





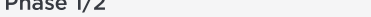



Takeda Oncology Pipeline

Our research and development efforts focus on advancing medicines for thoracic, gastrointestinal and hematologic cancers by leveraging modalities best suited to make a difference in the treatment of these diseases. Our core modalities include antibody drug conjugates (ADCs), complex biologics, small molecules and gamma delta T cell therapies.

These therapies in development are investigational and have not been approved by any regulatory bodies for use in the investigational indications listed below.

INVESTIGATIONAL THERAPY	MODALITY	INVESTIGATIONAL INDICATION	PHASE	ADDITIONAL INFORMATION
Brentuximab vedotin*	ADC	<i>Newly diagnosed Hodgkin lymphoma (EU)</i>	Phase 3 	Pfizer and Takeda fund joint development costs on a 50:50 basis, except in Japan where Takeda is solely responsible for development costs. Takeda has commercialization rights for the treatment outside of U.S. and Canada.
Cabozantinib*	Small molecule	<i>Metastatic castration-resistant prostate cancer (JP)</i>	Phase 3 	Tyrosine kinase inhibitor including MET/AXL/VEGFR Cabozantinib is being developed in Japan in collaboration with Exelixis.
Dazostinag (TAK-676)	Small molecule	<i>Solid tumors</i>	Phase 1/2 	STING agonist
Mirvetuximab soravtansine-gynx (TAK-853)	ADC	<i>Folate receptor alpha (FRα)-positive ovarian cancer (JP)</i>	Phase 1/2 	Takeda has development and commercialization rights for the treatment in Japan.
TAK-012	Gamma delta T cell therapy	<i>Acute myeloid leukemia</i>	Phase 1/2a 	
TAK-186	Complex Biologic	<i>Solid tumors</i>	Phase 1/2 	EGFR x CD3 targeting COBRA (COnditional Bispecific Redirected Activation) T cell engager immunotherapy
TAK-280	Complex Biologic	<i>Solid tumors</i>	Phase 1/2 	B7-H3 x CD3 targeting COBRA T cell engager immunotherapy
TAK-500	ADC	<i>Solid tumors</i>	Phase 1/2 	STING agonist immunostimulatory ADC (iADC)

*Marketed products have received approval in one or more jurisdictions.

All programs have global development rights unless otherwise noted.

GL = global (USA, Europe, Japan, China) / EU = Europe / JP = Japan / CN = China / GEM = global emerging markets

 @TakedaOncology

Learn more at [TakedaOncology.com](https://www.takedaoncology.com)

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ONCOLOGY