after Cycle 1 dosing. |
| v. | Informed consent for future biomedical research samples must be obtained before the DNA sample. DNA sample for analysis should be obtained predose, on Day 1 (or with the next scheduled blood draw) as the last sample drawn or at a later date as soon as the informed consent is obtained. Detailed instructions for the collection and management of specimens for FBR are provided in the Procedures Manual and Section 12.2. |
| w. | The initial tumor imaging will be performed within 28 days prior to the first dose of trial treatment. Scans performed as part of routine clinical management are acceptable for use as the screening scan if they are of diagnostic quality and performed within 28 days prior to the first dose of trial treatment. On-study imaging will be performed every 8 weeks (± 7 days) after the first dose of trial treatment or more frequently if clinically indicated. Imaging timing should follow calendar days and should not be adjusted for delays in cycle starts or extension of MK-3475 cycle frequencies. The same imaging technique should be used in a subject throughout the trial. Local reading (investigator assessment with site radiology reading) will be used to determine eligibility and for subject management; Sponsor will collect radiological assessments for retrospective analysis by an independent central radiology review vendor. The processes for image collection and transmission to the independent central radiology review vendor are in the Site Imaging Manual. |
| x. | Per the modified RECIST 1.1 used in this protocol, if imaging shows progressive disease, the imaging assessment should be performed at a minimum of 4 weeks later in order to confirm progressive disease as described in Section 4.2.3.1. Please refer to the Procedure Manual for additional details on modifications to RECIST. |
| y. | In subjects who discontinue study therapy without confirmed disease progression, a radiological evaluation should be performed at the time of treatment discontinuation (i.e. date of discontinuation ± 4 week window). If a previous scan was obtained within 4 weeks prior to the date of discontinuation, then a scan at treatment discontinuation isn’t mandatory. |
| z. | Detailed instructions for tissue collection, processing and shipment are provided in the Procedures Manual. If the subject signs the Future Biomedical Research (FBR) consent, any leftover tissue that would ordinarily be discarded at the end of the main study will be retained for FBR. |
| aa. | Baseline tumor tissue for biomarker analysis from an archival tissue sample or newly obtained core or excisional biopsy (FNA not adequate) of a tumor lesion not previously irradiated must be provided and received by the independent central radiology review vendor before enrollment for characterization of PD-L1 status. These samples are not required to be obtained within 28 days of enrollment. Subjects with H/N cancer may provide tissue from a previously irradiated lesion. |
| bb. | Newly obtained tumor biopsy is required for subjects enrolled into Cohort B (H/N cancer) and Cohort D (gastric cancer) of the study. Tumor biopsies that are inaccessible or contraindicated due to subject safety concerns are exempt from this requirement. The pre-dose newly obtained biopsy is not required for PD-L1 characterization and may be performed just prior to the first dose of study treatment after all eligibility criteria has been met. |
| cc. | Tumor biopsy for clinically stable subjects at treatment discontinuation is highly encouraged. |
| dd. | Tumor biopsy is highly encouraged for all subjects. If activity within the Cohort is observed (at least 2 responders within the Cohort) the tumor biopsy will become mandatory. Tumor biopsies that are inaccessible or contraindicated due to subject safety concerns are exempt from this requirement. The pre-dose newly obtained biopsy is not required for PD-L1 characterization and may be performed just prior to the first dose of study treatment after all eligibility criteria has been met. |
| ee. | Blood for correlative studies should be collected prior to Cycle 1, at Cycle 5 and again at treatment discontinuation. |
| ff. | Following Cycle 8, the directed physical exam is only required at Cycle 11, 15, 19, and every 4 cycles thereafter. |
| gg. | Following Cycle 8, the ECOG performance status should be determined only in conjunction with a protocol-specified full or directed physical exam (Cycle 9, 11, 13, 15, 17, 19 and every 2 cycles thereafter. |
### 6.2 Study Flow Chart for Cohort B2 (200 mg Q3W dosing)

<table>
<thead>
<tr>
<th>Trial Period:</th>
<th>Screening Phase</th>
<th>Treatment Cycles&lt;sup&gt;a&lt;/sup&gt;</th>
<th>End of Treatment</th>
<th>Post-Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>To be repeated beyond 6 cycles</td>
<td>Post-Treatment Safety Follow-up</td>
<td>Follow Up Visits&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Treatment Cycle/Title:</td>
<td>Main Study Screening (Visit 2)</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Scheduling Window (Days)&lt;sup&gt;d&lt;/sup&gt;:</td>
<td>-28 to -1</td>
<td>± 3</td>
<td>± 3</td>
<td>± 3</td>
</tr>
</tbody>
</table>

- **Informed Consent**: X<sup>e</sup>
- **Informed Consent for Future Biomedical Research**: X<sup>e</sup>
- **Inclusion/Exclusion Criteria**: X
- **Subject Identification Card**: X
- **Demographics and Medical History**: X
- **Prior and Concomitant Medication Review**: X<sup>e</sup> X X X X X X X X<sup>f</sup>
- **Trial Treatment Administration**: X X X X X X X X X X
- **Post-study anticancer therapy status**: X
- **Survival Status**: X
- **Review Adverse Events**: X X X X X X X X X X X X X X
- **Full Physical Examination**: X X
- **Directed Physical Examination**: X X X X X<sup>e</sup> X<sup>e</sup> X X
- **Vital Signs and Weight**: X X X X X X X X
- **ECOG Performance Status**: X X X X X X X X
- **Pregnancy Test – Urine or Serum β-HCG**: X
- **PT/INR and aPTT**: X<sup>e</sup>
- **CBC with Differential**: X<sup>e</sup> X X X X X X X X<sup>e</sup>
- **Comprehensive Chemistry Panel**: X<sup>e</sup> X X X X X X X
- **Urinalysis**: X<sup>e</sup> X<sup>e</sup> X<sup>e</sup>
- **T3, TT4 and TSH**: X<sup>e</sup> X<sup>e</sup> X
- **Blood for Future Biomedical Research**: X
- **Tumor Imaging**: X X X X X X X X X X X X X
- **Archival Tissue Collection**: X<sup>e</sup>
- **Newly Obtained Biopsy Collection**: X<sup>e</sup> X<sup>e</sup> X<sup>e</sup>
- **Correlative Studies Blood Collection**: X<sup>e</sup> X<sup>e</sup> X<sup>e</sup> X<sup>e</sup>
a. In general, assessments/procedures are to be performed on Day 1 and prior to the first dose of treatment for each cycle unless otherwise specified. Treatment cycles are 3 weeks; Imaging should always be performed every 8 weeks (±7 days) regardless of any treatment delays.
b. In subjects who discontinue study therapy without documented disease progression, every effort should be made to continue monitoring their disease status by radiologic imaging every 8 weeks (±7 days) until (1) the start of new anti-cancer treatment, (2) documented disease progression, (3) death, or (4) the end of the study, whichever occurs first. If imaging assessment beyond the protocol-required imaging time frame is necessary, the reason for the extended imaging schedule should be documented in the subject’s chart and recorded in the ORST eCRF for each imaging assessment completed within the extended time frame. The maximum imaging time frame allowed is no greater than twice the imaging time frame noted (i.e., for every 8 weeks’ imaging, the maximum imaging schedule allowance is every 16 weeks). Every effort should be made to target the protocol-required imaging schedule as closely as possible; however, in the interest of the subject’s continued participation in the follow-up phase of the study, the extended imaging time frame may be considered at the discretion of the investigator.
c. After the start of new anti-cancer treatment or documented disease progression, the subject should be contacted by telephone approximately every 12 weeks to assess for clinical management.
d. In general, the window for each visit is ±3 days unless otherwise noted.
e. Written consent must be obtained prior to performing any protocol specific procedure. Results of a test performed prior to the subject signing consent as part of routine clinical management are acceptable in lieu of a screening test if performed within the specified time frame (e.g., within 28 days prior to the first dose of trial treatment). Screening number will be assigned when the study informed consent is signed.
f. Signing the informed consent for future biomedical research (FBR) samples is optional. Detailed instructions for the collection and management of specimens for FBR are provided in the Procedures Manual and Section 12.2.
g. Prior medications – Record all medications taken within 28 days of screening visit. Concomitant medications – Enter new medications started during the trial through the Safety Follow-up visit. Record all medications taken for SAEs as defined in Section 7.2.
h. AEs and laboratory safety measurements will be graded per NCI CTCAE version 4.0. All AEs, whether gradable by CTCAE or not, will also be evaluated for seriousness.
i. Record all AEs occurring within 30 days after the last dose of trial treatment. Report all SAEs (related and unrelated to trial treatment) and ECIs occurring up until 90 days after the last dose of trial treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier. Afterwards, report only SAEs and ECIs that are related to trial treatment.
j. To be repeated every 3 cycles after cycle 4.
k. Following Cycle 6, the directed physical exam is only required as clinically appropriate as long as a physical exam is performed every 6 weeks.
l. Vital signs to include temperature, pulse, respiratory rate, weight and blood pressure. Height will be measured at the screening visit (visit 2) only.
m. Following Cycle 6, the ECOG performance status should be determined only in conjunction with a protocol-specified full or directed physical exam.
n. For women of reproductive potential, a urine pregnancy test should be performed within 72 hours prior to first dose of trial treatment. If urine pregnancy results cannot be confirmed as negative, a serum pregnancy test performed by the local study site laboratory will be required. Pregnancy tests (serum and/or urine tests) should be repeated if required by local guidelines.
o. Coagulation factors (PT/INR and aPTT) should be tested as part of the screening procedures for all subjects. Any subject receiving anticoagulant therapy should have coagulation factors monitored closely throughout the trial.
p. Laboratory tests for screening are to be performed within 10 days prior to the first dose of trial treatment. See Section 7.1.3 for details regarding laboratory tests.
q. After Cycle 1, lab samples can be collected up to 72 hours prior to the scheduled time point. See Section 7.1.3 for details regarding laboratory tests.
r. Unresolved abnormal labs that are drug related AEs should be followed until resolution. Labs do not need to be repeated after the end of treatment if labs are within normal range.
s. Informed consent for future biomedical research samples must be obtained before the DNA sample. DNA sample for analysis should be obtained predose, on Day 1 (or with the next scheduled blood draw) as the last sample drawn or at a later date as soon as the informed consent is obtained. Detailed instructions for the collection and management of specimens for FBR are provided in the Procedures Manual and Section 12.2.
1. The initial tumor imaging will be performed within 28 days prior to the first dose of trial treatment. Scans performed as part of routine clinical management are acceptable for use as the screening scan if they are of diagnostic quality and performed within 28 days prior to the first dose of trial treatment. On-study imaging will be performed every 8 weeks (±7 days) after the first dose of trial treatment or more frequently if clinically indicated. Imaging timing should follow calendar days and should not be adjusted for delays in cycle starts or extension of MK-3475 cycle frequencies. The same imaging technique should be used in a subject throughout the trial. Local reading (investigator assessment with site radiology reading) will be used to determine eligibility and for subject management; Sponsor will collect radiological assessments for retrospective analysis by an independent central radiology review vendor. The processes for image collection and transmission to the independent central radiology review vendor are in the Site Imaging Manual.

u. Per the modified RECIST 1.1 used in this protocol, if imaging shows progressive disease, the imaging assessment should be performed at a minimum of 4 weeks later in order to confirm progressive disease as described in Section 4.2.3.1. Please refer to the Procedure Manual for additional details on modifications to RECIST.

v. In subjects who discontinue study therapy without confirmed disease progression, a radiological evaluation should be performed at the time of treatment discontinuation (i.e. date of discontinuation ± 4 week window). If a previous scan was obtained within 4 weeks prior to the date of discontinuation, then a scan at treatment discontinuation isn’t mandatory.

w. Detailed instructions for tissue collection, processing and shipment are provided in the Procedures Manual. If the subject signs the Future Biomedical Research (FBR) consent, any leftover tissue that would ordinarily be discarded at the end of the main study will be retained for FBR.

x. Baseline tumor tissue for biomarker analysis from an archival tissue sample of a tumor lesion must be provided and submitted to the independent central radiology review vendor before enrollment. These samples are not required to be obtained within 28 days of enrollment. Exceptions to the archival tissue requirement may be granted after discussion with the Sponsor if a newly obtained biopsy is performed at baseline.

y. Newly obtained tumor biopsies are mandatory for subjects prior to Cycle 1 initiation of MK-3475 and again at Cycle 3. Exemptions to the tumor biopsy require Sponsor approval and appropriate justification.

z. Tumor biopsy for clinically stable subjects at treatment discontinuation is highly encouraged.

aa. Blood for correlative studies should be collected prior to Cycle 1, at Cycle 3 and again at treatment discontinuation.
6.3 Second Course Phase (Retreatment ONLY) for Cohorts A, B, C and D

<table>
<thead>
<tr>
<th>Trial Period:</th>
<th>Treatment Cycles&lt;sup&gt;a&lt;/sup&gt;</th>
<th>End of Treatment</th>
<th>Post-Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Treatment Cycle/Title:</td>
<td>Second Course Phase</td>
<td>Discon</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Scheduling Window (Days)&lt;sup&gt;d&lt;/sup&gt;</td>
<td>±3</td>
<td>±3</td>
<td>±3</td>
</tr>
</tbody>
</table>

### Administrative Procedures

| Eligibility Criteria<sup>e</sup> | X |
| Concomitant Medication Review<sup>f</sup> | X X X X X X X X X X |
| Trial Treatment Administration<sup>g</sup> | X X X X X X X X X X |
| Post-study anticancer therapy status | X X |
| Survival Status<sup>h</sup> | X X |

### Clinical Procedures/Assessments

| Review Adverse Events<sup>i</sup> | X X X X X X X X X X X |
| Full Physical Examination | X |
| Directed Physical Examination | X X X X X X X X |
| Vital Signs and Weight<sup>k</sup> | X X X X X X X X X X |
| ECOG Performance Status | X X X X X X X X X X |
| Laboratory Procedures/Assessments: analysis performed by local laboratory |
| Pregnancy Test – Urine or Serum β-HCG<sup>j</sup> | X X |
| PT/INR and aPTT<sup>m</sup> | X X |
| CBC with Differential<sup>n</sup> | X X X X X X X X X X X |
| Comprehensive Chemistry Panel<sup>o</sup> | X X X X X X X X X X X |
| T3, FT4 and TSH<sup>p</sup> | X X X X X X X X X X |
| Laboratory Procedures/Assessments: analysis performed by central laboratory |
| Anti-MK-3475 Antibodies | X X X X X X X X X X X |
| Pharmacokinetics | X X X X X X X X |

### Efficacy Measurements

| Tumor Imaging<sup>q</sup> | X X X X X X X X |

---

<sup>a</sup> Treatment cycles to be repeated beyond 8 cycles.

<sup>b</sup> Follow-Up Visits:
- Every 8 weeks post discon
- Approx. every 12 weeks

<sup>c</sup> Survival Follow-Up:
- 30 days post discon
- Every 8 weeks post discon
- Approx. every 12 weeks

<sup>d</sup> Scheduling Window (Days): ±3 ±3 ±3 ±3 ±3 ±3 ±3 ±3

<sup>e</sup> Eligibility Criteria:
- X

<sup>f</sup> Concomitant Medication Review:
- X X X X X X X X X X

<sup>g</sup> Trial Treatment Administration:
- X X X X X X X X X X

<sup>h</sup> Post-study anticancer therapy status:
- X X

<sup>i</sup> Survival Status:
- X X

<sup>j</sup> Vital Signs and Weight:
- X X X X X X X X X X

<sup>k</sup> ECOG Performance Status:
- X X X X X X X X X X

<sup>l</sup> Pregnancy Test – Urine or Serum β-HCG:
- X

<sup>m</sup> PT/INR and aPTT:
- X

<sup>n</sup> CBC with Differential:
- X X X X X X X X X X

<sup>o</sup> Comprehensive Chemistry Panel:
- X X X X X X X X X X

<sup>p</sup> T3, FT4 and TSH:
- X X X X X X X X X X

<sup>q</sup> Tumor Imaging:
- X X X X X X X X

<sup>r</sup> Tumor Imaging:
- X X X X X X X X
a. In general, assessments/procedures are to be performed on Day 1 and prior to the first dose of treatment for each cycle unless otherwise specified. Treatment cycles are 2 weeks Imaging should always be performed every 8 weeks (56 days ± 7 days) regardless of any treatment delays.

b. In subjects who discontinue study therapy without documented disease progression, every effort should be made to continue monitoring their disease status by radiologic imaging until (1) the start of new anti-cancer treatment, (2) documented disease progression, (3) death, or (4) the end of the study, whichever occurs first. If imaging assessment beyond the protocol-required imaging time frame is necessary, the reason for the extended imaging schedule should be documented in the subject’s chart and recorded in the ORST eCRF for each imaging assessment completed within the extended time frame. The maximum imaging time frame allowed is no greater than twice the imaging time frame noted (ie, for every 8 weeks’ imaging, the maximum imaging schedule allowance is every 16 weeks). Every effort should be made to target the protocol-required imaging schedule as closely as possible; however, in the interest of the subject’s continued participation in the follow-up phase of the study, the extended imaging time frame may be considered at the discretion of the investigator.

c. After the start of new anti-cancer treatment or documented disease progression, the subject should be contacted by telephone approximately every 12 weeks to assess for survival status. Updated survival status may be requested by the Sponsor at any time during the course of the study. Upon Sponsor notification, all subjects who do not/will not have a scheduled study visit or study contact during the Sponsor-defined time period will be contacted for their survival status (excluding participants that have a death event previously recorded).

d. In general, the window for each visit is ± 3 days unless otherwise noted.

e. Subjects who either a) attain a CR and discontinue treatment or b) discontinue treatment after 24 months on MK-3475 for reasons other than disease progression or intolerability may restart trial treatment if they meet the criteria specified in Section 7.1.5.2.1.

f. Concomitant medications – Enter new medications started during the trial through the Safety Follow-up visit. Record all medications taken for SAEs as defined in Section 7.2.

g. Subjects who restart treatment should resume at the same dose and cycle interval which they were receiving prior to discontinuation.

h. AEs and laboratory safety measurements will be graded per NCI CTCAE version 4.0. All AEs, whether gradable by CTCAE or not, will also be evaluated for seriousness.

i. Record all AEs occurring within 30 days after the last dose of trial treatment. Report all SAEs (related and unrelated to trial treatment) and ECIs occurring up until 90 days after the last dose of trial treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier. Afterwards, report only SAEs and ECIs that are related to trial treatment.

j. To be repeated every 4 cycles after cycle 5.

k. Vital signs to include temperature, pulse, respiratory rate, weight and blood pressure.

l. For women of reproductive potential, a urine pregnancy test will be performed within 72 hours prior to first dose of retreatment. If urine pregnancy results cannot be confirmed as negative, a serum pregnancy test performed by the local study site laboratory will be required. Pregnancy tests (serum and/or urine tests) should be repeated if required by local guidelines.

m. Coagulation factors (PT/INR and aPTT) should be monitored closely throughout the trial for any subject receiving anticoagulant therapy.

n. Laboratory tests for determining eligibility for retreatment are to be performed within 10 days prior to the first retreatment dose of MK-3475. See Section 7.1.3 for details regarding laboratory tests.

o. After the first dose, lab samples can be collected up to 72 hours prior to the scheduled time point. See Section 7.1.3 for details regarding laboratory tests.

p. A scan must be performed within 28 days prior to restarting treatment with MK-3475. Imaging should continue to be performed every 8 weeks (56 ± 7 days) from the first dose of trial treatment or more frequently if clinically indicated. Imaging timing should follow calendar days and should not be adjusted for delays in cycle starts or extension of MK-3475 cycle frequencies. The same imaging technique should be used in a subject throughout the trial. Local reading (investigator assessment with site radiology reading) will be used to determine eligibility and for subject management. The Sponsor will collect radiological assessments for retrospective analysis by an independent central radiology review vendor. The processes for image collection and transmission to the independent central radiology review vendor are in the Site Imaging Manual.

q. Per the modified RECIST 1.1 used in this protocol, if imaging shows progressive disease, the imaging assessment should be performed at a minimum of 4 weeks later in order to confirm progressive disease as described in Section 4.2.3.1. Please refer to the Procedure Manual for additional details on modifications to RECIST.
r. In subjects who discontinue study therapy without confirmed disease progression, a radiological evaluation should be performed at the time of treatment discontinuation (i.e. date of discontinuation ± 4 week window). If a previous scan was obtained within 4 weeks prior to the date of discontinuation, then a scan at treatment discontinuation isn’t mandatory.
s. Unresolved labs that are drug related AEs should be followed until resolution. Labs do not need to be repeated after the end of treatment if labs are within normal range.
t. Pre-dose trough PK samples will be collected at Cycles 1 and 2. Pre-dose trough samples will be collected every 4 cycles starting with Cycle 5 and through Cycle 37, 30 days after discontinuation of study drug, and 3 months and 6 months after discontinuation of study drug (or until the subject starts new anti-neoplastic therapy). All trough samples should be drawn within 24 hours before infusion of MK-3475. Anti-MK-3475 antibodies should be drawn with all pre-dose trough PK samples, the 30 day discontinuation draw and 3 months and 6 months after discontinuation of study drug (or until the subject starts new anti-neoplastic therapy). Procedures for sample collection are described in the Procedures Manual.
u. Following Cycle 8, the directed physical exam is only required at Cycle 11, 15, 19, and every 4 cycles thereafter.
v. Following Cycle 8, the ECOG performance status should be determined only in conjunction with a protocol-specified full or directed physical exam (Cycle 9, 11, 13, 15, 17, 19 and every 2 cycles thereafter.
### 6.4 Second Course Phase (Retreatment ONLY) for Cohort B2

<table>
<thead>
<tr>
<th>Trial Period:</th>
<th>Treatment Cycles&lt;sup&gt;a&lt;/sup&gt;</th>
<th>End of Treatment</th>
<th>Post-Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>To be repeated beyond 6 cycles</td>
<td>Discon</td>
<td>Post-Treatment Safety Follow-up</td>
</tr>
<tr>
<td></td>
<td>Second Course Phase</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Treatment Cycle/Title:</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Scheduling Window (Days)&lt;sup&gt;d&lt;/sup&gt;:</td>
<td>±3</td>
<td>±3</td>
<td>±3</td>
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</table>

#### Administrative Procedures

<p>| | |</p>
<table>
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<tbody>
<tr>
<td>Eligibility Criteria&lt;sup&gt;e&lt;/sup&gt;</td>
<td>X</td>
</tr>
<tr>
<td>Concomitant Medication Review&lt;sup&gt;f&lt;/sup&gt;</td>
<td>X X X X X X X X X X X X</td>
</tr>
<tr>
<td>Trial Treatment Administration&lt;sup&gt;g&lt;/sup&gt;</td>
<td>X X X X X X X X X X X X</td>
</tr>
<tr>
<td>Post-study anticancer therapy status</td>
<td>X</td>
</tr>
<tr>
<td>Survival Status&lt;sup&gt;e&lt;/sup&gt;</td>
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</tr>
</tbody>
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#### Clinical Procedures/Assessments

<p>| | |</p>
<table>
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<tbody>
<tr>
<td>Review Adverse Events&lt;sup&gt;h&lt;/sup&gt;</td>
<td>X X X X X X X X X X X X</td>
</tr>
<tr>
<td>Full Physical Examination</td>
<td>X</td>
</tr>
<tr>
<td>Directed Physical Examination</td>
<td>X X X X X X X X X X X X</td>
</tr>
<tr>
<td>Vital Signs and Weight&lt;sup&gt;i&lt;/sup&gt;</td>
<td>X X X X X X X X X X X X</td>
</tr>
<tr>
<td>ECOG Performance Status</td>
<td>X X X X X X X X X X X X</td>
</tr>
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#### Laboratory Procedures/Assessments: analysis performed by local laboratory

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
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<tbody>
<tr>
<td>Pregnancy Test – Urine or Serum β-HCG&lt;sup&gt;j&lt;/sup&gt;</td>
<td>X</td>
</tr>
<tr>
<td>PT/INR and aPTT&lt;sup&gt;l&lt;/sup&gt;</td>
<td>X</td>
</tr>
<tr>
<td>CBC with Differential&lt;sup&gt;m&lt;/sup&gt;</td>
<td>X&lt;sup&gt;n&lt;/sup&gt; X X X X X X X X X X X X X X X</td>
</tr>
<tr>
<td>Comprehensive Chemistry Panel&lt;sup&gt;n&lt;/sup&gt;</td>
<td>X&lt;sup&gt;n&lt;/sup&gt; X X X X X X X X X X X X X X X</td>
</tr>
<tr>
<td>T3, FT4 and TSH&lt;sup&gt;n&lt;/sup&gt;</td>
<td>X&lt;sup&gt;n&lt;/sup&gt; X X X X X X X X X X X X X X X</td>
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#### Efficacy Measurements

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<table>
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<tbody>
<tr>
<td>Tumor Imaging&lt;sup&gt;n&lt;/sup&gt;</td>
<td>X</td>
</tr>
</tbody>
</table>

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<sup>a</sup> Treatments are to be administered at a single site and will be repeated beyond 6 cycles.

<sup>b</sup> Post-Treatment Safety Follow-up visits to be performed at time of Discon.

<sup>c</sup> Post-Treatment Safety Follow-up visits to be performed 30 days post Discon.

<sup>d</sup> Scheduling windows are ±3 days post Discon.

<sup>e</sup> Survival status will be followed every 8 weeks post Discon.

<sup>f</sup> Review of concomitant medications will be performed prior to each cycle.

<sup>g</sup> Review of trial treatment administration will be performed prior to each cycle.

<sup>h</sup> Review of adverse events will be performed prior to each cycle.

<sup>i</sup> Vital signs and weight will be monitored prior to each cycle.

<sup>j</sup> Pregnancy test will be performed prior to each cycle.

<sup>k</sup> Monitoring of ECOG performance status will be performed prior to each cycle.

<sup>l</sup> PT/INR and aPTT will be monitored prior to each cycle.

<sup>m</sup> CBC with differential will be monitored prior to each cycle.

<sup>n</sup> Comprehensive chemistry panel will be monitored prior to each cycle.

<sup>o</sup> T3, FT4 and TSH will be monitored prior to each cycle.

<sup>p</sup> Tumor imaging will be performed prior to each cycle.
a. In general, assessments/procedures are to be performed on Day 1 and prior to the first dose of treatment for each cycle unless otherwise specified. Treatment cycles are 3 weeks long. Imaging should always be performed every 8 weeks (56 days ± 7 days) regardless of any treatment delays.

b. In subjects who discontinue study therapy without documented disease progression, every effort should be made to continue monitoring their disease status by radiologic imaging until (1) the start of new anti-cancer treatment, (2) documented disease progression, (3) death, or (4) the end of the study, whichever occurs first. If imaging assessment beyond the protocol-required imaging time frame is necessary, the reason for the extended imaging schedule should be documented in the subject’s chart and recorded in the ORST eCRF for each imaging assessment completed within the extended time frame. The maximum imaging time frame allowed is no greater than twice the imaging time frame noted (i.e., for every 8 weeks’ imaging, the maximum imaging schedule allowance is every 16 weeks). Every effort should be made to target the protocol-required imaging schedule as closely as possible; however, in the interest of the subject’s continued participation in the follow-up phase of the study, the extended imaging time frame may be considered at the discretion of the investigator.

c. After the start of new anti-cancer treatment or documented disease progression, the subject should be contacted by telephone approximately every 12 weeks to assess for survival status. Updated survival status may be requested by the Sponsor at any time during the course of the study. Upon Sponsor notification, all subjects who do not/will not have a scheduled study visit or study contact during the Sponsor-defined time period will be contacted for their survival status (excluding participants that have a death event previously recorded).

d. In general, the window for each visit is ± 3 days unless otherwise noted.

e. Subjects who either a) attain a CR and discontinue treatment or b) discontinue treatment after 24 months on MK-3475 for reasons other than disease progression or intolerability may restart treatment if they meet the criteria specified in Section 7.1.5.2.1.

f. Concomitant medications – Enter new medications started during the trial through the Safety Follow-up visit. Record all medications taken for SAEs as defined in Section 7.2.

g. Subjects who restart treatment should resume at the same dose and cycle interval which they were receiving prior to discontinuation.

h. Ongoing treatment – Enter ongoing treatment as described in Section 7.1.5.2.2. Record all medications taken for SAEs as defined in Section 7.2.

i. Vital signs to include temperature, pulse, respiratory rate, weight and blood pressure.

j. Record all AEs occurring within 30 days after the last dose of trial treatment. Report all SAEs (related and unrelated to trial treatment) and ECIs occurring up until 90 days after the last dose of trial treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier. Afterwards, report only SAEs and ECIs that are related to trial treatment.

k. To be repeated every 3 cycles after cycle 4.

l. Laboratory tests for determining eligibility for retreatment are to be performed within 10 days prior to the first retreatment dose of MK-3475. See Section 7.1.3 for details regarding laboratory tests.

m. For women of reproductive potential, a urine pregnancy test will be performed within 72 hours prior to first dose of retreatment. If urine pregnancy results cannot be confirmed as negative, a serum pregnancy test performed by the local study site laboratory will be required. Pregnancy tests (serum and/or urine tests) should be repeated if required by local guidelines.

n. Coagulation factors (PT/INR and aPTT) should be monitored closely throughout the trial for any subject receiving anticoagulant therapy.

o. Laboratory safety measurements will be graded per NCI CTCAE version 4.0. All AEs, whether gradable by CTCAE or not, will also be evaluated for seriousness.

p. Imaging should continue to be performed every 8 weeks (56 days ± 7 days) from the first dose of trial treatment or more frequently if clinically indicated. Imaging timing should follow calendar days and should not be adjusted for delays in cycle starts or extension of MK-3475 cycle frequencies. The same imaging technique should be used in a subject throughout the trial. Local reading (investigator assessment with site radiology reading) will be used to determine eligibility and for subject management. The Sponsor will collect radiological assessments for retrospective analysis by an independent central radiology review vendor. The processes for image collection and transmission to the independent central radiology review vendor are in the Site Imaging Manual.

q. Imaging should be performed within 28 days prior to restarting treatment with MK-3475. Imaging should continue to be performed every 8 weeks (56 days ± 7 days) from the first dose of trial treatment or more frequently if clinically indicated. Imaging timing should follow calendar days and should not be adjusted for delays in cycle starts or extension of MK-3475 cycle frequencies. The same imaging technique should be used in a subject throughout the trial. Local reading (investigator assessment with site radiology reading) will be used to determine eligibility and for subject management. The Sponsor will collect radiological assessments for retrospective analysis by an independent central radiology review vendor. The processes for image collection and transmission to the independent central radiology review vendor are in the Site Imaging Manual.

r. Imaging should be performed within 28 days prior to restarting treatment with MK-3475. Imaging should continue to be performed every 8 weeks (56 days ± 7 days) from the first dose of trial treatment or more frequently if clinically indicated. Imaging timing should follow calendar days and should not be adjusted for delays in cycle starts or extension of MK-3475 cycle frequencies. The same imaging technique should be used in a subject throughout the trial. Local reading (investigator assessment with site radiology reading) will be used to determine eligibility and for subject management. The Sponsor will collect radiological assessments for retrospective analysis by an independent central radiology review vendor. The processes for image collection and transmission to the independent central radiology review vendor are in the Site Imaging Manual.

s. Imaging should always be performed every 8 weeks (56 days ± 7 days) regardless of any treatment delays.

t. Imaging should always be performed every 8 weeks (56 days ± 7 days) regardless of any treatment delays.
7.0 TRIAL PROCEDURES

7.1 Trial Procedures

The Trial Flow Chart - Section 6.0 summarizes the trial procedures to be performed at each visit. Individual trial procedures are described in detail below. It may be necessary to perform these procedures at unscheduled time points if deemed clinically necessary by the investigator.

Furthermore, additional evaluations/testing may be deemed necessary by the investigator and or the Sponsor for reasons related to subject safety. In some cases, such evaluation/testing may be potentially sensitive in nature (e.g., HIV, Hepatitis C, etc.), and thus local regulations may require that additional informed consent be obtained from the subject. In these cases, such evaluations/testing will be performed in accordance with those regulations.

7.1.1 Administrative Procedures

7.1.1.1 Informed Consent

The investigator must obtain documented consent from each potential subject prior to participating in a clinical trial or Future Biomedical Research.

7.1.1.1.1 General Informed Consent

Consent must be documented by the subject’s dated signature or by the subject’s legally acceptable representative’s dated signature on a consent form along with the dated signature of the person conducting the consent discussion.

A copy of the signed and dated consent form should be given to the subject before participation in the trial.

The initial informed consent form, any subsequent revised written informed consent form and any written information provided to the subject must receive the IRB/ERC’s approval/favorable opinion in advance of use. The subject or his/her legally acceptable representative should be informed in a timely manner if new information becomes available that may be relevant to the subject’s willingness to continue participation in the trial. The communication of this information will be provided and documented via a revised consent form or addendum to the original consent form that captures the subject’s dated signature or by the subject’s legally acceptable representative’s dated signature.

Specifics about a trial and the trial population will be added to the consent form template at the protocol level.

The informed consent will adhere to IRB/ERC requirements, applicable laws and regulations and Sponsor requirements.
7.1.1.2 Consent and Collection of Specimens for Future Biomedical Research

The investigator or qualified designee will explain the Future Biomedical Research consent to the subject, answer all of his/her questions, and obtain written informed consent before performing any procedure related to the Future Biomedical Research sub-trial. A copy of the informed consent will be given to the subject.

7.1.1.2 Inclusion/Exclusion Criteria

All inclusion and exclusion criteria will be reviewed by the investigator or qualified designee to ensure that the subject qualifies for the trial.

7.1.1.3 Subject Identification Card

All subjects will be given a Subject Identification Card identifying them as participants in a research trial. The card will contain trial site contact information (including direct telephone numbers) to be utilized in the event of an emergency. The investigator or qualified designee will provide the subject with a Subject Identification Card after the subject provides written informed consent.

7.1.1.4 Medical History

A medical history will be obtained by the investigator or qualified designee. Medical history will include all active conditions, and any condition diagnosed within the prior 10 years that are considered to be clinically significant by the Investigator. Details regarding the disease for which the subject has enrolled in this study will be recorded separately and not listed as medical history.

7.1.1.5 Prior and Concomitant Medications Review

7.1.1.5.1 Prior Medications

The investigator or qualified designee will review prior medication use, including any protocol-specified washout requirement, and record prior medication taken by the subject within 28 days before starting the trial. Treatment for the disease for which the subject has enrolled in the study will be recorded separately and not listed as a prior medication.

7.1.1.5.2 Concomitant Medications

The investigator or qualified designee will record medication, if any, taken by the subject during the trial. All medications related to reportable SAEs and ECIs should be recorded as defined in Section 7.2.
7.1.1.6 Disease Details and Treatments

7.1.1.6.1 Disease Details

The investigator or qualified designee will obtain prior and current details regarding disease status. HPV status will be collected for those subjects enrolled in Cohort B and B2.

7.1.1.6.2 Prior Treatment Details

The investigator or qualified designee will review all prior cancer treatments including systemic treatments, radiation and surgeries.

7.1.1.6.3 Subsequent Anti-neoplastic Therapy Status

The investigator or qualified designee will review all new anti-neoplastic therapy initiated after the last dose of trial treatment. If a subject initiates a new anti-neoplastic therapy within 30 days after the last dose of trial treatment, the 30 day Safety Follow-up visit must occur before the first dose of the new therapy. Once new anti-neoplastic therapy has been initiated the subject will move into survival follow-up.

7.1.1.7 Assignment of Screening Number

All consented subjects will be given a unique screening number that will be used to identify the subject for all procedures that occur prior to randomization or treatment allocation. Each subject will be assigned only one screening number. Screening numbers must not be re-used for different subjects. Any subject who is screened multiple times will retain the original screening number assigned at the initial screening visit.

7.1.1.8 Assignment of Randomization Number

All eligible subjects will be allocated, by non-random assignment, to trial treatment and will receive a unique number. This unique number is termed a randomization number throughout the protocol for operational purposes. The randomization number identifies the subject for all procedures occurring after treatment allocation. Once a randomization number is assigned to a subject, it can never be re-assigned to another subject.

A single subject cannot be assigned more than 1 randomization number.

7.1.1.9 Trial Compliance (Medication/Diet/Activity/Other)

Interruptions from the protocol specified treatment plan for greater than 12 weeks between MK-3475 doses due to toxicity require consultation between the investigator and the Sponsor and written documentation of the collaborative decision on subject management.

Administration of trial medication will be witnessed by the investigator and/or trial staff. The total volume of trial treatment infused will be compared to the total volume prepared to determine compliance with each dose administered.
The instructions for preparing and administering MK-3475 will be provided in the Pharmacy Manual.

7.1.2 Clinical Procedures/Assessments

7.1.2.1 Adverse Event (AE) Monitoring

The investigator or qualified designee will assess each subject to evaluate for potential new or worsening AEs as specified in the Trial Flow Chart and more frequently if clinically indicated. Adverse experiences will be graded and recorded throughout the study and during the follow-up period according to NCI CTCAE Version 4.0 (see Section 12.5). Toxicities will be characterized in terms regarding seriousness, causality, toxicity grading, and action taken with regard to trial treatment.

All AEs of unknown etiology associated with MK-3475 exposure should be evaluated to determine if it is possibly an event of clinical interest (ECI) of a potentially immunologic etiology (irAE). See Section 5.6.1.1 in the administrative binder regarding the identification, evaluation and management of AEs of a potential immunological etiology.

Please refer to section 7.2 for detailed information regarding the assessment and recording of AEs.

7.1.2.2 Physical Exam

7.1.2.2.1 Full Physical Exam

The investigator or qualified designee will perform a complete physical exam during the screening period. Clinically significant abnormal findings should be recorded as medical history. A full physical exam should be performed as specified in the Trial Flow Chart (Section 6.0) for the appropriate cohort. After the first dose of trial treatment new clinically significant abnormal findings should be recorded as AEs.

7.1.2.2.2 Directed Physical Exam

For cycles that do not require a full physical exam per the Trial Flow Chart, the investigator or qualified designee will perform a directed physical exam as clinically indicated prior to trial treatment administration. Directed physical exams should be performed as specified in the Trial Flow Chart (Section 6.0) for the appropriate cohort. For Cohorts A, B, C and D after Cycle 8 directed physical exams should occur at Cycle 11 and every 4 cycles thereafter. For Cohort B2 after Cycle 6 a physical exam (either full or directed) is required to be performed at least every 6 weeks but should be performed more frequently if clinically indicated. New clinically significant abnormal findings should be recorded as AEs.
7.1.2.3 Vital Signs

The investigator or qualified designee will take vital signs at screening, prior to the administration of each dose of trial treatment and at treatment discontinuation as specified in the Trial Flow Chart (Section 6.0). Vital signs should include temperature, pulse, respiratory rate, weight and blood pressure. Height will be measured at screening only.

7.1.2.4 Eastern Cooperative Oncology Group (ECOG) Performance Scale

The investigator or qualified designee will assess ECOG status (see Section 12.4) at screening, prior to the administration of each dose of trial treatment and discontinuation of trial treatment as specified in the Trial Flow Chart. After Cycle 8 (Cohorts A, B, C or D) or Cycle 6 (Cohort B2) assessment of ECOG status will be performed in conjunction with the directed or full physical exam.

7.1.2.5 Tumor Imaging and Assessment of Disease

Processes for image collection and transmission to the independent central radiology review vendor can be found in the Site Imaging Manual.

7.1.2.5.1 Assessment of Disease

RECIST 1.1 will be applied by the site as the primary measure for assessment of tumor response and as a basis for all protocol guidelines related to disease status (e.g., discontinuation of study therapy).

RECIST 1.1 will be adapted as follows to account for the unique tumor response seen in this class of therapeutics.

If imaging shows PD, tumor assessment should be repeated ≥ 4 weeks later in order to confirm PD with the option of continuing treatment for clinically stable subjects as discussed below in Table 6. Clinically stable is defined by the following criteria:

- Absence of signs and symptoms indicating disease progression
- No decline in ECOG performance status
- Absence of rapid progression of disease
- Absence of progressive tumor at critical anatomical sites (e.g., cord compression) requiring urgent alternative medical intervention
### Table 6  Imaging and Treatment after 1st radiologic evidence of PD

<table>
<thead>
<tr>
<th>1st radiologic evidence of PD</th>
<th>Clinically Stable</th>
<th>Clinically Unstable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imaging</td>
<td>Treatment</td>
<td>Imaging</td>
</tr>
<tr>
<td>Repeat imaging at ≥ 4 weeks to confirm PD</td>
<td>May continue study treatment at the Investigator’s discretion while awaiting confirmatory scan</td>
<td>Repeat imaging at ≥ 4 weeks to confirm PD if possible</td>
</tr>
<tr>
<td>Repeat scan confirms PD</td>
<td>No additional imaging required</td>
<td>Discontinue treatment (exception noted below)</td>
</tr>
<tr>
<td>Repeat scan shows SD, PR or CR</td>
<td>Continue regularly scheduled imaging assessments every 8 weeks</td>
<td>Continue study treatment at the Investigator’s discretion</td>
</tr>
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</table>

In determining whether or not the tumor burden has increased or decreased, Investigators should consider all target lesions as well as non-target lesions (please refer to the Site Imaging Manual). Subjects that are deemed clinically unstable are not required to have repeat imaging for confirmation. If radiologic progression is confirmed, then the subject will be discontinued from trial treatment. If radiologic progression is not confirmed, then the subject should resume/continue trial treatment and have their next scan according to the every 8 week (56 ± 7 days) schedule.

**NOTE:** If a subject with confirmed radiographic progression (i.e. 2 scans at least 28 days apart demonstrating progressive disease) is clinically stable or clinically improved, and there is no further increase in the tumor dimensions at the confirmatory scan, an exception may be considered to continue treatment upon consultation with the Sponsor. Clinically stable subjects should also have at the confirmatory scan no further increase in the target lesions, no unequivocal increase in non-target lesions, and no additional new lesions develop (non-worsening PD) to continue study treatment.

Imaging during the follow-up period is to be repeated every 8 weeks (56 ± 7 days) for subjects who discontinue trial treatment for reasons other than disease progression until the subject experiences confirmed disease progression or starts a new anti-neoplastic therapy.
If imaging assessment beyond the protocol-required imaging time frame is necessary for the subject’s continued participation in the follow-up phase of the study, an extended imaging schedule may be considered at the discretion of the investigator. Please refer to the applicable Trial Flow Chart in Section 6.0.

Local reading (investigator assessment with site radiology reading) based on RECIST 1.1 will be used to determine subject eligibility and for subject management. The Sponsor will also receive radiologic images for a retrospective analysis of subject eligibility and treatment response to be performed by an independent central radiology review vendor, including RECIST 1.1 and evaluation of volumetric tumor response.

7.1.2.5.2 Initial Tumor Imaging

Initial tumor imaging must be performed within 28 days prior to the first dose of trial treatment. The site study team must review pre-trial images to confirm the subject has measurable disease per RECIST 1.1. The baseline imaging scan should be submitted to the independent central radiology review vendor for a retrospective analysis of this eligibility criterion.

Scans performed as part of routine clinical management are acceptable for use as the screening scan if they are of diagnostic quality and performed within 28 days prior to the first dose of trial treatment. The same imaging technique should be used in a subject throughout the study.

7.1.2.5.3 Tumor Imaging During Trial

Tumor imaging may be performed by CT or magnetic resonance imaging (MRI), but the same imaging technique should be used in a subject throughout the trial. Imaging should be performed every 8 weeks (56 days ± 7 days) from the first dose of trial treatment or more frequently if clinically indicated. Imaging should not be delayed for delays in cycle starts or extension of MK-3475 cycle intervals.

Per RECIST 1.1, response should be confirmed by a repeat radiographic assessment not less than 4 weeks from the date the response was first documented. The scan for confirmation of response may be performed at the earliest 4 weeks after the first indication of response, or at the next scheduled scan (8 weeks later), whichever is clinically indicated.

Imaging should continue to be performed until documented disease progression, the start of new anti-cancer treatment, withdrawal of consent, death, or the end of the study, whichever occurs first. Disease progression should be confirmed at least 4 weeks after the first scan indicating progressive disease in clinically stable subjects. Subjects who have unconfirmed disease progression may continue on treatment until progression is confirmed provided they have met the conditions detailed in Section 7.1.2.5.1.
7.1.2.6 Tumor Tissue Collection and Correlative Studies Blood Sampling

Enrollment in this study is limited to those subjects with tumors who have submitted archival FFPE tumor sample or newly obtained core or excisional biopsy (FNA not adequate) to a central lab for PD-L1 characterization. These samples are not required to be obtained within 28 days of enrollment, however, a biopsy for screening purposes cannot be performed until the main consent is signed.

Enrollment in Cohorts A, B, C and D of this study is limited to those subjects with tumors characterized as PD-L1 positive by IHC at a central laboratory.

Subjects in Cohort B2 should submit both archival FFPE tumor samples and newly obtained biopsy samples (as described below) to the central laboratory. PD-L1 expression by IHC will be evaluated retrospectively. The tumor biopsy specimen must be sufficient for assessment of PD-L1 expression by the central lab or an additional sample will be required. Exceptions to this requirement must be discussed with the Sponsor prior to enrollment in the study.

Biopsy sites should be selected so that subsequent biopsies can be performed at the same location. Exceptions from the mandatory tumor biopsy requirement that allow subjects to continue receiving trial treatment must occur in consultation with the Sponsor.

Blood for correlative biomarker studies should be collected at baseline, on treatment and upon treatment discontinuation. The treatment sample should be collected at Cycle 5 (subjects enrolled in Cohorts A, B, C and D) or Cycle 3 (subjects enrolled in Cohort B2) as appropriate.

Serial tumor biopsy requirements for each cohort are detailed below:

Cohort A – TNBC

Every effort should be made to obtain an optional biopsy prior to Cycle 1 initiation of MK-3475, at Cycle 5, and upon disease progression. If clinical activity of MK-3475 is observed (2 responders in Cohort A), the pre-dose Cycle 1 biopsy and the biopsy at Cycle 5 will become mandatory for all remaining subjects enrolled into the cohort. A tumor biopsy at treatment discontinuation is highly encouraged for all subjects. Tumor biopsies that are inaccessible or contraindicated due to subject safety concerns are exempt from this requirement.

Cohort B – H/N Cancer

Tumor biopsies are mandatory for subjects prior to Cycle 1 initiation of MK-3475 and again at Cycle 5. A tumor biopsy at treatment discontinuation is highly encouraged. Tumor biopsies that are inaccessible or contraindicated due to subject safety concerns are exempt from this requirement.
Cohort B2 – H/N Cancer

Newly obtained tumor biopsies are mandatory for subjects prior to Cycle 1 initiation of MK-3475 and again at Cycle 3. Exemptions to the mandatory tumor biopsy require Sponsor approval and appropriate justification. A tumor biopsy at the time of treatment discontinuation is highly encouraged.

Cohort C – Urothelial Tract Cancer

Every effort should be made to obtain an optional biopsy prior to Cycle 1 initiation of MK-3475, at Cycle 5, and upon disease progression. If clinical activity of MK-3475 is observed (2 responders in Cohort C), the pre-dose Cycle 1 biopsy and the biopsy at Cycle 5 will become mandatory for all remaining subjects enrolled into the cohort. A tumor biopsy at treatment discontinuation is highly encouraged for all subjects. Tumor biopsies that are inaccessible or contraindicated due to subject safety concerns are exempt from this requirement.

Cohort D – Gastric Cancer

Tumor biopsies are mandatory for subjects prior to Cycle 1 initiation of MK-3475 and again at Cycle 5. A tumor biopsy at treatment discontinuation is highly encouraged. Tumor biopsies that are inaccessible or contraindicated due to subject safety concerns are exempt from this requirement.

Detailed instructions for tissue collection, process and shipment are provided in the Procedures Manual.

7.1.3 Laboratory Procedures/Assessments

Details regarding specific laboratory procedures/assessments to be performed in this trial are provided below. The total amount of blood/tissue to be drawn/collected over the course of the trial (from pre-trial to post-trial visits), including approximate blood/tissue volumes drawn/collected by visit and by sample type per subject can be found in the Procedures Manual.

7.1.3.1 Laboratory Safety Evaluations (Hematology, Chemistry and Urinalysis)

Laboratory tests for hematology, chemistry, urinalysis, and others are specified in Table 7.
### Table 7  
#### Laboratory Tests

<table>
<thead>
<tr>
<th>Hematology‡</th>
<th>Chemistry</th>
<th>Urinalysis</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hematocrit</td>
<td>Albumin</td>
<td>Blood</td>
<td>Serum β-human chorionic gonadotropin (β-hCG)†</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>Alkaline phosphatase</td>
<td>Glucose</td>
<td>PT (INR)</td>
</tr>
<tr>
<td>Platelet count</td>
<td>Alanine aminotransferase (ALT)</td>
<td>Protein</td>
<td>aPTT</td>
</tr>
<tr>
<td>WBC (total and differential)</td>
<td>Aspartate aminotransferase (AST)</td>
<td>Specific gravity</td>
<td>Total triiodothyronine (T3)¶</td>
</tr>
<tr>
<td>Red Blood Cell Count</td>
<td>Lactate dehydrogenase (LDH)</td>
<td>Microscopic exam, if abnormal results are notedǁ</td>
<td>Free thyroxine (T4)</td>
</tr>
<tr>
<td>Absolute Neutrophil Count</td>
<td>Carbon Dioxide§ (CO₂ or bicarbonate)</td>
<td>Urine pregnancy test*</td>
<td>Thyroid stimulating hormone (TSH)</td>
</tr>
<tr>
<td>Absolute Lymphocyte Count</td>
<td>Creatinine††</td>
<td>Anti-MK-3475 Antibodies</td>
<td></td>
</tr>
<tr>
<td>Uric Acid</td>
<td></td>
<td>PK</td>
<td></td>
</tr>
<tr>
<td>Calcium</td>
<td></td>
<td>Blood for FBR</td>
<td></td>
</tr>
<tr>
<td>Chloride</td>
<td></td>
<td>Blood for correlative studies</td>
<td></td>
</tr>
<tr>
<td>Glucose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phosphorus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Potassium</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnesium</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Bilirubin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct Bilirubin, if total bilirubin is elevated above the upper limit of normal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total protein</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood Urea Nitrogen</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

‡ Perform on women of childbearing potential only. If urine pregnancy results cannot be confirmed as negative, a serum pregnancy test will be required.

‡ Either differential or absolute value is acceptable

§ If considered standard of care in your region

ǁ Institutional standards are acceptable

¶ Free T3 may be performed in place of Total T3 per local standards

†† GFR (measured or calculated) or CrCl can be used in place of creatinine

Laboratory tests for screening or entry into the Second Course Phase should be performed within 10 days prior to the first dose of treatment. After Cycle 1, pre-dose laboratory procedures can be conducted up to 72 hours prior to dosing. Results must be reviewed by the investigator or qualified designee and found to be acceptable prior to each dose of trial treatment.
7.1.3.2 Pharmacokinetic/Pharmacodynamic Evaluations

7.1.3.2.1 Blood Collection for Serum MK-3475

Sample collection, storage and shipment instructions for serum samples will be provided in the Procedures Manual.

The time points for PK blood sampling are described in Section 6 – Trial Flow Chart.

Please note that PK blood sampling will occur in Cohorts A, B, C, and D only. Subjects in Cohort B2 (H/N expansion) will not undergo PK blood sampling.

7.1.3.2.2 Blood Collection for Anti-MK-3475 Antibodies

Sample collection, storage and shipment instructions for blood samples will be provided in the Procedures Manual.

Please note that blood collection for anti-MK-3475 antibodies will occur in Cohorts A, B, C, and D only. Subjects in Cohort B2 (H/N expansion) will not undergo blood collection for anti-MK-3475 antibodies.

7.1.3.3 Future Biomedical Research

The following specimens are to be obtained as part of Future Biomedical Research:

- Blood for genomics use
- Leftover archival tumor tissue or leftover newly obtained biopsy samples taken throughout the study

7.1.4 Other Procedures

7.1.4.1 Withdrawal/Discontinuation

When a subject discontinues/withdraws prior to trial completion, all applicable activities scheduled for the final trial visit should be performed at the time of discontinuation. Any adverse events which are present at the time of discontinuation/withdrawal should be followed in accordance with the safety requirements outlined in Section 7.2 - Assessing and Recording Adverse Events. Subjects who a) attain a CR or b) complete 24 months of treatment with MK-3475 may discontinue treatment with the option of restarting treatment if they meet the criteria specified in Section 7.1.5.2.1. After discontinuing treatment following assessment of CR, these subjects should return to the site for a Safety Follow-up Visit (described in Section 7.1.5.3.1) and then proceed to the Follow-Up Period of the study (described in Section 7.1.5.4).
7.1.4.1.1 Withdrawal From Future Biomedical Research

Subjects may withdraw their consent for Future Biomedical Research and have their specimens and all derivatives destroyed. Subjects may withdraw consent at any time by writing to the principal investigator for the main trial. If medical records for the main trial are still available, the investigator will contact the Sponsor using the designated mailbox (clinical.specimen.management@merck.com), and a form will be provided by the Sponsor to obtain appropriate information to complete specimen withdrawal. Subsequently, the subject's specimens will be removed from the biorepository and be destroyed. A letter will be sent from the Sponsor to the investigator confirming the destruction. It is the responsibility of the investigator to inform the subject of completion of destruction. Any analyses in progress at the time of request for destruction or already performed prior to the request being received by the Sponsor will continue to be used as part of the overall research trial data and results. No new analyses would be generated after the request is received.

In the event that the medical records for the main trial are no longer available (e.g., if the investigator is no longer required by regulatory authorities to retain the main trial records) or the specimens have been completely anonymized, there will no longer be a link between the subject’s personal information and their specimens. In this situation, the request for specimen destruction cannot be processed.

7.1.4.2 Blinding/Unblinding

This is an open label trial; there is no blinding for this trial.

7.1.5 Visit Requirements

Visit requirements are outlined in Section 6.0 - Trial Flow Chart. Specific procedure-related details are provided above in Section 7.1 - Trial Procedures.

7.1.5.1 Screening

7.1.5.1.1 Pre-screening Period

The Pre-screening period may be utilized by subjects in Cohorts A, B, C and D to determine biomarker eligibility based on PD-L1 status using an archival tumor biopsy sample. After providing a pre-screening consent, subjects will be assigned a screening number. Characterization of PD-L1 status will be performed at a pre-screening visit for subjects with an available archival tumor biopsy sample.

Subjects that do not have an archival tumor biopsy sample available must provide written consent for the main study before the tumor biopsy or any other protocol-specified procedures can occur. These subjects will not enter the pre-screening period as eligibility based on PD-L1 expression will be determined in the main study screening period.

As PD-L1 tumor expression is not an inclusion criterion for subjects in Cohort B2, the Pre-screening period is not relevant for this cohort and all subjects in this cohort will begin the study in the Screening Period.
7.1.5.1.2 Screening Period

Approximately 28 days prior to enrollment, potential subjects will be evaluated to determine that they fulfill the entry requirements as set forth in Section 5.1. Visit requirements are outlined in Section 6.0 – Trial Flow Chart.

Written consent for the main study must be obtained prior to performing any protocol specific procedure. Results of a test performed prior to the subject signing consent as part of routine clinical management are acceptable in lieu of a screening test if performed within the specified time frame. Screening procedures are to be completed within 28 days prior to the first dose of trial treatment except for the following:

- Laboratory tests are to be performed within 10 days prior to the first dose of trial treatment
- For women of reproductive potential, a urine pregnancy test will be performed within 72 hours prior to the first dose of trial treatment. If urine pregnancy results cannot be confirmed as negative, a serum pregnancy test will be required (performed by the local study site laboratory)
- Biopsy for PD-L1 characterization is not required to be obtained within 28 days prior to the first dose of trial treatment

Subjects may be rescreened after initially failing to meet the inclusion/exclusion criteria. Results from assessments performed during the initial screening period are acceptable in lieu of a repeat screening test if performed within the specified time frame and the inclusion/exclusion criteria is met.

7.1.5.2 Treatment Period

7.1.5.2.1 Second Course Phase (Retreatment Period)

Subjects who stop MK-3475 with SD or better may be eligible for up to one year of additional MK-3475 therapy if they progress after stopping MK-3745. This retreatment is termed the Second Course Phase of this study and is only available if the study remains open and the subject meets the following conditions:

- Either
  - Stopped initial treatment with MK-3475 after attaining an investigator-determined confirmed CR according to RECIST 1.1
    - Was treated for at least 24 weeks with MK-3475 before discontinuing therapy
    - Received at least two treatments with MK-3475 beyond the date when the initial CR was declared
OR
- Subject had SD, PR or CR and stopped MK-3475 treatment after 24 months of study therapy for reasons other than disease progression or intolerability

AND
- Experienced an investigator-determined confirmed radiographic disease progression after stopping their initial treatment with MK-3475
- Did not receive any anti-cancer treatment since the last dose of MK-3475
- Have a performance status of 0 or 1 on the ECOG Performance Scale
- Demonstrate adequate organ function as detailed in Section 5.1.2
- Female subject of childbearing potential should have a negative urine or serum pregnancy test within 72 hours prior to receiving retreatment with study medication.
- Female subjects of childbearing potential should be willing to use 2 methods of birth control or be surgically sterile, or abstain from heterosexual activity for the course of the study through 120 days after the last dose of study medication (Reference Section 5.7.2). Subjects of child bearing potential are those who have not been surgically sterilized or have been free from menses for > 1 year.
- Male subjects should agree to use an adequate method of contraception starting with the first dose of study therapy through 120 days after the last dose of study therapy.
- Does not have a history or current evidence of any condition, therapy, or laboratory abnormality that might interfere with the subject’s participation for the full duration of the trial or is not in the best interest of the subject to participate, in the opinion of the treating investigator.

Subjects who restart treatment will be retreated at the same dose frequency as when they last received MK-3475. Treatment will be administered for up to one additional year.

Visit requirements are outlined in Section 6.0 – Trial Flow Chart.

7.1.5.3 Post-Treatment Visits

7.1.5.3.1 Safety Follow-Up Visit

The mandatory Safety Follow-Up Visit should be conducted approximately 30 days after the last dose of trial treatment or before the initiation of a new anti-neoplastic treatment, whichever comes first. All AEs that occur prior to the Safety Follow-Up Visit should be recorded. Subjects with an AE of Grade > 1 will be followed until the resolution of the AE to Grade 0-1 or until the beginning of a new anti-neoplastic therapy, whichever occurs first.
SAEs that occur within 90 days of the end of treatment or 30 days after the end of treatment if the subject initiates new anticancer therapy should also be followed and recorded.

Subjects who are eligible for retreatment with MK-3475 (as described in Section 7.1.5.2.1) may have up to two safety follow-up visits, one after the Treatment Period and one after the Second Course Phase.

7.1.5.4 Follow-up Visits

Subjects who discontinue trial treatment for a reason other than disease progression will move into the Follow-Up Phase and should be assessed every 8 weeks (56 ± 7 days) by radiologic imaging to monitor disease status. If imaging assessment beyond the protocol-required imaging time frame is necessary for the subject’s continued participation in the follow-up phase of the study, an extended imaging schedule may be considered at the discretion of the investigator. Please refer to the applicable Trial Flow Chart in Section 6.0.

Every effort should be made to collect information regarding disease status until the start of new anti-neoplastic therapy, disease progression, death, end of the study or if the subject begins retreatment with MK-3475 as detailed in Section 7.1.5.2.1. Information regarding post-study anti-neoplastic treatment will be collected if new treatment is initiated.

Subjects who are eligible to receive retreatment with MK-3475 according to the criteria in Section 7.1.5.2.1 will move from the follow-up phase to the Second Course Phase when they experience disease progression. Details are provided in Section 6.2 – Trial Flow Chart for Retreatment.

7.1.5.4.1 Survival Follow-up

Once a subject experiences confirmed disease progression or starts a new anti-neoplastic therapy, the subject moves into the survival follow-up phase and should be contacted by telephone approximately every 12 weeks (+/- 4 weeks) to assess for survival status until death, withdrawal of consent, or the end of the study, whichever occurs first.

7.1.5.5 Survival Status

To ensure current and complete survival data are available at the time of database locks, updated survival data may be requested during the course of the trial by the Sponsor. For example, updated survival status may be requested prior to but not limited to an external Data Monitoring Committee review, interim, and/or final analysis. Upon Sponsor notification, all subjects who do not/will not have a scheduled study visit or study contact during the Sponsor-defined time period will be contacted for their survival status (excluding subjects who have a previously recorded death event in the collection tool).
7.2 Assessing and Recording Adverse Events

An adverse event is defined as any untoward medical occurrence in a patient or clinical investigation subject administered a pharmaceutical product and which does not necessarily have to have a causal relationship with this treatment. An adverse event can therefore be any unfavourable and unintended sign (including an abnormal laboratory finding, for example), symptom, or disease temporally associated with the use of a medicinal product or protocol-specified procedure, whether or not considered related to the medicinal product or protocol-specified procedure. Any worsening (i.e., any clinically significant adverse change in frequency and/or intensity) of a preexisting condition that is temporally associated with the use of the Sponsor’s product, is also an adverse event.

Changes resulting from normal growth and development that do not vary significantly in frequency or severity from expected levels are not to be considered adverse events. Examples of this may include, but are not limited to, teething, typical crying in infants and children and onset of menses or menopause occurring at a physiologically appropriate time.

Sponsor’s product includes any pharmaceutical product, biological product, device, diagnostic agent or protocol-specified procedure, whether investigational (including placebo or active comparator medication) or marketed, manufactured by, licensed by, provided by or distributed by the Sponsor for human use.

Adverse events may occur during clinical trials, or as prescribed in clinical practice, from overdose (whether accidental or intentional), from abuse and from withdrawal.

Progression of the cancer under study is not considered an adverse event.

All adverse events that occur after the consent form is signed but before randomization/treatment allocation must be reported by the investigator if they cause the subject to be excluded from the trial, or are the result of a protocol-specified intervention, including but not limited to washout or discontinuation of usual therapy, diet, placebo treatment or a procedure.

From the time of randomization/treatment allocation through 30 days following cessation of treatment, all adverse events must be reported by the investigator. Such events will be recorded at each examination on the Adverse Event case report forms/worksheets. The reporting timeframe for adverse events meeting any serious criteria is described in section 7.2.3.1. The investigator will make every attempt to follow all subjects with non-serious adverse events for outcome.

Electronic reporting procedures can be found in the EDC data entry guidelines. Paper reporting procedures can be found in the Investigator Trial File Binder (or equivalent).

Adverse events will not be collected for subjects during the pre-screening period (for determination of archival tissue status) as long as that subject has not undergone any protocol-specified procedure or intervention. If the subject requires a blood draw, fresh tumor biopsy etc., the subject is first required to provide consent to the main study and AEs will be captured according to guidelines for standard AE reporting.

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Confidential

11-Dec-2017
7.2.1 Definition of an Overdose for This Protocol and Reporting of Overdose to the Sponsor

For purposes of this trial, an overdose will be defined as any dose exceeding the prescribed dose for MK-3475 by 20% over the prescribed dose. No specific information is available on the treatment of overdose of MK-3475. In the event of overdose, MK-3475 should be discontinued and the subject should be observed closely for signs of toxicity. Appropriate supportive treatment should be provided if clinically indicated.

If an adverse event(s) is associated with ("results from") the overdose of Sponsor's product or vaccine, the adverse event(s) is reported as a serious adverse event, even if no other seriousness criteria are met.

If a dose of Sponsor's product or vaccine meeting the protocol definition of overdose is taken without any associated clinical symptoms or abnormal laboratory results, the overdose is reported as a non-serious Event of Clinical Interest (ECI), using the terminology “accidental or intentional overdose without adverse effect.”

All reports of overdose with and without an adverse event must be reported within 24 hours to the Sponsor either by electronic media or paper. Sponsor Contact information can be found in the Investigator Trial File Binder.

7.2.2 Reporting of Pregnancy and Lactation to the Sponsor

Although pregnancy and lactation are not considered adverse events, it is the responsibility of investigators or their designees to report any pregnancy or lactation in a subject (spontaneously reported to them) that occurs during the trial.

Pregnancies and lactations that occur after the consent form is signed but before randomization/treatment allocation must be reported by the investigator if they cause the subject to be excluded from the trial, or are the result of a protocol-specified intervention, including but not limited to washout or discontinuation of usual therapy, diet, placebo treatment or a procedure.

Pregnancies and lactations that occur from the time of randomization/treatment allocation through 120 days following cessation of Sponsor’s product, or 30 days following cessation of treatment if the subject initiates new anticancer therapy, whichever is earlier, must be reported by the investigator. All reported pregnancies must be followed to the completion/termination of the pregnancy. Pregnancy outcomes of spontaneous abortion, missed abortion, benign hydatidiform mole, blighted ovum, fetal death, intrauterine death, miscarriage and stillbirth must be reported as serious events (Important Medical Events). If the pregnancy continues to term, the outcome (health of infant) must also be reported.

Such events must be reported within 24 hours to the Sponsor either by electronic media or paper. Electronic reporting procedures can be found in the EDC data entry guidelines. Paper reporting procedures can be found in the Investigator Trial File Binder (or equivalent).
7.2.3 Immediate Reporting of Adverse Events to the Sponsor

7.2.3.1 Serious Adverse Events

A serious adverse event is any adverse event occurring at any dose or during any use of Sponsor's product that:

- Results in death;
- Is life threatening;
- Results in persistent or significant disability/incapacity;
- Results in or prolongs an existing inpatient hospitalization;
- Is a congenital anomaly/birth defect;
- Is an other important medical event

Note: In addition to the above criteria, adverse events meeting either of the below criteria, although not serious per ICH definition, are reportable to the Sponsor in the same timeframe as SAEs to meet certain local requirements. Therefore, these events are considered serious by the Sponsor for collection purposes.

- Is a new cancer (that is not a condition of the study);
- Is associated with an overdose.

Refer to Table 8 for additional details regarding each of the above criteria.

For the time period beginning when the consent form is signed until randomization/treatment allocation, any serious adverse event, or follow up to a serious adverse event, including death due to any cause other than progression of the cancer under study (reference Section 7.2.3.3 for additional details), that occurs to any subject must be reported within 24 hours to the Sponsor if it causes the subject to be excluded from the trial, or is the result of a protocol-specified intervention, including but not limited to washout or discontinuation of usual therapy, diet, placebo treatment or a procedure.

For the time period beginning at randomization/treatment allocation through 90 days following cessation of treatment, or 30 days following cessation of treatment if the subject initiates new anticancer therapy, whichever is earlier, any serious adverse event, or follow up to a serious adverse event, including death due to any cause other than progression of the cancer under study (reference Section 7.2.3.3 for additional details), whether or not related to the Sponsor's product, must be reported within 24 hours to the Sponsor either by electronic media or paper. Electronic reporting procedures can be found in the EDC data entry guidelines. Paper reporting procedures can be found in the Investigator Trial File Binder (or equivalent).

Additionally, any serious adverse event, considered by an investigator who is a qualified physician to be related to the Sponsor's product that is brought to the attention of the investigator at any time outside of the time period specified in the previous paragraph also must be reported immediately to the Sponsor.
All subjects with serious adverse events must be followed up for outcome.

7.2.3.2 Events of Clinical Interest

Selected non-serious and serious adverse events are also known as Events of Clinical Interest (ECI) and must be reported to the Sponsor.

For the time period beginning when the consent form is signed until randomization/treatment allocation, any ECI, or follow up to an ECI, that occurs to any subject must be reported within 24 hours to the Sponsor if it causes the subject to be excluded from the trial, or is the result of a protocol-specified intervention, including but not limited to washout or discontinuation of usual therapy, diet, placebo treatment or a procedure.

For the time period beginning at randomization/treatment allocation through 90 days following cessation of treatment, or 30 days following cessation of treatment if the subject initiates new anticancer therapy, whichever is earlier, any ECI, or follow up to an ECI, whether or not related to the Sponsor’s product, must be reported within 24 hours to the Sponsor, either by electronic media or paper. Electronic reporting procedures can be found in the EDC data entry guidelines. Paper reporting procedures can be found in the Investigator Trial File Binder (or equivalent).

Events of clinical interest for this trial include:

1. an overdose of Sponsor's product, as defined in Section 7.2.1 - Definition of an Overdose for This Protocol and Reporting of Overdose to the Sponsor, that is not associated with clinical symptoms or abnormal laboratory results.

2. an elevated AST or ALT lab value that is greater than or equal to 3X the upper limit of normal and an elevated total bilirubin lab value that is greater than or equal to 2X the upper limit of normal and, at the same time, an alkaline phosphatase lab value that is less than 2X the upper limit of normal, as determined by way of protocol-specified laboratory testing or unscheduled laboratory testing.*

*Note: These criteria are based upon available regulatory guidance documents. The purpose of the criteria is to specify a threshold of abnormal hepatic tests that may require an additional evaluation for an underlying etiology. The trial site guidance for assessment and follow up of these criteria can be found in the Investigator Trial File Binder.
3. Additional adverse events:

A separate guidance document has been provided entitled Event of Clinical Interest Guidance Document” (previously entitled “Event of Clinical Interest and Immune-Related Adverse Event Guidance Document”). This document can be found in the administrative binder and provides guidance regarding identification, evaluation and management of ECIs and irAEs.

Subjects should be assessed for possible ECIs prior to each dose. Lab results should be evaluated and subjects should be asked for signs and symptoms suggestive of an immune-related event. Subjects who develop an ECI thought to be immune-related should have additional testing to rule out other etiologic causes. If lab results or symptoms indicate a possible immune-related ECI, then additional testing should be performed to rule out other etiologic causes. If no other cause is found, then it is assumed to be immune-related.

Events of clinical interest that occur to any subject from the date of first dose through 90 days following cessation of treatment, or 30 days after the end of treatment if the subject initiates new anticancer therapy, whichever is earlier, whether or not related to the Sponsor's product, must be reported within 24 hours to the Sponsor either by electronic media or paper. Sponsor Contact information can be found in the administrative binder.

7.2.3.3 Protocol-Specific Exceptions to Serious Event Reporting

Efficacy endpoints as outlined in this section will not be reported to the Sponsor as described in Section 7.2.3 - Immediate Reporting of Adverse Events to the Sponsor.

Specifically, the suspected/actual events covered in this exception include any event that is disease progression of the cancer under study.

The Sponsor will monitor unblinded aggregated efficacy endpoint events and safety data to ensure the safety of the subjects in the trial. Any suspected endpoint which upon review is not progression of the cancer under study will be forwarded to global safety as a SAE within 24 hours of determination that the event is not progression of the cancer under study.

7.2.4 Evaluating Adverse Events

An investigator who is a qualified physician will evaluate all adverse events according to the NCI Common Terminology for Adverse Events (CTCAE), version 4.0. Any adverse event which changes CTCAE grade over the course of a given episode will have each change of grade recorded on the adverse event case report forms/worksheets.

All adverse events regardless of CTCAE grade must also be evaluated for seriousness.
### Evaluating Adverse Events

An investigator who is a qualified physician, will evaluate all adverse events as to:

<table>
<thead>
<tr>
<th>V4.0 CTCAE Grading</th>
<th>Grade 1</th>
<th>Mild; asymptomatic or mild symptoms; clinical or diagnostic observations only; intervention not indicated.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Grade 2</td>
<td>Moderate; minimal, local or noninvasive intervention indicated; limiting age-appropriate instrumental ADL.</td>
</tr>
<tr>
<td></td>
<td>Grade 3</td>
<td>Severe or medically significant but not immediately life-threatening; hospitalization or prolongation of hospitalization indicated;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>disabling; limiting self-care ADL.</td>
</tr>
<tr>
<td></td>
<td>Grade 4</td>
<td>Life threatening consequences; urgent intervention indicated.</td>
</tr>
<tr>
<td></td>
<td>Grade 5</td>
<td>Death related to AE</td>
</tr>
</tbody>
</table>

#### Seriousness

A serious adverse event is any adverse event occurring at any dose or during any use of Sponsor’s product that:

- Results in death; or
- Is life threatening; or
- Places the subject, in the view of the investigator, at immediate risk of death from the event as it occurred (Note: This does not include an adverse event that, had it occurred in a more severe form, might have caused death); or
- Results in a persistent or significant disability/incapacity (substantial disruption of one’s ability to conduct normal life functions); or
- Results in or prolongs an existing inpatient hospitalization (hospitalization is defined as an inpatient admission, regardless of length of stay, even if the hospitalization is a precautionary measure for continued observation. (Note: Hospitalization [including hospitalization for an elective procedure] for a preexisting condition which has not worsened does not constitute a serious adverse event.); or
- Is a congenital anomaly/birth defect (in offspring of subject taking the product regardless of time to diagnosis); or
- Other important medical events that may not result in death, not be life threatening, or not require hospitalization may be considered a serious adverse event when, based upon appropriate medical judgment, the event may jeopardize the subject and may require medical or surgical intervention to prevent one of the outcomes listed previously (designated above by a †).

#### Duration

Record the start and stop dates of the adverse event. If less than 1 day, indicate the appropriate length of time and units.

#### Action Taken

Did the adverse event cause the Sponsor’s product to be discontinued?

#### Relationship to Sponsor’s Product

Did the Sponsor’s product cause the adverse event? The determination of the likelihood that the Sponsor’s product caused the adverse event will be provided by an investigator who is a qualified physician. The investigator’s signed/dated initials on the source document or worksheet that supports the causality noted on the AE form, ensures that a medically qualified assessment of causality was done. This initialed document must be retained for the required regulatory time frame. The criteria below are intended as reference guidelines to assist the investigator in assessing the likelihood of a relationship between the test drug and the adverse event based upon the information.

The following components are to be used to assess the relationship between the Sponsor’s product and the AE; the greater the correlation with the components and their respective elements (in number and/or intensity), the more likely the Sponsor’s product caused the adverse event (AE):

<table>
<thead>
<tr>
<th>Component</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure</td>
<td>Is there evidence that the subject was actually exposed to the Sponsor’s product such as: reliable history, acceptable compliance assessment (pill count, diary, etc.), expected pharmacologic effect, or measurement of drug/metabolite in bodily specimen?</td>
</tr>
<tr>
<td>Time Course</td>
<td>Did the AE follow in a reasonable temporal sequence from administration of the Sponsor’s product? Is the time of onset of the AE compatible with a drug-induced effect (applies to trials with investigational medicinal product)?</td>
</tr>
<tr>
<td>Likely Cause</td>
<td>Is the AE not reasonably explained by another etiology such as underlying disease, other drug(s)/vaccine(s), or other host or environmental factors</td>
</tr>
</tbody>
</table>
Relationship to Sponsor’s Product (continued)

<table>
<thead>
<tr>
<th>Relationship to Sponsor’s Product (continued)</th>
<th>The following components are to be used to assess the relationship between the test drug and the AE: (continued)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dechallenge</td>
<td>Was the Sponsor’s product discontinued or dose/exposure/frequency reduced? If yes, did the AE resolve or improve? If yes, this is a positive dechallenge. If no, this is a negative dechallenge. (Note: This criterion is not applicable if: (1) the AE resulted in death or permanent disability; (2) the AE resolved/improved despite continuation of the Sponsor’s product; or (3) the trial is a single-dose drug trial; or (4) Sponsor’s product(s) is/are only used one time.)</td>
</tr>
<tr>
<td>Rechallenge</td>
<td>Was the subject re-exposed to the Sponsor’s product in this study? If yes, did the AE recur or worsen? If yes, this is a positive rechallenge. If no, this is a negative rechallenge. (Note: This criterion is not applicable if: (1) the initial AE resulted in death or permanent disability, or (2) the trial is a single-dose drug trial); or (3) Sponsor’s product(s) is/are used only one time).</td>
</tr>
</tbody>
</table>

Consistency with Trial Treatment Profile

| Consistency with Trial Treatment Profile | Is the clinical/pathological presentation of the AE consistent with previous knowledge regarding the Sponsor’s product or drug class pharmacology or toxicology? |

The assessment of relationship will be reported on the case report forms/worksheets by an investigator who is a qualified physician according to his/her best clinical judgment, including consideration of the above elements.

Record one of the following

Yes, there is a reasonable possibility of Sponsor's product relationship.

No, there is not a reasonable possibility of Sponsor's product relationship.

Use the following scale of criteria as guidance (not all criteria must be present to be indicative of a Sponsor’s product relationship).

| Yes, there is a reasonable possibility of Sponsor's product relationship | There is evidence of exposure to the Sponsor's product. The temporal sequence of the AE onset relative to the administration of the Sponsor's product is reasonable. The AE is more likely explained by the Sponsor's product than by another cause. |
| No, there is not a reasonable possibility of Sponsor's product relationship | Subject did not receive the Sponsor's product OR temporal sequence of the AE onset relative to administration of the Sponsor's product is not reasonable OR the AE is more likely explained by another cause than the Sponsor’s product. (Also entered for a subject with overdose without an associated AE.) |
7.2.5 Sponsor Responsibility for Reporting Adverse Events

All Adverse Events will be reported to regulatory authorities, IRB/IECs and investigators in accordance with all applicable global laws and regulations, i.e., per ICH Topic E6 (R1) Guidelines for Good Clinical Practice.

8.0 STATISTICAL ANALYSIS PLAN

8.1 Statistical Analysis Plan Summary

This section contains a brief summary of the statistical analyses for this trial. Full detail is in the Statistical Analysis Plan (SAP) (Section 8.2).

The primary purpose of this study is to:

- investigate the safety, tolerability, and anti-tumor activity of MK-3475 10 mg/kg Q2W administered intravenously to subjects with PD-L1 positive triple negative advanced breast cancer (Cohort A), subjects with PD-L1 positive advanced head and neck cancer (Cohort B), subjects with PD-L1 positive advanced urothelial tract cancer (Cohort C), and subjects with PD-L1 positive advanced gastric cancer (Cohort D).

- investigate the safety, tolerability, and anti-tumor activity of MK-3475 200 mg Q3W administered intravenously to subjects with advanced head and neck cancer irrespective of PD-L1 expression (Cohorts B2).

8.1.1 Efficacy Analyses

The full analysis set (FAS) population (defined as all subjects with a baseline scan with measurable disease; by independent central radiology review for each cohort, and who either have a post baseline scan or discontinue the trial due to progressive disease or a drug-related AE) will serve as the primary population for the analyses of efficacy data in this trial. Supportive analyses of efficacy will be conducted in the all subjects as treated (ASaT) population and the FAS population by Investigator review. For each cohort, overall response rate will be used as the primary endpoint for efficacy assessment. A 95% confidence interval along with a one-sided p-value for testing the null hypothesis based on the binomial distribution will be provided for the response rate in each cohort. The respective cohort is considered to have reached the efficacy objective if the corresponding p-value for testing the respective null hypothesis is less than 2.5%. An outline of the efficacy analysis strategy is presented in Table 9 below.
<table>
<thead>
<tr>
<th>Endpoint/Variable(^1) (Description, Time Point)</th>
<th>Statistical Method</th>
<th>Analysis Population</th>
<th>Missing Data Approach</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review (Cohort A, Cohort B HPV negative subjects, Cohort C, and Cohort D evaluated separately)</td>
<td>Exact test of binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review for subjects in Cohort B2</td>
<td>Exact test of binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Secondary:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review, Cohort B HPV positive subjects,</td>
<td>Exact test of binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review, Cohort D AP subjects,</td>
<td>Exact test of binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review, for subjects previously treated with cetuximab and platinum in Cohorts B and B2</td>
<td>Exact test of binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on investigator assessment for cohorts A, B, C and D</td>
<td>Exact methods for binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on investigator assessment for Cohort B2</td>
<td>Exact methods for binomial parameter</td>
<td>FAS</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Progression-free survival</td>
<td>Summary statistics using Kaplan-Meier method</td>
<td>FAS</td>
<td>Censored at last assessment</td>
</tr>
<tr>
<td>Overall survival</td>
<td>Summary statistics using Kaplan-Meier method</td>
<td>FAS</td>
<td>Censored at last assessment</td>
</tr>
<tr>
<td>Response duration</td>
<td>Summary statistics using Kaplan-Meier method</td>
<td>All responders</td>
<td>Non-responders are excluded in analysis</td>
</tr>
</tbody>
</table>

\(^1\) For Cohort D, the analyses for the Asia Pacific (AP) population will be performed as appropriate.
8.1.2 Safety Analyses

The All-Subjects-as-Treated population will be employed for safety analyses.

8.1.3 Power and Sample Size

The calculation of power and sample size for each cohort does not account for interim analysis. Cohort A (triple negative breast cancer subjects): With approximately 26 evaluable PD-L1 positive subjects with triple negative breast cancer, the study has approximately 80% power to detect a 25% difference in ORR under the null hypothesis of ORR=20% with a type I error rate of 2.5% if the true ORR is 45%. Success for this hypothesis requires at least 10/26 responses. The actual number of subjects enrolled may be larger than 26 to ensure that at least 26 subjects are evaluable for analysis.

Cohort B (head and neck cancer subjects): HPV negative head and neck cancer subjects will be evaluated separately from HPV positive head and neck cancer subjects. With a maximum of 22 evaluable PD-L1 positive subjects with HPV negative head and neck cancer, the study has approximately 80% power to detect a 25% difference in ORR under the null hypothesis of ORR=10% with a type I error rate of 2.5% if the true ORR is 35%. Success for this hypothesis requires at least 6/22 responses. The actual number of subjects enrolled may be larger than 22 to ensure that at least 22 subjects are evaluable for analysis.

With a maximum of 12 evaluable PD-L1 positive subjects with HPV positive head and neck cancer, the study has approximately 73% power to detect a 35% difference in ORR under the null hypothesis of ORR=20% with a type I error rate of 5% if the true ORR is 55%. Success for this hypothesis requires at least 6 responses. The actual number of subjects enrolled may be larger than 12 to ensure that at least 12 subjects are evaluable for analysis. However, if 6 responses are observed prior to 12 subjects with HPV positive head and neck cancer enrolling in the trial, enrollment may be stopped in this group as the hypothesis success criterion will have been reached.

Cohort C (urothelial tract cancer subjects): With a maximum of 22 evaluable PD-L1 positive subjects with urothelial tract cancer, the study has approximately 80% power to detect a 25% difference in ORR under the null hypothesis of ORR=10% with a type I error rate of 2.5% if the true ORR is 35%. Success for this hypothesis requires at least 6/22 responses. The actual number of subjects enrolled may be larger than 22 to ensure that at least 22 subjects are evaluable for analysis.

Cohort D (gastric cancer subjects): With a maximum of 32 evaluable PD-L1 positive subjects with gastric cancer, stratified to include 16 Asia Pacific (AP) and 16 non-AP, the study has approximately 90% power to detect a 25% difference in ORR under the null hypothesis of ORR=15% with a type I error rate of 2.5% if the true ORR is 40%. Success for this hypothesis requires at least 10/32 responses. A total of 16 AP subjects will provide approximately 80% power with a type I error rate of 0.1 to detect a 25% difference in ORR under the null hypothesis of ORR=15% if the true ORR is 40%. At least 4/16 responses will be needed to claim the success of this hypothesis for AP population. The actual number of subjects enrolled may be larger than 32 to ensure that at least 32 subjects are evaluable for analysis.
Cohort B2 (head and neck cancer subjects, expansion cohort): With 100 evaluable subjects in Cohort B2, the study provides >99% power to detect a 15% difference in ORR under the null hypothesis of ORR=5% with a type I error rate of 2.5% if the true ORR is 20%. Success for this hypothesis requires at least 11/100 responses. The actual number of subjects enrolled may be larger than 100 to ensure at least 100 subjects are evaluable for analysis.

Cohorts B and B2 previously treated with cetuximab and platinum: With 60 evaluable head and neck cancer subjects previously treated with cetuximab and platinum, the study has 93% power to detect a 15% difference in ORR under the null hypothesis of ORR=5% with a type I error rate of 2.5% if the true ORR is 20%. Success for this hypothesis requires at least 8/60 responses.

### 8.1.4 Interim Analysis

No efficacy interim analyses are planned for Cohort B2 or Cohort D in this trial. However, an interim analysis for Cohort A, B or C may be performed if the rate of enrollment is much slower than anticipated during the course of the trial. An interim analysis for each cohort would only be performed in this study when ≥10 subjects in the respective cohort have had post-baseline scans through Week 16. Results will be reviewed by the study team. Table 10 summarizes the strategy and timing of the potential interim analysis for each cohort.

For Cohort A, if an interim analysis is conducted and 2 or fewer subjects out of the first 10 subjects with post-baseline imaging scans (Week 16) have a confirmed or unconfirmed response, then enrollment may be paused until response data for the subsequent imaging scan (Week 24) are reviewed for subjects already enrolled in the trial. Enrollment may be resumed if the binomial probability of the observed response rate is ≥ 80% under the assumption of a ≥ 20% true response rate (e.g., ≥4/16 confirmed or unconfirmed response).

For Cohorts B (HPV negative subjects only) and C, if an interim analysis is conducted and 1 or fewer subjects out of the first 10 subjects with post-baseline imaging scans (Week 16) have a confirmed or unconfirmed response, then enrollment may be paused until response data for the subsequent imaging scan (Week 24) are reviewed for subjects already enrolled in the trial. Enrollment may be resumed if the binomial probability of the observed response rate is ≥ 80% under the assumption of a ≥ 10% true response rate (e.g., ≥2/16 confirmed or unconfirmed response). No interim analysis will be performed for the subjects with HPV positive head and neck cancer in Cohort B.

Although safety monitoring will occur continuously in the study, an assessment of Grade 4/5 drug-related immunologic AEs will be performed after 10 subjects have been enrolled within each cohort for 1 cycle of therapy. Enrollment will be paused if 4 or more subjects out of the first 10 subjects within a cohort experience a Grade 4/5 drug-related immunologic adverse event. Enrollment may be resumed within a cohort only after a full safety evaluation is performed in consultation between the study investigators and the Sponsor.
Table 10 Summary of Interim Analysis Strategy

<table>
<thead>
<tr>
<th>Interim Analysis Number</th>
<th>Key Endpoints for Interim Analysis</th>
<th>Timing (Sample Size) for Analysis</th>
<th>Purpose of Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interim Analysis Cohort A</td>
<td>ORR</td>
<td>≥10 subjects with post-baseline scans (Week 16)</td>
<td>Pause enrollment until further evidence of efficacy &lt;br&gt; Pause enrollment for safety based on Grade 4+ drug-related immunologic AEs out of first 10 subjects</td>
</tr>
<tr>
<td>Interim Analysis Cohort B (HPV negative subjects only)</td>
<td>ORR</td>
<td>≥10 subjects with post-baseline scans (Week 16)</td>
<td>Pause enrollment until further evidence of efficacy &lt;br&gt; Pause enrollment for safety based on Grade 4+ drug-related immunologic AEs out of first 10 subjects</td>
</tr>
<tr>
<td>Interim Analysis Cohort C</td>
<td>ORR</td>
<td>≥10 subjects with post-baseline scans (Week 16)</td>
<td>Pause enrollment until further evidence of efficacy &lt;br&gt; Pause enrollment for safety based on Grade 4+ drug-related immunologic AEs out of first 10 subjects</td>
</tr>
</tbody>
</table>

8.2 Statistical Analysis Plan

8.2.1 Responsibility for Analyses/In-House Blinding

The statistical analysis of the data obtained from this study will be the responsibility of the Clinical Biostatistics department of the SPONSOR. This trial is being conducted as an open-label study, i.e., subjects, investigators, and SPONSOR personnel will be aware of subject treatment assignments after each subject is enrolled and treatment is assigned. Access to the PD-L1 subject-level biomarker results for subjects in Cohort B2 for the purpose of summary reporting of PD-L1 level until the time that a clinical study report (CSR) is generated will be limited to an unblinded SPONSOR statistician and unblinded SPONSOR statistical programmer who will be responsible for data review to ensure validity of results and summary reporting of clinical response by biomarker status, but who will have no other responsibilities associated with the study.

The Clinical Biostatistics department will generate the allocation schedule(s) for study treatment assignment. Allocation will be implemented in an interactive voice response system (IVRS).
8.2.2 Hypotheses/Estimation
Objectives and hypotheses of the study are stated in Section 3.0.

8.2.3 Analysis Endpoints
Efficacy and safety endpoints that will be evaluated for within- and/or between-treatment differences are listed below, followed by the descriptions of the derivations of selected endpoints.

8.2.3.1 Efficacy/Immunogenicity/Pharmacokinetics Endpoints
Efficacy and safety endpoints that will be evaluated for are listed below, followed by the descriptions of the derivations of selected endpoints.

The primary efficacy endpoint is response rate, defined as the proportion of subjects in the analysis population who have complete response (CR) or partial response (PR) using RECIST 1.1 criteria at any time during the study. Response for the primary analysis for cohorts A, B, C, D and B2 will be determined by a central independent radiology review. Secondary analyses will be based on the responses as assessed by the investigator.

Key secondary efficacy endpoints include:

- RECIST 1.1 response rate among HPV positive subjects.
- Response rate based on site assessments using RECIST 1.1.

Other secondary efficacy endpoints include: (1) duration of response, defined as time from first RECIST 1.1 response to disease progression in subjects who achieve a PR or better; (2) progression-free survival (PFS), defined as the time from allocation to the first documented disease progression according to RECIST 1.1 or death due to any cause, whichever occurs first; and (3) overall survival (OS).

8.2.3.2 Safety Endpoints
A description of safety measures is provided in Section 4.2.3.2.

The primary safety endpoints are AEs graded using CTCAE (Version 4.0) criteria. Safety will be assessed by quantifying the toxicities and grades experienced by subjects who have received MK-3475, including serious adverse events (SAEs) and events of clinical interest (ECIs). Immune-related ECIs, as described in Section 7.2.3.2 will be collected. Other safety endpoints include laboratory safety assessments, ECOG performance status, vital signs and physical examinations.
8.2.4 Analysis Populations

8.2.4.1 Efficacy Analysis Populations

The Full Analysis Set (FAS) population will serve as the primary population for the analysis of efficacy data in this study. The FAS population consists of all randomized subjects in each Cohort who:

- Receive at least one dose of study treatment,
- Have a baseline scan with measurable disease per RECIST 1.1 by independent central radiology review, and
- Have a post baseline scan OR discontinue the trial due to progressive disease/drug related AE

For secondary by Investigator assessment, measurable disease for inclusion in the FAS population will be defined using Investigator assessment of the baseline scan.

A supportive analysis using the All Subjects as Treated (ASaT) population, defined as all randomized subjects, who received at least one dose of study treatment, will be performed for the primary efficacy endpoint(s).

Subjects will be included in the treatment group to which they are randomized for the analysis of efficacy data. Details on the approach to handling missing data are provided in Section 8.2.5 Statistical Methods.

8.2.4.2 Safety Analysis Populations

The All Subjects as Treated (ASaT) population will be used for the analysis of safety data in this study. The ASaT population consists of all allocated subjects who received at least one dose of study treatment.

At least one laboratory or vital sign measurement obtained subsequent to at least one dose of study treatment is required for inclusion in the analysis of each specific parameter. To assess change from baseline, a baseline measurement is also required.

Details on the approach to handling missing data for safety analyses are provided in Section 8.2.5 Statistical Methods.

8.2.5 Statistical Methods

Statistical testing and inference for safety analyses are described in 8.2.5.2. Efficacy results that will be considered to be statistically significant after consideration of the strategy for controlling the Type I error are described in Section 8.2.6, Multiplicity. Nominal p-values may be computed for other efficacy analyses as a measure of strength of association between the endpoint and the treatment effect rather than formal tests of hypotheses. Unless otherwise stated, all statistical tests will be conducted at the α=0.05 (2-sided) level.
8.2.5.1 Statistical Methods for Efficacy Analyses

Efficacy will be evaluated separately in each cohort. For the primary efficacy endpoint independent centrally reviewed RECIST 1.1 response rate, the point estimate, exact 95% Clopper-Pearson confidence interval, and p-value for testing the RECIST 1.1 response rate is greater than the historical control for each cohort will be provided using exact binomial distribution. Subjects in the primary analysis population (FAS) without response data will be counted as non-responder.

Secondary efficacy evaluations of RECIST 1.1 response based on Investigator assessment will also be conducted using the same methodology as for the primary efficacy analysis. Efficacy for the secondary analysis of RECIST 1.1 response by independent central radiology review among subjects with HPV positive head and neck cancer will be evaluated in the same manner as in the primary analysis.

For PFS endpoint, Kaplan-Meier (KM) curves and median estimates from the KM curves will be provided as appropriate. Subjects without efficacy evaluation data or without survival data will be censored at Day 1. Table 11 summarizes the key efficacy analyses.

Table 11 Analysis Strategy for Key Efficacy Variables

<table>
<thead>
<tr>
<th>Endpoint/Variable</th>
<th>Statistical Method</th>
<th>Analysis Population</th>
<th>Missing Data Approach</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review (Cohort A, Cohort B HPV negative subjects, Cohort C, and Cohort D evaluated separately)</td>
<td>Exact test of binomial parameter</td>
<td>FAS (Primary) ASaT (Supportive)</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review for subjects in Cohort B2</td>
<td>Exact test of binomial parameter</td>
<td>FAS (Primary) ASaT (Supportive)</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review, Cohort B HPV positive subjects</td>
<td>Exact test of binomial parameter</td>
<td>FAS (Primary) ASaT (Supportive)</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on independent central radiology review, Cohort D AP subjects</td>
<td>Exact test of binomial parameter</td>
<td>FAS (Primary) ASaT (Supportive)</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on based on independent central radiology review, for subjects previously treated with cetuximab and platinum in Cohorts B and B2</td>
<td>Exact test of binomial parameter</td>
<td>FAS (Primary) ASaT (Supportive)</td>
<td>Subjects with missing data are considered non-responders</td>
</tr>
<tr>
<td>Overall RECIST 1.1 response rate based on investigator assessment for cohorts A,B, C and D</td>
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</tr>
<tr>
<td>Endpoint/Variable(^c) (Description, Time Point)</td>
<td>Statistical Method</td>
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</tr>
<tr>
<td>-----------------------------------------------</td>
<td>-------------------</td>
<td>---------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>Progression-free survival</td>
<td>Summary statistics using Kaplan-Meier method</td>
<td>FAS(Primary) ASaT (Supportive)</td>
<td>Censored at last assessment</td>
</tr>
<tr>
<td>Overall survival</td>
<td>Summary statistics using Kaplan-Meier method</td>
<td>FAS(Primary) ASaT (Supportive)</td>
<td>Censored at last assessment</td>
</tr>
<tr>
<td>Response duration</td>
<td>Summary statistics using Kaplan-Meier method</td>
<td>All responders</td>
<td>Non-responders are excluded in analysis</td>
</tr>
</tbody>
</table>

The strategy to address multiplicity issues with regard to multiple efficacy endpoints is described in Section 8.2.6, Multiplicity and Section 8.2.9, Interim Analyses.

8.2.5.2 Statistical Methods for Safety Analyses

Safety and tolerability will be assessed by clinical review of all relevant parameters including adverse experiences (AEs), laboratory tests, and vital signs.

Summary statistics (counts, percentage, mean, standard deviation, etc.) will be provided for the safety endpoints as appropriate. The 80% confidence interval for the incidence rate of Grade 2 or higher adverse events with an immune etiology and the incidence rate of Grade 4/5 AEs will be provided as appropriate.

8.2.5.3 Summaries of Baseline Characteristics, Demographics, and Other Analyses

8.2.5.3.1 Demographic and Baseline Characteristics

Baseline characteristics will be assessed by the use of tables and/or graphs for each cohort. No statistical hypothesis tests will be performed on these characteristics. The number and percentage of subjects screened, randomized, the primary reasons for screening failure, and the primary reason for discontinuation will be displayed. Demographic variables (e.g., age, gender), baseline characteristics, primary and secondary diagnoses, and prior and concomitant therapies will be summarized by treatment either by descriptive statistics or categorical tables.

8.2.6 Multiplicity

The false positive rate for testing the primary efficacy endpoint in each cohort (Cohort A, Cohort B, Cohort C, Cohort D, and Cohort B2) is controlled at 0.025 (1-sided) for each cohort.

If the primary hypothesis in Cohort B2 is successful, then alpha=0.025 (1-sided) will be passed to the secondary evaluation of subjects in Cohort B and Cohort B2 with prior exposure to cetuximab and platinum therapy in a step down approach.
8.2.7 Sample Size and Power Calculations

The calculation of power and sample size for each cohort does not account for interim analysis. Cohort A (triple negative breast cancer subjects): With approximately 26 evaluable PD-L1 positive subjects with triple negative breast cancer, the study has approximately 80% power to detect a 25% difference in ORR under the null hypothesis of ORR=20% with a type I error rate of 2.5% if the true ORR is 45%. The null hypothesis rate of 20% is based on the historic response rate in large phase III trials for standard single agent chemotherapy in 2nd line breast cancer utilizing WHO-based bi-dimensional responses [34, 35, 36]. Success for this hypothesis requires at least 10/26 responses. The actual number of subjects enrolled may be larger than 26 to ensure that at least 26 subjects are evaluable for analysis.

Cohort B (head and neck cancer subjects): HPV negative head and neck cancer subjects will be evaluated separately from HPV positive head and neck cancer subjects. With a maximum of 22 evaluable PD-L1 positive subjects with HPV negative head and neck cancer, the study has approximately 80% power to detect a 25% difference in ORR under the null hypothesis of ORR=10% with a type I error rate of 2.5% if the true ORR is 35%. Success for this hypothesis requires at least 6/22 responses. The actual number of subjects enrolled may be larger than 22 to ensure that at least 22 subjects are evaluable for analysis.

With a maximum of 12 evaluable PD-L1 positive subjects with HPV positive head and neck cancer, the study has approximately 73% power to detect a 35% difference in ORR under the null ORR=20% with a type I error rate of 5% if the true ORR is 55%. The null hypothesis rate of 20% is based on historic response of chemotherapy in head and neck cancer trials [37, 38]. Success for this hypothesis requires at least 6 responses. The actual number of subjects enrolled may be larger than 12 to ensure that at least 12 subjects are evaluable for analysis. However, if 6 responses are observed prior to 12 subjects with HPV positive head and neck cancer enrolling in the trial, enrollment may be stopped in this group as the hypothesis success criterion will have been reached.

Cohort C (urothelial tract cancer subjects): With a maximum of 22 evaluable PD-L1 positive subjects with urothelial tract cancer, the study has approximately 80% power to detect a 25% difference in ORR under the null hypothesis of ORR=10% with a type I error rate of 2.5% if the true ORR is 35%. The null hypothesis rate of 10% is based on the historic response rate of single agent pemetrexed (RR 6%) or vinflunine (RR 9%) chemotherapy for bladder cancer [39, 40]. Success for this hypothesis requires at least 6/22 responses. The actual number of subjects enrolled may be larger than 22 to ensure that at least 22 subjects are evaluable for analysis.

Cohort D (gastric cancer subjects): With a maximum of 32 evaluable PD-L1 positive subjects with gastric cancer, stratified to include 16 Asia Pacific (AP) and 16 non-AP, the study has approximately 90% power to detect a 25% difference in ORR under the null hypothesis of ORR=15% with a type I error rate of 2.5% if the true ORR is 40%. Success for this hypothesis requires at least 10/32 responses. A total of 16 AP subjects will provide approximately 80% power with type I error rate of 0.1 to detect a 25% difference in ORR under the null hypothesis of ORR=15% if the true ORR is 40%. At least 4/16 responses will
be needed to claim the success of this hypothesis for the AP population. The null hypothesis rate of 15% is based on the response rate in large (n > 100), trials for standard of care chemotherapy irinotecan/mFOLFIRI (RR 12%), and irinotecan/docetaxel (RR 13%) for 2nd line metastatic gastric cancer [28, 29]. The actual number of subjects enrolled may be larger than 32 to ensure that at least 32 subjects are evaluable for analysis.

Cohort B2 (head and neck cancer subjects, expansion cohort): With 100 evaluable subjects in Cohort B2, the study provides >99% power to detect a 15% difference in ORR under the null hypothesis of ORR=5% with a type I error rate of 2.5% if the true ORR is 20%. Success for this hypothesis requires at least 11/100 responses. The null hypothesis rate of 5% is based on historic response of chemotherapy in head and neck cancer trials [37]. The actual number of subjects enrolled may be larger than 100 to ensure at least 100 subjects are evaluable for analysis.

Cohorts B and B2 previously treated with cetuximab and platinum: With 60 evaluable head and neck cancer subjects previously treated with cetuximab and platinum, the study has 93% power to detect a 15% difference in ORR under the null hypothesis of ORR=5% with a type I error rate of 2.5% if the true ORR is 20%. Success for this hypothesis requires at least 8/60 responses. The null hypothesis rate of 5% is based on historic response of chemotherapy in head and neck cancer trials [37].

8.2.8 Subgroup Analyses and Effect of Baseline Factors

Efficacy will be summarized for the key endpoints for the following subgroups:

- HPV-positive vs. HPV negative head/neck cancer
- Subjects with head/neck cancer enrolled in Cohort B and Cohort B2 with prior cetuximab and platinum exposure.

8.2.9 Interim Analyses

8.2.9.1 Efficacy Interim Analyses

No efficacy interim analyses are planned in this trial.

However, an interim analysis for Cohort A, B, or C may be performed if the rate of enrollment is much slower than anticipated during the course of the trial. An interim analysis for each cohort would only be performed in this study when ≥10 subjects in the respective cohort have had scans through Week 16. Results will be reviewed by the study team.

For Cohort A, if an interim analysis is conducted and 2 or fewer subjects out of the first 10 subjects with post-baseline imaging scans (Week 16) have a confirmed or unconfirmed response, then enrollment may be paused until response data for the subsequent imaging scan (Week 24) are reviewed for subjects already enrolled in the trial. Enrollment may be resumed if the binomial probability of the observed response rate is ≥ 80% under the assumption of a ≥ 20% true response rate (e.g., ≥4/16 confirmed or unconfirmed response).
The operating characteristics of the interim analysis rule are provided in Table 12. The power in Cohort A (assuming true ORR of 45%) decreases to ~77% if the interim analysis is conducted.

Table 12  Operating Characteristics of Interim Analysis Rule for Triple Negative Breast Cancer – Pause Enrollment if ≤ 2/10 Subjects Respond

<table>
<thead>
<tr>
<th>True ORR</th>
<th>Probability of Pausing Enrollment</th>
<th>Probability of Study Success</th>
</tr>
</thead>
<tbody>
<tr>
<td>10%</td>
<td>0.93</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>15%</td>
<td>0.82</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>20%</td>
<td>0.68</td>
<td>0.02</td>
</tr>
<tr>
<td>25%</td>
<td>0.53</td>
<td>0.08</td>
</tr>
<tr>
<td>30%</td>
<td>0.38</td>
<td>0.21</td>
</tr>
<tr>
<td>35%</td>
<td>0.26</td>
<td>0.39</td>
</tr>
<tr>
<td>40%</td>
<td>0.17</td>
<td>0.60</td>
</tr>
<tr>
<td>45%</td>
<td>0.10</td>
<td>0.77</td>
</tr>
<tr>
<td>50%</td>
<td>0.06</td>
<td>0.88</td>
</tr>
</tbody>
</table>

For Cohorts B (HPV negative) and C, if an interim analysis is conducted and 1 or fewer subjects out of the first 10 subjects with post-baseline imaging scans (Week 16) have a confirmed or unconfirmed response, then enrollment may be paused until response data for the subsequent imaging scan (Week 24) are reviewed for subjects already enrolled in the trial. Enrollment may be resumed if the binomial probability of the observed response rate is ≥ 80% under the assumption of a ≥ 10% true response rate (e.g., ≥2/16 confirmed or unconfirmed response). The operating characteristics of the interim analysis rule are provided in Table 13. The power in Cohort B (HPV negative) and Cohort C (assuming true ORR of 35%) does not substantially decrease from 80% as a result of conducting the interim analysis.

Table 13  Operating Characteristics of Interim Analysis Rule for Head/Neck Cancer (HPV negative) and Urothelial Cancer– Pause Enrollment if ≤ 1/10 subjects respond

<table>
<thead>
<tr>
<th>True ORR</th>
<th>Probability of Pausing Enrollment</th>
<th>Probability of Study Success</th>
</tr>
</thead>
<tbody>
<tr>
<td>5%</td>
<td>0.91</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>10%</td>
<td>0.74</td>
<td>0.02</td>
</tr>
<tr>
<td>15%</td>
<td>0.54</td>
<td>0.09</td>
</tr>
<tr>
<td>20%</td>
<td>0.38</td>
<td>0.25</td>
</tr>
<tr>
<td>25%</td>
<td>0.25</td>
<td>0.45</td>
</tr>
<tr>
<td>30%</td>
<td>0.15</td>
<td>0.65</td>
</tr>
<tr>
<td>35%</td>
<td>0.09</td>
<td>0.80</td>
</tr>
<tr>
<td>40%</td>
<td>0.05</td>
<td>0.90</td>
</tr>
<tr>
<td>45%</td>
<td>0.02</td>
<td>0.95</td>
</tr>
</tbody>
</table>
8.2.9.2 Safety Interim Analyses

Although safety monitoring will occur continuously in the study, an assessment of Grade 4/5 drug-related immunologic AEs will be performed after 10 subjects have been enrolled within each cohort for 1 cycle of therapy. Enrollment will be paused if 4 or more subjects out of the first 10 subjects within a cohort experience a Grade 4/5 drug-related immunologic adverse event. Enrollment may be resumed within a cohort only after a full safety evaluation is performed in consultation between the study investigators and the Sponsor.

The estimate of and the upper bound of the 95% confidence interval for the underlying percentage of subjects with a Grade 4/5 drug-related immunologic adverse event given various hypothetical observed number of subjects are provided in Table 14. These calculations are based on the exact binomial method proposed by Clopper and Pearson [41].

Table 14 Estimate of Incidence of Grade 4/5 events and 95% Upper Confidence Bound Based on Hypothetical Number of Subjects with Event Out of 10 Subjects Evaluated

<table>
<thead>
<tr>
<th>Hypothetical Number of Subjects With Grade 4/5 Event</th>
<th>Estimate of Incidence</th>
<th>95% Upper Confidence Bound†</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0%</td>
<td>30.9%</td>
</tr>
<tr>
<td>1</td>
<td>10%</td>
<td>44.5%</td>
</tr>
<tr>
<td>2</td>
<td>20%</td>
<td>55.6%</td>
</tr>
<tr>
<td>3</td>
<td>30%</td>
<td>65.2%</td>
</tr>
<tr>
<td>4</td>
<td>40%</td>
<td>73.8%</td>
</tr>
<tr>
<td>5</td>
<td>50%</td>
<td>81.3%</td>
</tr>
</tbody>
</table>

† Based on the two-tailed exact confidence interval of a binomial proportion [41].

8.2.10 Compliance (Medication Adherence)

A day within the study will be considered an On-Therapy day if the subject receives the study medication infusion. The number of days on therapy is the total number of days from the first day of study medication to the date of the last dose of study medication. Compliance with trial treatment administration will be measured by the number of administered infusions divided by the number of infusions that were supposed to be administered as determined by the number of days on therapy.

Summary statistics for the number of Days on Therapy will be provided by treatment group for the FAS population.

8.2.11 Extent of Exposure

Extent of Exposure for a subject is defined as number of cycles in which the subject receives the study medication infusion. Summary statistics will be provided on Extent of Exposure for ASaT population.
9.0 LABELING, PACKAGING, STORAGE AND RETURN OF CLINICAL SUPPLIES

9.1 Investigational Product

The investigator shall take responsibility for and shall take all steps to maintain appropriate records and ensure appropriate supply, storage, handling, distribution and usage of investigational product in accordance with the protocol and any applicable laws and regulations.

Clinical Supplies will be provided by the Sponsor as summarized in Table 15.

Table 15  Product Descriptions

<table>
<thead>
<tr>
<th>Product Name &amp; Potency</th>
<th>Dosage Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>MK-3475 50 mg</td>
<td>Lyophilized Powder for Injection</td>
</tr>
<tr>
<td>MK-3475 100 mg/ 4mL</td>
<td>Solution for Injection</td>
</tr>
</tbody>
</table>

9.2 Packaging and Labeling Information

Clinical supplies will be affixed with a clinical label in accordance with regulatory requirements.

Vials will be provided in an open label fashion for subject dosing.

9.3 Clinical Supplies Disclosure

This trial is open-label; therefore, the subject, the trial site personnel, the Sponsor and/or designee are not blinded to treatment. Drug identity (name, strength) is included in the label text; random code/disclosure envelopes or lists are not provided.

9.4 Storage and Handling Requirements

Clinical supplies must be stored in a secure, limited-access location under the storage conditions specified on the label.

Receipt and dispensing of trial medication must be recorded by an authorized person at the trial site.

Clinical supplies may not be used for any purpose other than that stated in the protocol.

9.5 Returns and Reconciliation

The investigator is responsible for keeping accurate records of the clinical supplies received from the Sponsor or designee, the amount dispensed to and returned by the subjects and the amount remaining at the conclusion of the trial.

For all trial sites, the local country Sponsor personnel or designee will provide appropriate documentation that must be completed for drug accountability and return.
9.6 Standard Policies

Trial site personnel will have access to a central electronic randomization system (IVRS/IWRS system) to allocate subjects, to assign drug to subjects and to manage the distribution of clinical supplies. Each person accessing the IVRS system must be assigned an individual unique PIN. They must use only their assigned PIN to access the system, and they must not share their assigned PIN with anyone.

10.0 ADMINISTRATIVE AND REGULATORY DETAILS

10.1 Confidentiality

10.1.1 Confidentiality of Data

By signing this protocol, the investigator affirms to the Sponsor that information furnished to the investigator by the Sponsor will be maintained in confidence, and such information will be divulged to the institutional review board, ethics review committee (IRB/ERC) or similar or expert committee; affiliated institution and employees, only under an appropriate understanding of confidentiality with such board or committee, affiliated institution and employees. Data generated by this trial will be considered confidential by the investigator, except to the extent that it is included in a publication as provided in the Publications section of this protocol.

10.1.2 Confidentiality of Subject Records

By signing this protocol, the investigator agrees that the Sponsor (or Sponsor representative), IRB/ERC, or regulatory authority representatives may consult and/or copy trial documents in order to verify worksheet/case report form data. By signing the consent form, the subject agrees to this process. If trial documents will be photocopied during the process of verifying worksheet/case report form information, the subject will be identified by unique code only; full names initials will be masked prior to transmission to the Sponsor.

By signing this protocol, the investigator agrees to treat all subject data used and disclosed in connection with this trial in accordance with all applicable privacy laws, rules and regulations.

10.1.3 Confidentiality of Investigator Information

By signing this protocol, the investigator recognizes that certain personal identifying information with respect to the investigator, and all subinvestigators and trial site personnel, may be used and disclosed for trial management purposes, as part of a regulatory submissions, and as required by law. This information may include:

1. name, address, telephone number and e-mail address;
2. hospital or clinic address and telephone number;
3. curriculum vitae or other summary of qualifications and credentials; and
4. other professional documentation.
Consistent with the purposes described above, this information may be transmitted to the Sponsor, and subsidiaries, affiliates and agents of the Sponsor, in your country and other countries, including countries that do not have laws protecting such information. Additionally, the investigator’s name and business contact information may be included when reporting certain serious adverse events to regulatory authorities or to other investigators. By signing this protocol, the investigator expressly consents to these uses and disclosures.

If this is a multicenter trial, in order to facilitate contact between investigators, the Sponsor may share an investigator’s name and contact information with other participating investigators upon request.

10.1.4 Confidentiality of IRB/IEC Information

The Sponsor is required to record the name and address of each IRB/IEC member that reviews and approves this trial. The Sponsor is also required to document that each IRB/IEC meets regulatory and ICH GCP requirements by requesting and maintaining records of the names and qualifications of the IRB/IEC members and to make these records available for regulatory agency review upon request by those agencies.

10.2 Compliance with Financial Disclosure Requirements

Financial Disclosure requirements are outlined in the US Food and Drug Administration Regulations, Financial Disclosure by Clinical Investigators (21 CFR Part 54). It is the Sponsor's responsibility to determine, based on these regulations, whether a request for Financial Disclosure information is required. It is the investigator's/subinvestigator's responsibility to comply with any such request.

The investigator/subinvestigator(s) agree, if requested by the Sponsor in accordance with 21 CFR Part 54, to provide his/her financial interests in and/or arrangements with the Sponsor to allow for the submission of complete and accurate certification and disclosure statements. The investigator/subinvestigator(s) further agree to provide this information on a Certification/Disclosure Form, commonly known as a financial disclosure form, provided by the Sponsor or through a secure password-protected electronic portal provided by the Sponsor. The investigator/subinvestigator(s) also consent to the transmission of this information to the Sponsor in the United States for these purposes. This may involve the transmission of information to countries that do not have laws protecting personal data.

10.3 Compliance with Law, Audit and Debarment

By signing this protocol, the investigator agrees to conduct the trial in an efficient and diligent manner and in conformance with this protocol; generally accepted standards of Good Clinical Practice (e.g., International Conference on Harmonization of Technical Requirements for Registration of Pharmaceuticals for Human Use Good Clinical Practice: Consolidated Guideline and other generally accepted standards of good clinical practice); and all applicable federal, state and local laws, rules and regulations relating to the conduct of the clinical trial.
The Code of Conduct, a collection of goals and considerations that govern the ethical and scientific conduct of clinical investigations sponsored by Merck, is provided in Section 12.1 - Merck Code of Conduct for Clinical Trials.

The investigator also agrees to allow monitoring, audits, IRB/ERC review and regulatory authority inspection of trial-related documents and procedures and provide for direct access to all trial-related source data and documents.

The investigator agrees not to seek reimbursement from subjects, their insurance providers or from government programs for procedures included as part of the trial reimbursed to the investigator by the Sponsor.

The investigator shall prepare and maintain complete and accurate trial documentation in compliance with Good Clinical Practice standards and applicable federal, state and local laws, rules and regulations; and, for each subject participating in the trial, provide all data, and, upon completion or termination of the clinical trial, submit any other reports to the Sponsor as required by this protocol or as otherwise required pursuant to any agreement with the Sponsor.

Trial documentation will be promptly and fully disclosed to the Sponsor by the investigator upon request and also shall be made available at the trial site upon request for inspection, copying, review and audit at reasonable times by representatives of the Sponsor or any regulatory authorities. The investigator agrees to promptly take any reasonable steps that are requested by the Sponsor as a result of an audit to cure deficiencies in the trial documentation and worksheets/case report forms.

The investigator must maintain copies of all documentation and records relating to the conduct of the trial in compliance with all applicable legal and regulatory requirements. This documentation includes, but is not limited to, the protocol, worksheets/case report forms, advertising for subject participation, adverse event reports, subject source data, correspondence with regulatory authorities and IRBs/ERCs, consent forms, investigator’s curricula vitae, monitor visit logs, laboratory reference ranges, laboratory certification or quality control procedures and laboratory director curriculum vitae. By signing this protocol, the investigator agrees that documentation shall be retained until at least 2 years after the last approval of a marketing application in an ICH region or until there are no pending or contemplated marketing applications in an ICH region or until at least 2 years have elapsed since the formal discontinuation of clinical development of the investigational product. Because the clinical development and marketing application process is variable, it is anticipated that the retention period can be up to 15 years or longer after protocol database lock. The Sponsor will determine the minimum retention period and notify the investigator when documents may be destroyed. The sponsor also recognizes that documents may need to be retained for a longer period if required by local regulatory requirements. All trial documents shall be made available if required by relevant regulatory authorities. The investigator must consult with and obtain written approval by the Sponsor prior to discarding trial and/or subject files.
ICH Good Clinical Practice guidelines recommend that the investigator inform the subject’s primary physician about the subject’s participation in the trial if the subject has a primary physician and if the subject agrees to the primary physician being informed.

The investigator will promptly inform the Sponsor of any regulatory authority inspection conducted for this trial.

Persons debarred from conducting or working on clinical trials by any court or regulatory authority will not be allowed to conduct or work on this Sponsor’s trials. The investigator will immediately disclose in writing to the Sponsor if any person who is involved in conducting the trial is debarred or if any proceeding for debarment is pending or, to the best of the investigator’s knowledge, threatened.

In the event the Sponsor prematurely terminates a particular trial site, the Sponsor will promptly notify that trial site’s IRB/IEC.

According to European legislation, a Sponsor must designate an overall coordinating investigator for a multi-center trial (including multinational). When more than one trial site is open in an EU country, Merck, as the Sponsor, will designate, per country, a national principal coordinator (Protocol CI), responsible for coordinating the work of the principal investigators at the different trial sites in that Member State, according to national regulations. For a single-center trial, the Protocol CI is the principal investigator. In addition, the Sponsor must designate a principal or coordinating investigator to review the trial report that summarizes the trial results and confirm that, to the best of his/her knowledge, the report accurately describes the conduct and results of the trial [Clinical Study Report (CSR) CI]. The Sponsor may consider one or more factors in the selection of the individual to serve as the Protocol CI and or CSR CI (e.g., availability of the CI during the anticipated review process, thorough understanding of clinical trial methods, appropriate enrollment of subject cohort, timely achievement of trial milestones). The Protocol CI must be a participating trial investigator.

10.4 Compliance with Trial Registration and Results Posting Requirements

Under the terms of the Food and Drug Administration Modernization Act (FDAMA) and the Food and Drug Administration Amendments Act (FDAAA), the Sponsor of the trial is solely responsible for determining whether the trial and its results are subject to the requirements for submission to the Clinical Trials Data Bank, http://www.clinicaltrials.gov. Merck, as Sponsor of this trial, will review this protocol and submit the information necessary to fulfill these requirements. Merck entries are not limited to FDAMA/FDAAA mandated trials. Information posted will allow subjects to identify potentially appropriate trials for their disease conditions and pursue participation by calling a central contact number for further information on appropriate trial locations and trial site contact information.

By signing this protocol, the investigator acknowledges that the statutory obligations under FDAMA/FDAAA are that of the Sponsor and agrees not to submit any information about this trial or its results to the Clinical Trials Data Bank.
10.5 Quality Management System

By signing this protocol, the Sponsor agrees to be responsible for implementing and maintaining a quality management system with written development procedures and functional area standard operating procedures (SOPs) to ensure that trials are conducted and data are generated, documented, and reported in compliance with the protocol, accepted standards of Good Clinical Practice, and all applicable federal, state, and local laws, rules and regulations relating to the conduct of the clinical trial.

10.6 Data Management

The investigator or qualified designee is responsible for recording and verifying the accuracy of subject data. By signing this protocol, the investigator acknowledges that his/her electronic signature is the legally binding equivalent of a written signature. By entering his/her electronic signature, the investigator confirms that all recorded data have been verified as accurate.

Detailed information regarding Data Management procedures for this protocol will be provided by the Sponsor.

10.7 Publications

This trial is intended for publication, even if terminated prematurely. Publication may include any or all of the following: posting of a synopsis online, abstract and/or presentation at a scientific conference, or publication of a full manuscript. The Sponsor will work with the authors to submit a manuscript describing trial results within 12 months after the last data become available, which may take up to several months after the last subject visit in some cases such as vaccine trials. However, manuscript submission timelines may be extended on OTC trials. For trials intended for pediatric-related regulatory filings, the investigator agrees to delay publication of the trial results until the Sponsor notifies the investigator that all relevant regulatory authority decisions on the trial drug have been made with regard to pediatric-related regulatory filings. Merck will post a synopsis of trial results for approved products on www.clinicaltrials.gov by 12 months after the last subject's last visit for the primary outcome, 12 months after the decision to discontinue development, or product marketing (dispensed, administered, delivered or promoted), whichever is later.

These timelines may be extended for products that are not yet marketed, if additional time is needed for analysis, to protect intellectual property, or to comply with confidentiality agreements with other parties. Authors of the primary results manuscript will be provided the complete results from the Clinical Study Report, subject to the confidentiality agreement. When a manuscript is submitted to a biomedical journal, the Sponsor's policy is to also include the protocol and statistical analysis plan to facilitate the peer and editorial review of the manuscript. If the manuscript is subsequently accepted for publication, the Sponsor will allow the journal, if it so desires, to post on its website the key sections of the protocol that are relevant to evaluating the trial, specifically those sections describing the trial objectives and hypotheses, the subject inclusion and exclusion criteria, the trial design and procedures,
the efficacy and safety measures, the statistical analysis plan, and any amendments relating to those sections. The Sponsor reserves the right to redact proprietary information.

For multicenter trials, subsequent to the multicenter publication (or after public disclosure of the results online at www.clinicaltrials.gov if a multicenter manuscript is not planned), an investigator and his/her colleagues may publish their data independently. In most cases, publication of individual trial site data does not add value to complete multicenter results, due to statistical concerns. In rare cases, publication of single trial site data prior to the main paper may be of value. Limitations of single trial site observations in a multicenter trial should always be described in such a manuscript.

Authorship credit should be based on 1) substantial contributions to conception and design, or acquisition of data, or analysis and interpretation of data; 2) drafting the article or revising it critically for important intellectual content; and 3) final approval of the version to be published. Authors must meet conditions 1, 2 and 3. Significant contributions to trial execution may also be taken into account to determine authorship, provided that contributions have also been made to all three of the preceding authorship criteria. Although publication planning may begin before conducting the trial, final decisions on authorship and the order of authors’ names will be made based on participation and actual contributions to the trial and writing, as discussed above. The first author is responsible for defending the integrity of the data, method(s) of data analysis and the scientific content of the manuscript.

The Sponsor must have the opportunity to review all proposed abstracts, manuscripts or presentations regarding this trial 45 days prior to submission for publication/presentation. Any information identified by the Sponsor as confidential must be deleted prior to submission; this confidentiality does not include efficacy and safety results. Sponsor review can be expedited to meet publication timelines.
11.0 LIST OF REFERENCES


12.0 APPENDICES

12.1 Merck Code of Conduct for Clinical Trials

Merck*
Code of Conduct for Clinical Trials

I. Introduction

A. Purpose

Merck, through its subsidiaries, conducts clinical trials worldwide to evaluate the safety and effectiveness of our products. As such, we are committed to designing, implementing, conducting, analyzing and reporting these trials in compliance with the highest ethical and scientific standards. Protection of subject safety is the overriding concern in the design of clinical trials. In all cases, Merck clinical trials will be conducted in compliance with local and/or national regulations and in accordance with the ethical principles that have their origin in the Declaration of Helsinki.

B. Scope

Such standards shall be endorsed for all clinical interventional investigations sponsored by Merck irrespective of the party (parties) employed for their execution (e.g., contract research organizations, collaborative research efforts). This Code is not intended to apply to trials which are observational in nature, or which are retrospective. Further, this Code does not apply to investigator-initiated trials which are not under the control of Merck.

II. Scientific Issues

A. Trial Conduct

• Trial Design

Except for pilot or estimation trials, clinical trial protocols will be hypothesis-driven to assess safety, efficacy and/or pharmacokinetic or pharmacodynamic indices of Merck or comparator products. Alternatively, Merck may conduct outcomes research trials, trials to assess or validate various endpoint measures, or trials to determine subject preferences, etc.

The design (i.e., subject population, duration, statistical power) must be adequate to address the specific purpose of the trial. Research subjects must meet protocol entry criteria to be enrolled in the trial.

• Site Selection

Merck selects investigative sites based on medical expertise, access to appropriate subjects, adequacy of facilities and staff, previous performance in Merck trials, as well as budgetary considerations. Prior to trial initiation, sites are evaluated by Merck personnel to assess the ability to successfully conduct the trial.

• Site Monitoring/Scientific Integrity

Trial sites are monitored to assess compliance with the trial protocol and general principles of Good Clinical Practice. Merck reviews clinical data for accuracy, completeness and consistency. Data are verified versus source documentation according to standard operating procedures. Per Merck policies and procedures, if fraud, misconduct or serious GCP-non-Compliance are suspected, the issues are promptly investigated. When necessary, the clinical site will be closed, the responsible regulatory authorities and ethics review committees notified and data disclosed accordingly.

B. Publication and Authorship

To the extent scientifically appropriate, Merck seeks to publish the results of trials it conducts. Some early phase or pilot trials are intended to be hypothesis-generating rather than hypothesis testing. In such cases, publication of results may not be appropriate since the trial may be underpowered and the analyses complicated by statistical issues of multiplicity.

Merck’s policy on authorship is consistent with the requirements outlined in the ICH-Good Clinical Practice guidelines. In summary, authorship should reflect significant contribution to the design and conduct of the trial, performance or interpretation of the analysis, and/or writing of the manuscript. All named authors must be able to defend the trial results and conclusions. Merck funding of a trial will be acknowledged in publications.
III. Subject Protection

- IRB/ERC review
  All clinical trials will be reviewed and approved by an independent IRB/ERC before being initiated at each site. Significant changes or revisions to the protocol will be approved by the IRB/ERC prior to implementation, except that changes required urgently to protect subject safety and well-being may be enacted in anticipation of IRB/ERC approval. For each site, the IRB/ERC and Merck will approve the subject informed consent form.

- Safety
  The guiding principle in decision-making in clinical trials is that subject welfare is of primary importance. Potential subjects will be informed of the risks and benefits of, as well as alternatives to, trial participation. At a minimum, trial designs will take into account the local standard of care. Subjects are never denied access to appropriate medical care based on participation in a Merck clinical trial.

  All participation in Merck clinical trials is voluntary. Subjects are enrolled only after providing informed consent for participation. Subjects may withdraw from a Merck trial at any time, without any influence on their access to, or receipt of, medical care that may otherwise be available to them.

- Confidentiality
  Merck is committed to safeguarding subject confidentiality, to the greatest extent possible. Unless required by law, only the investigator, sponsor (or representative) and/or regulatory authorities will have access to confidential medical records that might identify the research subject by name.

- Genomic Research
  Genomic Research will only be conducted in accordance with informed consent and/or as specifically authorized by an Ethics Committee.

IV. Financial Considerations

- Payments to Investigators
  Clinical trials are time- and labor-intensive. It is Merck’s policy to compensate investigators (or the sponsoring institution) in a fair manner for the work performed in support of Merck trials. Merck does not pay incentives to enroll subjects in its trials. However, when enrollment is particularly challenging, additional payments may be made to compensate for the time spent in extra recruiting efforts.

  Merck does not pay for subject referrals. However, Merck may compensate referring physicians for time spent on chart review to identify potentially eligible subjects.

- Clinical Research Funding
  Informed consent forms will disclose that the trial is sponsored by Merck, and that the investigator or sponsoring institution is being paid or provided a grant for performing the trial. However, the local IRB/ERC may wish to alter the wording of the disclosure statement to be consistent with financial practices at that institution. As noted above, publications resulting from Merck trials will indicate Merck as a source of funding.

- Funding for Travel and Other Requests
  Funding of travel by investigators and support staff (e.g., to scientific meetings, investigator meetings, etc.) will be consistent with local guidelines and practices including, in the U.S., those established by the American Medical Association (AMA).

V. Investigator Commitment

  Investigators will be expected to review Merck’s Code of Conduct as an appendix to the trial protocol, and in signing the protocol, agree to support these ethical and scientific standards.

* In this document, "Merck" refers to Merck Sharp & Dohme Corp. and Schering Corporation, each of which is a subsidiary of Merck & Co., Inc. Merck is known as MSD outside of the United States and Canada. As warranted by context, Merck also includes affiliates and subsidiaries of Merck & Co., Inc."
12.2 Collection and Management of Specimens for Future Biomedical Research

1. Definitions

1. Biomarker: A biological molecule found in blood, other body fluids, or tissues that is a sign of a normal or abnormal process or of a condition or disease. A biomarker may be used to see how well the body responds to a treatment for a disease or condition.³

2. Pharmacogenomics: The investigation of variations of DNA and RNA characteristics as related to drug response.²

3. Pharmacogenetics: A subset of pharmacogenomics, pharmacogenetics is the influence of variations in DNA sequence on drug response.²

4. DNA: Deoxyribonucleic acid.

5. RNA: Ribonucleic acid.

2. Scope of Future Biomedical Research

The DNA and tumor tissue specimen(s) collected in the current trial will be used to study various causes for how subjects may respond to a drug. The DNA and tumor tissue specimen(s) will be stored to provide a resource for future trials conducted by Merck focused on the study of biomarkers responsible for how a drug enters and is removed by the body, how a drug works, other pathways a drug may interact with, or other aspects of disease. The specimen(s) may be used for future assay development and/or drug/vaccine development.

It is now well recognized that information obtained from studying and testing clinical specimens offers unique opportunities to enhance our understanding of how individuals respond to drugs, enhance our understanding of human disease and ultimately improve public health through development of novel treatments targeted to populations with the greatest need. All specimens will be used by Merck or designees and research will be monitored and reviewed by a committee of our scientists and clinicians.

3. Summary of Procedures for Future Biomedical Research

- Subjects for Enrollment

All subjects enrolled in the clinical trial will be considered for enrollment in the Future Biomedical Research sub-trial.

- Informed Consent

Informed consent for specimens (i.e., DNA, RNA, protein, etc.) will be obtained during screening for protocol enrollment from all subjects or legal guardians, at a trial visit by the investigator or his or her designate. Informed consent for Future Biomedical Research should be presented to the subjects on Visit 1. If delayed, present consent at next possible Subject Visit. Informed consent must be obtained prior to collection of all Future Biomedical Research specimens. Consent forms
signed by the subject will be kept at the clinical trial site under secure storage for regulatory reasons. Information contained on the consent form alone cannot be traced to any specimens, test results, or medical information once the specimens have been rendered de-identified.

Subjects are not required to participate in the Future Biomedical Research sub-trial in order to participate in the main trial. Subjects who decline to sign the Future Biomedical Research informed consent will not have the specimen collected nor will they be discontinued from the main trial.

A template of each trial site’s approved informed consent will be stored in the Sponsor’s clinical document repository. Each consent will be assessed for appropriate specimen permissions.

Each informed consent approved by an ethics committee is assigned a unique tracking number. The tracking number on this document will be used to assign specimen permissions for each specimen into the Entrusted Keyholder’s Specimen Database.

- **eCRF Documentation for Future Biomedical Research Specimens**
  
  Documentation of both consent and acquisition of Future Biomedical Research specimens will be captured in the electronic Case Report Forms (eCRFs). Reconciliation of both forms will be performed to assure that only appropriately-consented specimens are used for this sub-trial's research purposes. Any specimens for which such an informed consent cannot be verified will be destroyed.

- **Future Biomedical Research Specimen Collections**
  
  Blood specimens for DNA or RNA isolation will usually be obtained at a time when the subject is having blood drawn for other trial purposes. Specimens like tissue and bone marrow will usually be obtained at a time when the subject is having such a procedure for clinical purposes.

  Specimens will be collected and sent to the laboratory designated for the trial where they will be processed (e.g., DNA or RNA extraction, etc) following the Merck approved policies and procedures for specimen handling and preparation.

If specimens are collected for a specific genotype or expression analysis as an objective to the main trial, this analysis is detailed in the main body of this protocol (**Section 8.0 – Statistical Analysis Plan**). These specimens will be processed, analyzed, and the remainder of the specimen will be destroyed. The results of these analyses will be reported along with the other trial results. A separate specimen will be obtained from properly-consented subjects in this protocol for storage in the biorepository for Future Biomedical Research.
4. Confidential Subject Information for Future Biomedical Research

In order to optimize the research that can be conducted with Future Biomedical Research specimens, it is critical to link subject' clinical information with future test results. In fact little or no research can be conducted without connecting the clinical trial data to the specimen. The clinical data allow specific analyses to be conducted. Knowing subject characteristics like gender, age, medical history and treatment outcomes are critical to understanding clinical context of analytical results.

To maintain privacy of information collected from specimens obtained for Future Biomedical Research, Merck has developed secure policies and procedures. All specimens will be de-identified as described below.

At the clinical trial site, unique codes will be placed on the Future Biomedical Research specimens for transfer to the storage facility. This first code is a random number which does not contain any personally identifying information embedded within it. The link (or key) between subject identifiers and this first unique code will be held at the trial site. No personal identifiers will appear on the specimen tube.

This first code will be replaced with a second code at a Merck designated storage/lab facility. The second code is linked to the first code via a second key. The specimen is now double coded. Specimens with the second code are sometimes referred to as de-identified specimens. The use of the second code provides additional confidentiality and privacy protection for subjects over the use of a single code. Access to both keys would be needed to link any data or specimens back to the subject's identification.

The second code is stored separately from the first code and all associated personal specimen identifiers. A secure link, the second key, will be utilized to match the second code to the first code to allow clinical information collected during the course of the trial to be associated with the specimen. This second key will be transferred under secure procedures by the Merck designated facility to an Entrusted Keyholder at Merck. The second code will be logged into the primary biorepository database at Merck and, in this database, this identifier will not have identifying demographic data or identifying clinical information (i.e., race, sex, age, diagnosis, lab values) associated with it. The specimen will be stored in a designated biorepository site with secure policies and procedures for specimen storage and usage.

The second key can be utilized to reconstruct the link between the results of future biomedical research and the clinical information, at the time of analysis. This linkage would not be possible for the scientist conducting the analysis, but can only be done by the Merck Entrusted Keyholder under strict security policies and procedures. The Merck Entrusted Keyholder will link the information and then issue a de-identified data set for analysis. The only other circumstance by which future biomedical research data would be directly linked to the full clinical data set would be those situations mandated by regulatory authorities (e.g., EMEA, FDA), whereby this information would be directly transferred to the regulatory authority.
5. Biorepository Specimen Usage

Specimens obtained for the Merck Biorepository will be used for analyses using good scientific practices. However, exploratory analyses will not be conducted under the highly validated conditions usually associated with regulatory approval of diagnostics. The scope of research performed on these specimens is limited to the investigation of the variability in biomarkers that may correlate with a clinical phenotype in subjects.

Analyses utilizing the Future Biomedical Research specimens may be performed by Merck, or an additional third party (e.g., a university investigator) designated by Merck. The investigator conducting the analysis will be provided with double coded specimens. Re-association of analysis results with corresponding clinical data will only be conducted by the Merck Entrusted Keyholder. Any contracted third party analyses will conform to the specific scope of analysis outlined in this sub-trial. Future Biomedical Research specimens remaining with the third party after the specific analysis is performed will be returned to the sponsor or destroyed and documentation of destruction will be reported to Merck.

6. Withdrawal From Future Biomedical Research

Subjects may withdraw their consent for Future Biomedical Research and have their specimens and all derivatives destroyed. Subjects may withdraw consent at any time by writing to the principal investigator for the main trial. If medical records for the main trial are still available, the investigator will contact Merck using the designated mailbox (clinical.specimen.management@merck.com) and a form will be provided by Merck to obtain appropriate information to complete specimen withdrawal. Subsequently, the subject's specimens will be removed from the biorepository and be destroyed. A letter will be sent from Merck to the investigator confirming the destruction. It is the responsibility of the investigator to inform the subject of completion of destruction. Any analyses in progress at the time of request for destruction or already performed prior to the request being received by the Sponsor will continue to be used as part of the overall research trial data and results. No new analyses would be generated after the request is received.

In the event that the medical records for the main trial are no longer available (e.g., if the investigator is no longer required by regulatory authorities to retain the main trial records) or the specimens have been completely anonymized, there will no longer be a link between the subject’s personal information and their specimens. In this situation, the request for specimen destruction can not be processed.

7. Retention of Specimens

Future Biomedical Research specimens will be stored in the biorepository for potential analysis for up to 20 years from acquisition. Specimens may be stored for longer if a regulatory or governmental authority has active questions that are being answered. In this special circumstance, specimens will be stored until these questions have been adequately addressed.
Specimens from the trial site will be shipped to a central laboratory and then shipped to the Merck designated biorepository. The specimens will be stored under strict supervision in a limited access facility which operates to assure the integrity of the specimens. Specimens will be destroyed according to Merck policies and procedures and this destruction will be documented in the biorepository database.

8. **Data Security**

Separate databases for specimen information and for results from the Future Biomedical Research sub-trial will be maintained by Merck. This is done to separate the future exploratory test results (which include genetic data) from the clinical trial database thereby maintaining a separation of subject number and these results. The separate databases are accessible only to the authorized Sponsor and the designated trial administrator research personnel and/or collaborators. Database user authentication is highly secure, and is accomplished using network security policies and practices based in international standards (e.g., ISO17799) to protect against unauthorized access. The Merck Entrusted Keyholder maintains control over access to all specimen data. These data are collected for future biomedical research purposes only as specified in this sub-trial will not be used for any other purpose.

9. **Reporting of Future Biomedical Research Data to Subjects**

There is no definitive requirement in either authoritative ethical guidelines or in relevant laws/regulations globally that research results have to be, in all circumstances, returned to the trial participant. Some guidelines advocate a proactive return of data in certain instances. No information obtained from exploratory laboratory studies will be reported to the subject or family, and this information will not be entered into the clinical database maintained by Merck on subjects. Principle reasons not to inform or return results to the subject include: lack of relevance to subject health, limitations of predictive capability, concerns of misinterpretation and absence of good clinical practice standards in exploratory research typically used for diagnostic testing.

If any exploratory results are definitively associated with clinical significance for subjects while the clinical trial is still ongoing, investigators will be contacted with information as to how to offer clinical diagnostic testing (paid for by Merck) to subjects enrolled and will be advised that counseling should be made available for all who choose to participate in this diagnostic testing.

If any exploratory results are definitively associated with clinical significance after completion of a clinical trial, Merck will publish the results without revealing specific subject information, inform all trial sites who participated in the Merck clinical trial and post anonymized results on our website or other accredited website(s) that allow for public access (e.g., disease societies who have primary interest in the results) in order that physicians and patients may pursue clinical diagnostic testing if they wish to do so.
10. Gender, Ethnicity and Minorities

Although many diagnoses differ in terms of frequency by ethnic population and gender, every effort will be made to recruit all subjects diagnosed and treated on Merck clinical trials for future biomedical research. When trials with specimens are conducted and subjects identified to serve as controls, every effort will be made to group specimens from subjects and controls to represent the ethnic and gender population representative of the disease under current investigation.

11. Risks Versus Benefits of Future Biomedical Research

For future biomedical research, risks to the subject have been minimized. [Risks include those associated with venipuncture to obtain the whole blood specimen. This specimen will be obtained at the time of routine blood specimens drawn in the main trial. Therefore, there will not be an additional risk for the subject.]

Merck has developed strict security, policies and procedures to address subject data privacy concerns. Data privacy risks are largely limited to rare situations involving possible breach of confidentiality. In this highly unlikely situation there is risk that the information, like all medical information, may be misused.

It is necessary for subject-related data (i.e., ethnicity, diagnosis, drug therapy and dosage, age, toxicities, etc.) to be re-associated to double coded specimens at the time of data analysis. These subject data will be kept in a separate, secure Merck database, and all specimens will be stripped of subject identifiers. No information concerning results obtained from future biomedical research will be entered into clinical records, nor will it be released to outside persons or agencies, in any way that could be tied to an individual subject.

12. Self-Reported Ethnicity

Subjects who participate in future biomedical research will be asked to provide self-reported ethnicity. Subjects who do not wish to provide this data may still participate in future biomedical research.

13. Questions

Any questions related to the future biomedical research should be e-mailed directly to clinical.specimen.management@merck.com.

14. References


12.3 Understanding the Intent, Scope and Public Health Benefits of Exploratory Biomarker Research: A Guide for IRBs/IECs and Investigational Site Staff
This informational brochure is intended for IRBss/IECs and Investigational Site Staff. The brochure addresses issues relevant to specimen collection for biomarker research in the context of pharmaceutical drug and vaccine development.

1. What is a Biomarker and What is Biomarker Research?

A biomarker is a “characteristic that is objectively measured and evaluated as an indicator of normal biological processes, pathogenic processes, or pharmacologic responses to a therapeutic intervention”.

Biomarker research, including research on pharmacogenomic biomarkers, is a tool used to improve the development of pharmaceuticals and understanding of disease. It involves the analysis of biomolecules (such as DNA, RNA, proteins, and lipids), or other measurements (such as blood pressure or brain images) in relation to clinical endpoints of interest. Biomarker research can be influential across all phases of drug development, from drug discovery and preclinical evaluations to clinical development and post-marketing studies. This brochure focuses on biomarker research involving analysis of biomolecules from biological samples collected in clinical trials. Please refer to I-PWG Pharmacogenomic Informational Brochure and ICH Guidance E16 for additional information specific to pharmacogenomic biomarkers.

2. Why is Biomarker Research Important?

Importance to Patients and Public Health
Biomarker research is helping to improve our ability to predict, detect, and monitor diseases and improve our understanding of how individuals respond to drugs. This research underlies personalized medicine: a tailored approach to patient treatment based on the molecular analysis of genes, proteins, and metabolites. The goal of biomarker research is to aid clinical decision-making toward safer and more efficacious courses of treatment, improved patient outcomes, and overall cost-savings. It also allows for the continued development and availability of drugs that are effective in certain sub-populations when they otherwise might not have been developed due to insufficient efficacy in the broader population.

Recent advances in biomedical technology, including genetic and molecular medicine, have greatly increased the power and precision of analytical tools used in health research and have accelerated the drive toward personalized medicine. In some countries, highly focused initiatives have been created to promote biomarker research (e.g., in the US: www.fda.gov/oc/initiatives/criticalpath/; in the EU: www.imi.europa.eu/index_en.html).

Importance to Drug Development
Biomarker research is being used by the pharmaceutical industry to streamline the drug development process. Some biomarkers are used as substitutes or “surrogates” for safety or efficacy endpoints in clinical trials particularly where clinical outcomes or events cannot practically or ethically be measured (e.g., cholesterol as a surrogate for cardiovascular disease). By using biomarkers to assess patient response, ineffective drug candidates may be terminated earlier in the development process in favor of more promising drug candidates. Biomarkers are being used to optimize clinical trial designs and outcomes by identifying patient populations that are more likely to respond to a drug therapy or to avoid specific adverse events.
Biomarker research is also being used to enhance scientific understanding of the mechanisms of both treatment response and disease processes, which can help to identify future targets for drug development. Depending on the clinical endpoints in a clinical trial, biomarker sample collection may either be a required or optional component of the trial. However, both mandatory and optional sample collections are important for drug development.

3. Importance of Biomarkers to Regulatory Authorities

Regulatory health authorities are increasingly aware of the benefits of biomarkers and how they may be used for drug approval, clinical trial design, and clinical care. Biomarkers have been used to establish risk/benefit profiles. For example, the FDA has modified the US warfarin (Coumadin®) label to include the analysis of CYP2C9 and VKORC1 genes to guide dosing regimens. Health authorities such as the FDA (USA), EMEA (European Union), MHLW (Japan), and ICH (International) are playing a key role in advancing this scientific field as it applies to pharmaceutical development by creating the regulatory infrastructure to facilitate this research. Numerous regulatory guidances and concept papers have already been issued, many of which are available through www.i-pwg.org. Global regulatory authorities have highlighted the importance of biomarker research and the need for the pharmaceutical industry to take the lead in this arena.6-14

4. How are Biomarkers Being Used in Drug/Vaccine Development?

Biomarker research is currently being used in drug/vaccine development to:

- Explain variability in response among participants in clinical trials
- Better understand the mechanism of action or metabolism of investigational drugs
- Obtain evidence of pharmacodynamic activity (i.e., how the drug affects the body) at the molecular level
- Address emerging clinical issues such as unexpected adverse events
- Determine eligibility for clinical trials to optimize trial design
- Optimize dosing regimens to minimize adverse reactions and maximize efficacy
- Develop drug-linked diagnostic tests to identify patients who are more likely or less likely to benefit from treatment or who may be at risk of experiencing adverse events
- Provide better understanding of mechanisms of disease
- Monitor clinical trial participant response to medical interventions

Biomarker research, including research on banked samples, should be recognized as an important public health endeavor for the overall benefit of society, whether by means of advancement of medical science or by development of safer and more effective therapies.7 Since the value of collected samples may increase over time as scientific discoveries are made, investment in long-term sample repositories is a key component of biomarker research.
5. Biomarkers are Already a Reality in Health Care

A number of drugs now have biomarker information included in their labels. Biomarker tests are already being used in clinical practice to serve various purposes:

**Predictive biomarkers (efficacy)** – In clinical practice, predictive efficacy biomarkers are used to predict which patients are most likely to respond, or not respond, to a particular drug. Examples include: i) Her2 overexpression analysis required for prescribing trastuzumab (Herceptin®) to breast cancer patients, ii) c-kit expression analysis prior to prescribing imatinib mesylate (Gleevec®) to gastrointestinal stromal tumor patients, and iii) EMLI mutational status testing prior to prescribing panitumumab (Vedibix®) or cetuximab (Erbitux®) to metastatic colorectal cancer patients.

**Predictive biomarkers (safety)** – In clinical practice, predictive safety biomarkers are used to select the proper drug dose or to evaluate the appropriateness of continued therapy in the event of a safety concern. Examples include: i) monitoring of blood potassium levels in patients receiving drosperone and ethinyl estradiol (Yasmin®) together with daily long-term drug regimens that may increase serum potassium, and ii) prospective HLA-B*5701 screening to identify those at increased risk for hypersensitivity to abacavir (Ziagen®).

**Surrogate biomarkers** – In clinical practice, surrogate biomarkers may be used as alternatives to measures such as survival or irreversible morbidity. Surrogate biomarkers are measures that are reasonably likely, based on epidemiologic, therapeutic, pathophysiological, or other evidence, to predict clinical benefit. Examples include: i) LDL level as a surrogate for risk of cardiovascular diseases in patients taking lipid-lowering agents such as atorvastatin calcium (Liptor®), ii) blood glucose as a surrogate for clinical outcomes in patients taking anti-diabetic agents, and iii) HIV plasma viral load and CD4 cell counts as surrogates for time-to-clinical-events and overall survival in patients receiving antiretroviral therapy for HIV disease.

**Prognostic biomarkers** – Biomarkers can also help predict clinical outcomes independent of any treatment modality. Examples of prognostic biomarkers used in clinical practice include: i) CellSearch™ to predict progression-free survival in breast cancer, ii) anti-CCP (cyclic citrullinated protein) for the severity of rheumatoid arthritis, iii) estrogen receptor status for breast cancer, and iv) anti-DNA for the severity of systemic lupus erythematosus.

0. Biomarker Samples from Clinical Trials: An Invaluable Resource

Adequate sample sizes and high-quality data from controlled clinical trials are key to advancements in biomarker research. Samples collected in clinical trials create the opportunity for investigation of biomarkers related to specific drugs, drug classes, and disease areas. Clinical drug development programs are therefore an invaluable resource and a unique opportunity for highly productive biomarker research. In addition to conducting independent research, pharmaceutical companies are increasingly contributing to consortia efforts by pooling samples, data, and expertise in an effort to conduct rigorous and efficient biomarker research and to maximize the probability of success.

7. Informed Consent for Collection & Banking of Biomarker Samples

Collection of biological samples in clinical trials must be undertaken with voluntary informed consent of the participant (or legally-acceptable representative).
and regulations for legally-appropriate informed consent vary on national, state, and local levels, but are generally based on internationally recognized pillars of ethical conduct for research on human subjects.28-31

Optional vs. Required Subject Participation
Depending on the relevance of biomarker research to a clinical development program at the time of protocol development, the biomarker research may be a core required component of a trial (e.g., key to elucidating the drug mechanism of action or confirming that the drug is interacting with the target) or may be optional (e.g., to gain valuable knowledge that enhances the understanding of diseases and drugs). Informed consent for the collection of biomarker samples may be presented either in the main clinical informed consent form or as a separate informed consent form, with approaches varying somewhat across pharmaceutical companies. The relevance of biomarker research to a clinical development program may change over time as the science evolves. The samples may therefore increase in value after a protocol is developed.

Consent for Future Research Use
While it can be a challenge to specify the details of the research that will be conducted in the future, the I-PWG holds the view that future use of samples collected for exploratory biomarker research in clinical trials should be permissible when (i) the research is scientifically sound, (ii) participants are informed of the scope of the intended future research, even if this is broadly defined (see potential uses in Section 4 above), (iii) autonomy is respected by providing the option to consent separately to future use of samples or by providing the option to terminate further use of samples upon request (consent withdrawal / sample destruction), and (iv) industry standards for confidentiality protection per Good Clinical Practice guidelines are met.28, 91 Importantly, any research using banked samples should be consistent with the original informed consent, except where otherwise permitted by local law or regulation.

Important elements of informed consent for future use of samples include, but are not limited to:28

- The scope of research: Where the scope of the potential future research is broad, participants should be informed of the boundaries of the research. While it may not be possible to describe the exact analytical techniques that will be used, or specific molecules that will be analyzed, it is possible to clearly articulate in reasonable detail the type of research to be conducted and its purpose. Information regarding whether stored samples may be shared with other parties or utilized for commercialization purposes should also be addressed.

- Withdrawal of consent / sample destruction: The informed consent form should inform participants of their right to withdraw their consent / request destruction of their samples. This should include the mechanisms for exercising that right and any limitations to exercising that right. For example, participants should be informed that it is not possible to destroy samples that have been anonymized.28 In addition, according to industry standards and regulatory guidance, participants should be informed that data already generated prior to a consent withdrawal request are to be maintained as part of the study data.28

- The duration of storage: The permissible duration of storage may vary according to the nature and uses of the samples and may also vary on national, state, and local levels. The intended duration of storage, including indefinite storage, should be specified.
Biomarker Research in Clinical Trials

1. Clinical trial participants undergo the informed consent procedure and sign the informed consent form.

2. Biological samples are collected from clinical trial participants.

3. Scientists analyze the samples in the laboratory for biomarkers (e.g., DNA, RNA, proteins, lipids).

4. Test results are analyzed using various bioinformatic and statistical tools.

5. Biomarker research ultimately leads to the development of better drugs and treatment regimens.

6. With appropriate consent, biological samples are stored for future research.

7. As science evolves, research can be performed in the future on stored samples.

INFORMED CONSENT
SAMPLE COLLECTION
LABORATORY TESTS
DATA ANALYSIS
LONG-TERM STORAGE
DRUG DEVELOPMENT
8. Biomarker Sample Collection in Different Countries

Collection of biological samples for biomarker research is straightforward in most jurisdictions. Some countries have specific laws and regulations regarding collection, labeling, storage, export, and/or use of exploratory samples. In addition, some regulations distinguish between DNA and non-DNA samples or between samples used for diagnostic purposes and samples collected for scientific research. Processes for the collection, labeling, storage, export, and/or use of biomarker samples should always adhere to the laws and regulations of the country/region in which those samples are collected.

9. Return of Research Results to Study Participants

Policies for the return of biomarker research results to study participants who request them vary among pharmaceutical companies. There are many considerations that pharmaceutical companies weigh when determining their policy regarding the return of biomarker research results to study participants. These include:

i) the conditions under which biomarker research results were generated (i.e., exploratory research laboratory versus accredited diagnostic laboratory)

ii) whether the results will have an impact on the medical care of the participant or on a related person, if applicable

iii) whether genetic counseling is recommended for genetic results

iv) the ability to accurately link the result to the individual from whom the sample was collected

v) international, national, and local guidelines, policies, legislation, and regulations regarding participants’ rights to access data generated on them

Meng et al. 2000 and Article 29 Data Protection Working Party (an advisory committee to the European Commission on the European Data Protection Directive) have addressed these considerations in detail in relation to pharmacogenomic research data and provided a list of documents addressing the general issue of return of research results.29,30

10. Benefits and Risks Associated with Biomarker Research

Benefits

While it may not always directly benefit the study participant who is providing the samples, biomarker research can improve overall understanding of disease and treatment of future patients receiving therapies developed from such research. Patients are now benefiting from retrospective biomarker research conducted on samples collected from clinical trials and stored for exploratory research. One example is the recent label update to the EGFR antibody drugs cetuximab (Erbitux®) and panitumumab (Vectibix®) which highlights the value of KRAS status as a predictive biomarker for treatment of metastatic colorectal cancer with this class of drugs.

The humanitarian benefit of human research is recognized by the Nuremberg Code.28,33 Provided that the degree of risk does not exceed that determined by the humanitarian importance of the problem to be solved, research participants should not be denied the right to contribute to the greater common good.28,34

Risks

Risks associated with biomarker research are primarily related to the physical aspects of obtaining the sample and to patient privacy concerns.

Physical risks associated with biomarker sample collection in clinical trials can be characterized in two ways:

i) negligible additional risk when the biomarker sample is collected as part of a procedure conducted to support
other core trial objectives, and ii) some added risk where
the sampling procedure would otherwise have not been
performed as a core component of a trial. Risks are also
determined by the invasiveness of the sample collection
procedure.

Privacy risks are generally those associated with the inap-
propriate disclosure and misuse of data. Pharmaceutical
companies have policies and procedures for confidentiality
protection to minimize this risk for all data collected
and generated in clinical trials. These may vary across
companies, but are based on industry standards of confi-
dentiality and privacy protection highlighted in the fol-
lowing section. Importantly, privacy risks inherent to bio-
marker data are no greater than other data collected in a
clinical trial.

11. Privacy, Confidentiality, and
Patient Rights

Maintaining the privacy of study participants and the con-
fidentiality of information relating to them is of paramount
concern to industry researchers, regulators, and patients.
Good Clinical Practice (GCP), the standard adhered to
in pharmaceutical clinical research, is a standard that

"...provides assurance that the data and reported results
are credible and accurate, and that the rights, integ-
ity, and confidentiality of trial subjects are protected",\n
where confidentiality is defined as, "the prevention of dis-
closure, to other than authorized individuals, of a spon-
sor’s proprietary information or of a subject’s identity."

This standard dictates that "the confidentiality of
records that could identify subjects should be protect-
ed, respecting the privacy and confidentiality rules in
accordance with applicable regulatory requirements." \n
Exploratory biomarker research in pharmaceutical de-
velopment is commonly conducted in research laboratories
that are not accredited to perform diagnostic tests used
for healthcare decision making. Therefore, results from
exploratory biomarker research usually are not appro-
priate for use in making decisions about a trial par-
ticipant’s health. In addition, exploratory research data
should not be included as part of a participant’s med-
cal record accessible for use by insurance companies.
Legislation and policies to protect individuals against
discrimination based on genetic information continually
evolve based on social, ethical, and legal considerations.
Examples of such legislation include the Human Tissue
Act 2004 (UK) and the Genetic Information Nondiscrimina-
tion Act (GINA) 2008 (USA). 96,47

12. Where to Get More Information?

Educational resources related to biomarker and pharma-
cogenomic research that caters to health care profession-
als, IRBs/IECs, scientists, and patients are continually
being created and are publicly available. Links to many of
these resources are available through the I-PWG website:
www.i-pwg.org.

13. What is I-PWG?

The Industry Pharmacogenomics Working Group (I-PWG)
(formerly the Pharmacogenetics Working Group) is a vol-
untary association of pharmaceutical companies engaged in
pharmacogenomic research. The Group’s activities focus on
non-competitive educational, informational, ethical, legal,
and regulatory topics. The Group provides information and
expert opinions on these topics and sponsors educational/
informational programs to promote better understanding
of pharmacogenomic and other biomarker research for key
stakeholders. The I-PWG interacts with regulatory author-

I-PWG
INDUSTRY PHARMACOGENOMICS WORKING GROUP

MK-3475-012-05 Final Protocol
04TSOJ
Confidential
11-Dec-2017
Product: MK-3475 (SCH 900475)
Protocol/Amendment No.: 012-05

15. References


12.4 ECOG Performance Status

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<th>Description</th>
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<tr>
<td>0</td>
<td>Normal activity. Fully active, able to carry on all pre-disease performance without restriction.</td>
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<tr>
<td>1</td>
<td>Symptoms, but ambulatory. Restricted in physically strenuous activity, but ambulatory and able to carry out work of a light or sedentary nature (e.g., light housework, office work).</td>
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<tr>
<td>2</td>
<td>In bed &lt;50% of the time. Ambulatory and capable of all self-care, but unable to carry out any work activities. Up and about more than 50% of waking hours.</td>
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<tr>
<td>3</td>
<td>In bed &gt;50% of the time. Capable of only limited self-care, confined to bed or chair more than 50% of waking hours.</td>
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<tr>
<td>4</td>
<td>100% bedridden. Completely disabled. Cannot carry on any self-care. Totally confined to bed or chair.</td>
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<tr>
<td>5</td>
<td>Dead.</td>
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12.5 Common Terminology Criteria for Adverse Events V4.0 (CTCAE)

The descriptions and grading scales found in the revised NCI Common Terminology Criteria for Adverse Events (CTCAE) version 4.0 will be utilized for adverse event reporting. (http://ctep.cancer.gov/reporting/ctc.html)
12.6 Response Evaluation Criteria in Solid Tumors (RECIST) 1.1 Criteria For Evaluating Response in Solid Tumors

RECIST version 1.1* will be used in this study for assessment of tumor response. While either CT or MRI may be utilized, as per RECIST 1.1, CT is the preferred imaging technique in this study.

* As published in the European Journal of Cancer:


In addition, volumetric analysis will be explored by central review for response assessment.
13.0 SIGNATURES

13.1 Sponsor’s Representative

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13.2 Investigator

I agree to conduct this clinical trial in accordance with the design outlined in this protocol and to abide by all provisions of this protocol (including other manuals and documents referenced from this protocol). I agree to conduct the trial in accordance with generally accepted standards of Good Clinical Practice. I also agree to report any serious adverse events as defined in Section 7.0 – Assessing and Recording Adverse Events. I also agree to handle all clinical supplies provided by the Sponsor and handle all clinical specimens in accordance with the protocol. I understand that information that identifies me will be used and disclosed as described in the protocol, and that such information may be transferred to countries that do not have laws protecting such information. Since the information in this protocol and the referenced Investigator’s Brochure is confidential, I understand that its disclosure to any third parties, other than those involved in approval, supervision, or conduct of the trial is prohibited. I will ensure that the necessary precautions are taken to protect such information from loss, inadvertent disclosure or access by third parties.

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