

Merck Pipeline

Q2 2024 Reflecting Pipeline to May 1, 2024

Lead-in language

The chart below reflects the company's research pipeline as of May 1, 2024. Candidates shown in Phase 3 include specific products and the date such candidate entered into Phase 3 development. Candidates shown in Phase 2 include the most advanced compound with a specific mechanism or, if listed compounds have the same mechanism, they are each currently intended for commercialization in a given therapeutic area. Small molecules and biologics are given MK-number designations and vaccine candidates are given V-number designations. Except as otherwise noted, candidates in Phase 1, additional indications in the same therapeutic area (other than with respect to cancer and certain other indications) and additional claims, line extensions or formulations for in-line products are not shown.

Merck pipeline as of May 1, 2024

Being developed in a collaboration.

Being developed in combination with Keytruda Being developed as monotherapy and/or in combination with Keytruda

Moved forward since last pipeline update.

Phase 2	Phase 2	Phase 2	Phase 2	Phase 2
Cancer Gastric Melanoma patritumab deruxtecan MK-1022 ¹	Cancer NSCLC quavonlimab MK-1308 ²	Cancer CRC quavonlimab + pembrolizumab MK-1308A	Thrombosis MK-2060	Heme zilovertamab vedotin MK-2140
Cancer CRC SCLC ifinatamab deruxtecan MK-2400 ¹	Cancer Neoplasm Malignant sacituzumab tirumotecan MK-2870 ^{1, 3}	Cancer Advanced solid tumors Prostate KEYTRUDA® MK-3475	Cancer Cutaneous Squamous Cell pembrolizumab + hyaluronidase subcutaneous MK-3475A	Cancer NSCLC favezelimab MK-4280 ²
Cancer Bladder Cutaneous Squamous Cell Endometrial Esophageal Melanoma RCC favezelimab + pembrolizumab MK-4280A	PH-COPD MK-5475	Cancer Neoplasm Malignant boserolimab MK-5890²	Cancer Ovarian raludotatug deruxtecan MK-5909¹	NASH efinopegdutide MK-6024



Merck pipeline as of May 1, 2024

- Being developed in a collaboration.
- Being developed in combination with Keytruda Being developed as monotherapy and/or in combination with Keytruda
- On FDA clinical hold
- On partial clinical hold for higher doses than those used in current clinical trials
- Phase 2b development costs are being co-funded

Moved forward since last pipeline update.

Phase 2	Phase 2	Phase 2	Phase 2	Phase 2
Vitiligo MK-6194	Cancer Endometrial Esophageal HCC Prostate Rare cancers WELIREG [™] MK-6482 ³	Cancer Advanced solid tumors LYNPARZA® MK-7339 ^{1,3}	Cancer Biliary Bladder Breast Cervical CRC Endometrial Esophageal Gastric HNSCC HCC Ovarian Prostate RCC vibostolimab + pembrolizumab MK-7684A	Cancer HNSCC LENVIMA ® MK-7902 1,2
Pulmonary Hypertension due to Left Heart Disease WINREVAIR™ MK-7962	Schizophrenia MK-8189 ⁶	HIV-1 prevention MK-8527	HIV-1 Infection islatravir+MK-8507 MK-8591B ⁴	HIV-1 Infection islatravir+lenacapavir MK-8591D ^{1,5}
Dengue fever virus Vaccine V181	Cancer Cutaneous Squamous Cell Carcinoma Bladder RCC V940 ^{1,2}			



Merck pipeline as of May 1, 2024

- Being developed in a collaboration.
- 2. Being developed in combination with Keytruda
- 3. Being developed as monotherapy and/or in combination with Keytruda
- 4. On partial clinical hold for higher doses than those used in current clinical trials
- 5. Available in the U.S. under Emergency Use Authorization

Moved forward since last pipeline update.

Phase 3	Phase 3	Phase 3	Phase 3
Hypercholesterolemia MK-0616	Cancer NSCLC patritumab deruxtecan MK-1022¹ (EU)	Cancer Heme nemtabrutinib MK-1026	Cancer RCC quavonlimab + pembrolizumab MK-1308A
Respiratory syncytial virus clesrovimab MK-1654	Cancer Breast Endometrial NSCLC sacituzumab tirumotecan MK-2870 ^{1,3}	Cancer Cutaneous Squamous Cell Carcinoma (EU) Hepatocellular (EU) Mesothelioma Ovarian SCLC KEYTRUDA® MK-3475	Cancer NSCLC pembrolizumab + hyaluronidase subcutaneous MK-3475A
Cancer Myeloproliferative Disorders bomedemstat MK-3543	Cancer CRC Heme favezelimab + pembrolizumab MK-4280A	Anti-Viral COVID-19 molnupiravir MK-4482^{1,5} (US)	Cancer Prostate opevesostat MK-5684 ¹
Ulcerative Colitis tulisokibart MK-7240	Cancer NSCLC SCLC LYNPARZA® MK-7339 ^{1, 2}	Cancer Melanoma NSCLC SCLC vibostolimab + pembrolizumab MK-7684A	Cancer Esophageal Gastric LENVIMA® MK-7902 ^{1,2}
HIV-1 infection doravirine + islatravir MK-8591A ⁴	Cancer Melanoma NSCLC V940^{1,2}		



Being developed in a collaboration In Dec 2023, FDA issued a CLR for the NDA for gefapixant. Merck is reviewing the feedback to determine next steps.

Moved forward since last pipeline update.

Merck pipeline as of May 1, 2024

New Molecular Entities Under Review	New Molecular Entities Under Review
Previously Treated Locally Advanced or Metastatic EGFR- Mutated NSCLC (HERTHENA-Lung01) patritumab deruxtecan MK-1022 (US)	von Hippel-Lindau (VHL) disease (LITESPARK-004) WELIREG® MK-6482 (EU)
Cough gefapixant MK-7264 (US ²)	Pulmonary Arterial Hypertension (STELLAR) WINREVAIR™ MK-7962 (EU)
Pneumococcal Vaccine Adult V116 (US, EU)	

Certain Supplemental Filings Under Review	Certain Supplemental Filings Under Review	Certain Supplemental Filings Under Review
1L Locally Advanced Unresectable or Metastatic Biliary Tract Cancer (KN966) KEYTRUDA® MK-3475 (JPN)	Primary Advanced or Recurrent Endometrial Carcinoma (KN868) KEYTRUDA® MK-3475 (US, EU JPN)	1L HER2 negative Locally Advanced Unresectable or Metastatic Gastric Cancer (KN859) KEYTRUDA® MK-3475 (JPN)
Resectable Stage II, IIIA or IIIB NSCLC (KN671) KEYTRUDA® MK-3475 (JPN)	1L Locally Advanced or Metastatic Urothelial Cancer (KNA39) KEYTRUDA® MK-3475 (EU, JPN)	High-Risk Locally Advanced Cervical Cancer (KNA18) KEYTRUDA® MK-3475 (EU, JPN)
Previously Treated Advanced Renal Cell Carcinoma (LITESPARK-005) WELIREG® MK-6482 (EU)		



. Being developed in a collaboration



Merck pipeline as of May 1, 2024

New Molecular EntitiesApprovals¹

Adults with Pulmonary Arterial
Hypertension WHO Group 1
(STELLAR)
WINREVAIR™
MK-7962
(US)

Certain Supplemental Approvals¹

Neoadjuvant/Adjuvant
Resectable NSCLC at High Risk
of Recurrence in Adults
(KN671)
KEYTRUDA®
MK-3475
(EU)

Forward-looking statement

This presentation of Merck & Co., Inc., Rahway, N.J., USA (the "company") includes "forward-looking statements" within the meaning of the safe harbor provisions of the U.S. Private Securities Litigation Reform Act of 1995. These statements are based upon the current beliefs and expectations of the company's management and are subject to significant risks and uncertainties. There can be no guarantees with respect to pipeline candidates that the candidates will receive the necessary regulatory approvals or that they will prove to be commercially successful. If underlying assumptions prove inaccurate or risks or uncertainties materialize, actual results may differ materially from those set forth in the forward-looking statements.

Risks and uncertainties include but are not limited to, general industry conditions and competition; general economic factors, including interest rate and currency exchange rate fluctuations; the impact of pharmaceutical industry regulation and health care legislation in the United States and internationally; global trends toward health care cost containment; technological advances, new products and patents attained by competitors; challenges inherent in new product development, including obtaining regulatory approval; the company's ability to accurately predict future market conditions; manufacturing difficulties or delays; financial instability of international economies and sovereign risk; dependence on the effectiveness of the company's patents and other protections for innovative products; and the exposure to litigation, including patent litigation, and/or regulatory actions.

The company undertakes no obligation to publicly update any forward-looking statement, whether as a result of new information, future events or otherwise. Additional factors that could cause results to differ materially from those described in the forward-looking statements can be found in the company's Annual Report on Form 10-K for the year ended December 31, 2023 and the company's other filings with the Securities and Exchange Commission (SEC) available at the SEC's Internet site (www.sec.gov).

No duty to update

The information contained in the presentation set forth below was current as of May 1, 2024. While this presentation remains on the company's website the company assumes no duty to update the information to reflect subsequent developments. Consequently, the company will not update the information contained in the presentation and investors should not rely upon the information as current or accurate after May 1, 2024.

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Candidates shown in Phase 3 include specific products. Candidates shown in Phase 2 include the most advanced compound with a specific mechanism in a given therapeutic area. Phase 1 candidates are not shown.