## **Takeda Oncology Pipeline**

Our research and development efforts focus on advancing medicines for hematologic, thoracic and gastrointestinal 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
Elritercept (TAK-226)	Complex biologic	Second-line anemia- associated myelodysplastic syndromes	Phase 3	Takeda has development and commercialization rights for the treatment worldwide outside of mainland China, Hong Kong and Macau.  Activin A/B ligand trap
		Anemia-associated myelofibrosis	Phase 2	
Mirvetuximab soravtansine-gynx (TAK-853)	ADC	Folate receptor alpha (FRq)- positive platinum-sensitive ovarian cancer (JP)	Phase 3	Takeda has development and commercialization rights for the treatment in Japan.
		Folate receptor alpha (FRa)- positive platinum-resistant ovarian cancer (JP)	Phase 1/2	
Rusfertide (TAK-121)	Small molecule	Polycythemia vera (USA)	Phase 3	Collaboration with Protagonist Therapeutics. Protagonist is responsible for development in the U.S. through the completion of the Phase 3 VERIFY trial. Takeda has rights for ex-U.S. development and is responsible for leading global regulatory and commercialization activities.  Hepcidin mimetic peptide (injection)
TAK-168	Small molecule	Solid tumors	Phase 1	Collaboration with Kumquat Biosciences. Kumquat is responsible for development (KQB168) through the completion of the Phase 1 trial (NCT06994806).

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.











