



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**

- 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

Open 8/12/25

Biliary (K83.9)

1. First Line Therapy

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

- 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

Open 3/25/26

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**

- i. CLOSED: Phase 2 Dose Selection Arm A: BMS-986504 400mg + Nab-Pac + Gem
- ii. CLOSED: Phase 2 Dose Selection Arm B: BMS-986504 600mg + Nab-Pac + Gem
- iii. CLOSED: Phase 2 Dose Selection Arm C: BMS-986504 400mg Placebo + Nab-Pac + Gem
- iv. CLOSED: 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

On Hold 5/5/26

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) Punitamig in Combination with Chemotherapy Versus Bevacizumab in Combi**

- i. Closed-Phase 2 Arm A1: punitamig (BMS-986545) Dose Level 1, Day 1 + FOLFOX or FOLFIRI,Q2W
- ii. Closed-Phase 2 Arm A2: punitamig (BMS-986545) Dose Level 2, Day 1 + FOLFOX or FOLFIRI,Q2W
- iii. Closed-Phase 2 Arm B:bevacizumab, Day 1 + FOLFOX or FOLFIRI, Q2W
- iv. OPEN Phase 3 Arm C: punitamig RP3D, Day 1 + FOLFOX or FOLFIRI, Q2W; or punitamig RP3D + CAPOX, Q3W
- v. OPEN Phase 3 Arm D:bevacizumab, Day 1 + FOLFOX or FOLFIRI, Q2W or bevacizumab + CAPOX, Q3W

b. **AbbV M24-533 (AndroMETA-CRC-533) A Phase 2, Open-Label, Randomized, Master Protocol Study to Evaluate S**

- i. Stage 2 - SubStudy 1 Arm A: Folfox+Bevacizumab+Telisotuzumab Adizutecan High dose (2.4mg/kg)
- ii. Stage 2 - SubStudy 1 Arm B: Folfox+Bevacizumab+Telisotuzumab adizutecan Low dose (2.0mg/kg)
- iii. Stage 2 - SubStudy 1 Arm C: SOC: Folfox+Bevacizumab
- iv. Stage 2 - SubStudy 2 Arm A: 5FU/leucovorin+Panitumumab+Telisotuzumab Adizutecan High dose (2.4mg/kg)
- v. Stage 2 - SubStudy 2 Arm B: 5FU/leucovorin+Panitumumab+Telisotuzumab Adizutecan Low dose (2.0mg/kg)
- vi. Stage 2 - SubStudy 2 Arm C: FOLFOX+Panitumumab

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

Open 7/28/26

Open 8/27/26

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**

- 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 III, Locally Advanced, Nonresectable

a. **BMS CA266-0001 (ROSETTA Lung-201) Study of Punitamig Vs Durvalumab Following Concurrent Chemoradiation T**

- i. Arm A: Punitamig 1200 mg (if body weight < 50 kg) or 1500 mg (if body weight ≥ 50 kg) 60-minute IV Q3W
- ii. Arm B: Durvalumab 1500 mg 60-minute IV Q4W

5. Stage IIB-IIIB Resected and Chemotherapy-Naive Stage IV

6. Stage IIIB, IIIC (not amenable to curative surgery or radiotherapy), or Stage IV

a. **BNT327-06 (ROSETTA LUNG-02) Phase III, multisite, randomized master protocol for a global trial of**

- i. Phase 3 Substudy A Arm 3: Non-squamous NSCLC punitamig+carbo+pemetrexed for 4 cycles followed by punitamig+pemetrexed
- ii. Phase 3 Substudy A Arm 4: Non-squamous NSCLC Pembro+carbo+pemetrexed for 4 cycles followed by pembro+pemetrexed
- iii. Phase 3 Substudy B Arm 3: Squamous NSCLC punitamig+carbo+paclitaxel for 4 cycles followed by punitamig maintenance
- iv. Phase 3 Substudy B Arm 4: Squamous NSCLC Pembro+carbo+paclitaxel for 4 cycles followed by pembro maintenance

7. Stage IV, Metastatic

8. First Line- Metastatic

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

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

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

- i. Arm A: Punitamig 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)

c. **BMS CA239-0004 (KRYSTAL-4) Adagrasib plus Pembrolizumab plus Chemotherapy vs. Placebo plus Pembro**

- i. Arm A: Adagrasib 400mg BID + pemetrexed 500mg/m2 Q3W (until PD or unacceptable toxicity) + [cisplatin 75mg/m2 OR carboplatin AUC 5mg/mL/min Q3W (4 cycles)] + pembrolizumab 200mg Q3W (up to 24 months)
- ii. Arm B: Placebo + pemetrexed 500 mg/m2 Q3W (until PD or unacceptable toxicity) + [cisplatin 75 mg/m2 OR carboplatin AUC 5 mg/mL/min Q3W (4 cycles)] + pembrolizumab 200 mg Q3W (up to 24 months)

9. 2nd or 3rd Line

10. Metastatic, Previously Treated

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

- 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

Open 1/7/26

Open 4/15/26

Open 5/19/26

Pending 8/11/26

Open 7/22/26

Open 3/19/26

Open 10/20/25

b. AZ D763QC00001 (TROPION-Lung17) , Datopotamab Deruxtecan or Docetaxel in Previously Treated TROP2-posit	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	
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	
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/m2 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 W045654 (INAVO123). A PHASE III, MULTICENTER, RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED	
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	
2. Metastatic, First Line	
a. AZ D8532C00008 (SERAFA-1). Phase IIIb study to evaluate Camizestrant plus Ribociclib in ER-positi	
i. Camizestrant 75 mg PO QD for 28 days + Ribociclib 600 mg PO QD for 21 days of a 28 day cycle	
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	
a. MK 6482-034 A Phase 3, Randomized, Double-blind, Study of Belzutifan + Zanzalintin	
i. Arm 1: Belzutifan 120 mg qd+ Zanzalintinib 60 mg qd	
ii. Arm 2: Belzutifan 120 mg qd+Placebo	
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	
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/m2, 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	
7. Relapsed and Refractory Follicular or Marginal Zone Lymphoma	
8. Diffuse Large Cell, First Line	
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)	

Open 11/13/25

Open 1/24/25

Open 1/31/23

Open 5/23/25

Open 9/2/26

Open 8/12/26

Accrual = 3

Open 8/13/25

Accrual Goal = 3

- 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: 9/2/26