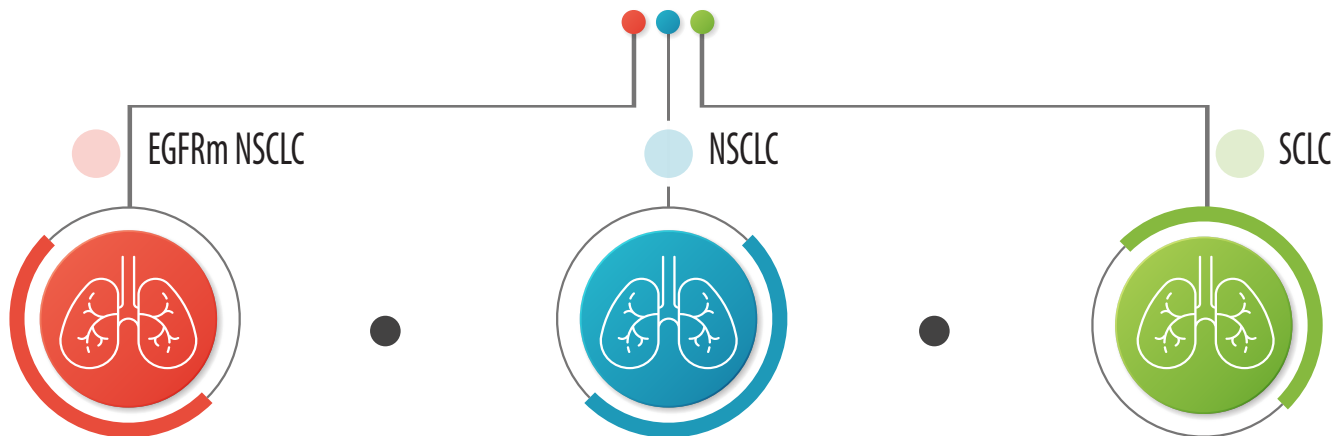


LUNG DATA REVIEW 2025

May 1, 2025



MARIPOSA: Head-to-head study demonstrated amivantamab + lazertinib benefits patients with previously untreated EGFR-mutant advanced NSCLC compared to single agent osimertinib and should be considered as a standard of care in the front line setting as a non-chemotherapy regimen

COCOON: Enhanced dermatologic management (COCOON DM) reduced dermatologic side effects compared to standard care in EGFR-mutant non-small cell lung cancer patients treated with amivantamab plus lazertinib, allowing more patients to remain on treatment

SKIPPirr: Incorporating prophylactic dexamethasone (8 mg, 5 doses) into treatment with amivantamab plus lazertinib can effectively prevent and reduce infusion related reactions

PALOMA-3 and -2: Incorporating prophylactic anticoagulation into treatment with amivantamab and lazertinib reduced venous thromboembolic events (VTEs)
More to come from **COPERNICUS**

TROPION-Lung05 / 01 Pooled Analysis: Dato-DXd has the potential to provide an additional treatment option for patients with previously treated EGFR-mutated NSCLC on tumor progression
Yet to be approved...
VENTANA TROP2 RxDx Device received breakthrough designation April 29, 2025

WU-KONG1: Sunvozertinib as a single oral drug has the potential to benefit patients with NSCLC and an EGFR exon20ins mutation who progressed on or after platinum-based chemotherapy
More to come...

eXALT3: The second-generation ALK inhibitor ensartinib represents an additional first-line treatment option for ALK-positive NSCLC patients... now FDA approved
Multiple options available... Optimal sequencing to be determined

eNRGy: Zenocutuzumab provides a treatment option for patients with NRG1+ NSCLC
Now FDA approved making it the first drug that targets tumors with the very rare NRG1 fusion genetic alteration

VISION: With long-term follow-up, tepotinib provides clinically meaningful benefit for patients with mNSCLC with METexon14 alterations, especially if treatment naïve

KRYSTAL-7: First-line adagrasib plus pembrolizumab has potential as a treatment option for patients with KRASG12C-mutated NSCLC and PD-L1 $\geq 50\%$ (KEYNOTE-024: 1L pembro alone, PFS 10.3 months). More to come... ongoing phase III trials...

BEAMION Lung-1: Zongertinib would be the first orally administered, targeted therapy for previously treated patients with HER2 (ERBB2)-mutant advanced non-small cell lung cancer (NSCLC) Not yet approved... PDUFA date is third quarter of 2025

KROCUS: Fulzerasib plus cetuximab provides benefit with an impressive objective response rate of 80% for patients with previously untreated advanced KRAS G12C mNSCLC and was superior to SoC established by the KEYNOTE-189 trial, which compared pemetrexed and platinum-based chemotherapy with or without pembrolizumab (Keytruda)... More to come...

LUNAR: Optune Lua is a portable medical device that could potentially offer a nontoxic supplemental treatment option for patients with metastatic NSCLC who have progressed on or after a platinum-based regimen

DeLLphi-300 / -301: Tarlatamab benefits patients with previously treated SCLC and could be considered a standard of care in the appropriate patient
Signal of intracranial activity... more to come
Monitoring and management of treatment related toxicities is key

ATLANTIS: Combination therapy with lurbinectedin plus doxorubicin did not improve overall survival versus control in patients with relapsed SCLC

LUPER: Small study of lurbinectedin plus pembrolizumab in patients with relapsed SCLC
More to come...

IMforte: Press release suggests statistically significant overall survival and progression-free survival results for combination of lurbinectedin and atezolizumab in first line maintenance therapy for ES-SCLC
More to come...

ADRIATIC: Now FDA approved...
Durvalumab consolidation therapy should be considered a new standard of care option for patients with LS-SCLC regardless of prior exposure to prophylactic cranial irradiation (PCI) or concurrent chemoradiotherapy (CRT) components

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