



## NGOC Clinical Trials List - Printable Version

### Head and Neck (C00-C14)

1. Locally Advanced
2. 1st Line Recurrent / Metastatic
3. Previously Treated

### Esophagus/Gastric (C15-C16)

1. Stage II- IV, Neo-adjuvant
2. First Line, Recurrent or Metastatic

**a. AZ D702AC00001 (Artemide-Gastric01) A Phase 3 Study of Rilvegostomig in Combination with Fluoropyrimidine**

Open 8/12/25

- i. Arm A: Rilvegostomig + T-Dxd + Investigator's Choice: 5-FU or Capecitabine
- ii. Arm B: Pembro + trastuzumab + Investigator's choice: Cisplatin+ 5FU or CAPOX+capecitabine
- iii. Arm C: Rilvegostomic (750mg IV Q3w) + Trastuzumab + Investigator's Choice: Cisplatin+5FU OR CAPOX+Capecitabine

3. Previously Treated

### Biliary (K83.9)

1. First Line Therapy

**a. AZ D702NC00001 A Global Phase III Study of Rilvegostomig Plus Chemotherapy for First-**

Open 3/25/26

- i. Arm A: Rilvegostomig 750mg on D1 of each Q3W+ Gemcitabine 1000 mg/m2 IV Q3W plus cisplatin 25 mg/m2 IV Q3W will be administered on D1 and D8 of each cycle starting on C1D1 for up to 8 cycles
- ii. Arm B: Durvalumab 1500mg on D1 of each Q4W cycle +Gemcitabine 1000 mg/m2 IV Q3W plus cisplatin 25 mg/m2 IV Q3W will be administered on D1 and D8 of each cycle starting on C1D1 for up to 8 cycles

### Pancreas (C25)

1. Adjuvant / NeoAdjuvant
2. 1st Line, Metastatic

**a. BMS CA240-0030 (MountainTAP) Study of BMS-986504 in Combination with Nab-p/Gem versus Placebo in Co**

Open 9/15/25

- i. Phase 2 Dose Selection Arm A: BMS-986504 400mg + Nab-Pac + Gem
- ii. Phase 2 Dose Selection Arm B: BMS-986504 600mg + Nab-Pac + Gem
- iii. Phase 2 Dose Selection Arm C: BMS-986504 400mg Placebo + Nab-Pac + Gem
- iv. Phase 2 Dose Selection Arm D: BMS-986504 600mg Placebo + Nab-Pac + Gem
- v. NOT YET OPEN Phase 3 Arm E: BMS-986504 Optimal Dose + Nab-Pac + Gem
- vi. NOT YET OPEN Phase 3 Arm F: BMS-986504 Optimal Dose Placebo + Nab-Pac + Gem

3. Previously Treated

### Colon & Rectum (C18-C20)

1. Stage III, Resected Colon, Adjuvant
2. Stage IV, Metastatic, 1st Line, Colon or Rectum
3. Metastatic, 1st Line, Colorectal

**a. BMS CA266-0003 (ROSETTA CRC-203) Pumitamig in Combination with Chemotherapy Versus Bevacizumab in Combi**

Open 2/24/26

- i. Phase 2 Arm A1: pumitamig (BMS-986545) Dose Level 1, Day 1 + FOLFOX or FOLFIRI, Q2W
- ii. Phase 2 Arm A2: pumitamig (BMS-986545) Dose Level 2, Day 1 + FOLFOX or FOLFIRI, Q2W
- iii. Phase 2 Arm B: bevacizumab, Day 1 + FOLFOX or FOLFIRI, Q2W
- iv. NOT OPEN Phase 3 Arm C: pumitamig RP3D, Day 1 + FOLFOX or FOLFIRI, Q2W; or pumitamig RP3D + CAPOX, Q3W
- v. NOT OPEN Phase 3 Arm D: bevacizumab, Day 1 + FOLFOX or FOLFIRI, Q2W or bevacizumab + CAPOX, Q3W

4. Metastatic, 3rd Line, Colon
5. Metastatic 2nd or 3rd Line, KRAS Mutated
6. Metastatic 2nd or 3rd Line, KRAS Wild Type
7. Refractory

### Lung, Squamous Cell (C34)

1. Stage IV, Metastatic
2. Second Line

### Lung, Non-Small Cell (C34)

1. Stage IB, II, IIIA; Adjuvant
2. Stage 1B, II, Resectable IIIA
3. Stage II-III Resectable or Resected

**a. BI 1479-0032 (Beamion Lung-3) A Phase 3, Beamion LUNG-3: A study to test whether zongertinib helps**

Open 3/19/26

- i. Arm A: Zongertinib (BI 1810631) 120mg po qd
- ii. Arm B: Physicians Choice( Nivolumab 480mg, pembrolizumab 200mg, atezolizumab 1200mg, or durvalumab 1500mg)

4. Stage IIIA-IIIIB, Locally Advanced, Unresectable
5. Stage IIB-IIIIB Resected and Chemotherapy-Naive Stage IV
6. Stage IV, Metastatic
7. First Line- Metastatic

**a. AZ D702GC00001 Artemide Lung-04 A Global Phase III Study of Rilvegostomig or Pembrolizumab Monotherapy**

Open 1/7/26

- i. Arm A: Rilvegostomig 750 mg IV Q3W
- ii. Arm B: Pembrolizumab 200 mg IV Q3W

**b. BMS CA266-0002(ROSETTA Lung-202) A Study of Pumitamig Versus Pembrolizumab in Participants With Previou**

Open 4/15/26

- i. Arm A: Pumitamig 1200 mg (if body weight < 50 kg) or 1500 mg (if body weight ≥ 50 kg) as a 60-minute IV infusion every 3 weeks (Q3W)
- ii. Arm B: Pembrolizumab 200 mg as a 60-minute IV infusion every 3 weeks (Q3W)

8. 2nd or 3rd Line
9. Metastatic, Previously Treated

**a. AbbV M25-274 (TEImet NSCLC-04) A Phase 2, Open-Label, Randomized, Global Study of Three Telisotuzumab**

Open 10/20/25

- i. Arm A: Telisotuzumab vedotin 1.9 mg/kg Q2W
- ii. Arm B: Telisotuzumab vedotin 1.6 mg/kg Q2W
- iii. Arm C: Telisotuzumab vedotin 1.6 mg/kg Q2W x 4 cycles then 1.9 mg/kg Q2W

**a. AZ D763QC00001 (TROPION-Lung17) Datopotamab Deruxtecan or Docetaxel in Previously Treated TROP2-posit**

Open 11/13/25

- i. Arm A: Dato-DXd 6mg/kg IV q3w
- ii. Arm B: Docetaxel 75mg/m2 IV q3w

## Mesothelioma (C45)

1.

## Lung, Small Cell (C34)

1. Extensive Stage, Small Cell Lung Ca

**a. BMS CA245-0001 A Randomized, Double-Blind, Multicenter Phase 3 Trial**

Open 1/24/25

- i. Arm A: BMS-986489 420mg/nivolumab 360 mg IV+carboplatin+etoposide
- ii. Arm B: atezolizumab 1200mg IV+carboplatin+etoposide

2. Previously Treated

## Breast, Adjuvant (C50)

1. NeoAdjuvant, Stage I, II, III

2. Adjuvant

**a. GS-US-595-6184 (ASCENT-05) A Randomized, Open-label, Phase 3 Study of Adjuvant Sacituzumab Govite**

Open 1/31/23

- i. Arm A: Sacituzumab Govitecan 10mg/kg IV Days 1 and 8 + Pembrolizumab 200mg IV Day 1 Q3W X 8 cycles
- ii. Arm B: Treatment of Physician's Choice: Pembrolizumab 200mg IV Day 1 Q3 weeks X 8 cycles OR Pembrolizumab 200 mg IV Day 1 and Capecitabine 1000 mg/m<sup>2</sup> PO BID D1-D14 Q3 Weeks X 8 cycles

## Breast, Her-2 Positive (C50)

1. NeoAdjuvant HER-2 Positive

2. Adjuvant, HER-2 Positive

3. Metastatic, HER-2 Positive, 1st Line

4. Metastatic, Previously Treated, Her-2 Positive

## Breast, Advanced (C50)

1. Stage IV, Metastatic, 1st Line

**a. RGN WO45654 (INAVO123) A PHASE III, MULTICENTER, RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED**

Open 5/23/25

- i. Arm A: Inavolisib 9mg tablet PO QD days 1-28 of 28 day cycle + + letrozole 2.5mg PO QD + Investigator's choice - palbociclib 125mg PO QD D1-21 of 28 or ribociclib 600mg D1-21 of 28d cycle
- ii. Arm B: Placebo tablet PO QD days 1-28 of 28 day cycle + + letrozole 2.5mg PO QD + Investigator's choice - palbociclib 125mg PO QD D1-21 of 28 or ribociclib 600mg D1-21 of 28d cycle

2. Stage IV, Metastatic, ER -

3. Stage IV, Metastatic, Previously Treated

4. Stage IV, Metastatic, Triple Negative Breast Cancer

## Breast, ER Positive (C50)

1. Adjuvant

**a. AZ D8535C00001 (Cambria-2) Adjuvant Endocrine-based Therapy Study of Camizestrant (AZD9833) in ER**

On Hold 4/1/26

- i. Arm A: Standard ET (AI or tamoxifen)+/-abemaciclib (+/-LHRH agonist\*)
- ii. Arm B: Camizestrant 75 mg/daily +/- abemaciclib (+/- LHRH agonist\*)

2. Metastatic, First Line

3. Metastatic, Prior Hormonal Therapy

## Melanoma (C43)

1. Adjuvant

2. Locally advanced or Metastatic

3. Previously Treated

## Ovarian (C56)

1. Previously Treated

## Bladder (C67)

1.

2. Urothelial

## Renal (C64)

1. Adjuvant, Phase III

2. First Line Treatment, Advanced or Metastatic

3. Advanced, Metastatic

4. Previously Treated

## Prostate (C61)

1. Hormone Refractory, 1st Line

2. 1st line Metastatic

3. Metastatic, 2nd Line Prostate

4. Hormone Refractory; Bone Predominant 3rd Line or Greater

## Sarcoma (C46-C49)

1. Sarcoma, Previously Treated

## Adrenal (C74)

1. Previously Treated

## Lymphoma (C82-C88)

1. Indolent, previously treated

2. Follicular, First Line

3. Indolent, Relapsed

4. Relapsed or Refractory; DLBCL, FL

5. Large Cell Lymphoma, Relapsed/Refractory

6. Relapsed and Refractory Follicular Lymphoma

**a. BMS CA073-1003 A Phase 3, Multicenter, Randomized, Open Label Study to Compare the Ef**

Open 8/13/25

- i. Arm A: R-Golca 0.4mg D1-14/48 days+Rituximab x5cycles followed by golcadomide monotherapy for up to 12 cycles of toral therapy
- ii. Arm B: R-Len 20mg, Days 1-21, cycles 1-12,rituximab 375mg/m<sup>2</sup>, Days 1,8,15,and 22 in cycle 1; Day 1 in cycles 2-5
- iii. Arm C: Inv. Choice for 6 cycles: R-CHOP or Rituximab-Bendamustine

Accrual Goal = 3

7. Relapsed and Refractory Follicular or Marginal Zone Lymphoma

8. Diffuse Large Cell, First Line

**a. CC-220-DLBCL-001 First line of Therapy**

Open 3/2/26

- i. Single arm extension cohort of CC-99282 with the Polatuzumab R-CHP regimen. CC-99282 dose will be determined upon completion of Part 2A and 2B.

9. Mantle Cell Lymphoma

**CNS (C71)**

1. Glioblastoma Multiforme

**Leukemia (C91-C92)**

1. CLL, First Line
2. CLL, Relapsed or Refractory
3. CML

**Myeloma (C90)**

1. 1st Line
2. Relapsed / Refractory 1-3 Lines
3. Relapsed / Refractory > or equal to 2 Lines

**MDS (D46)**

1. 1st Line
2. Observational Registry

**Unknown Primary (199)**

1. 1st Line

**Other**

1. Tissue Studies
2. Observational Studies

**Multiple Sites**

1. Phase I
2. EMD MS100070-0176 (EMR100070-001 Rollover)
3. Novartis Signature Series
4. Solid Tumors, Refractory
5. Solid Tumors, Advanced or Metastatic

**Supportive Care**

1. Bone Mets
2. Anemia
3. Thrombocytopenia

this list was last updated on: 4/28/26