Takeda Oncology Pipeline

Beginning with patients' needs, we build our pipeline by identifying the most promising science and collaborate with leading teams around the globe to accelerate innovation. Our oncology R&D activities are focused on translating science into curative or transformative potential treatments by targeting tumor vulnerabilities and through novel strategies that leverage the power of innate immunity.

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	PLATFORM	INVESTIGATIONAL INDICATION	PHASE	ADDITIONAL INFORMATION
Cabozantinib*	Targeted therapy	Metastatic castration- resistant prostate cancer (JP)	Phase 3	Tyrosine kinase inhibitor including MET/AXL/VEGFR Cabozantinib is being developed in Japan in collaboration with Exelixis.
Fruquintinib*	Targeted therapy	Previously treated metastatic colorectal cancer (EU, JP)	Filed	Oral VEGFR 1/2/3 tyrosine kinase inhibitor Takeda has development and commercialization rights for the treatment outside of mainland China, Hong Kong and Macau.
lxazomib*	Proteasome inhibitor	Maint. newly diagnosed multiple myeloma post- stem cell transplant (US, EU)	Phase 3	Proteasome inhibitor
Ponatinib*	Targeted therapy	Newly diagnosed Ph+ acute lymphocytic leukemia (US)	Phase 3	BCR-ABL inhibitor Takeda shares development rights with Incyte Corp. (Europe, Turkey and Israel) and Otsuka Pharm. (Asia Pacific territories).
		Pediatric Ph+ acute lymphoblastic leukemia	Phase 1	
Relugolix* (TAK-385)	Small molecule	Prostate cancer (JP, Asian countries)	Phase 3	GnRH antagonist Relugolix is being developed in Japan and Asian countries in collaboration with Myovant Sciences, Inc.
Dazostinag (TAK-676)	Innate immunity enhancer	Solid tumors	Phase 1/2	STING agonist
Subasumstat (TAK-981)	Innate immunity enhancer	Multiple cancers	Phase 1/2	SUMOylation inhibitor

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TAK-007	Redirected immunity	B-cell malignancies	Phase 2	CD19 CAR NK TAK-007 is being developed in collaboration with The University of Texas MD Anderson Cancer Center.
TAK-012	Redirected immunity	Acute myeloid leukemia	Phase 1/2a	γδ (gamma delta) T cell therapy
TAK-186	Redirected immunity	Solid tumors	Phase 1/2	EGFR x CD3 targeting COBRA (COnditional Bispecific Redirected Activation) T cell engager immunotherapy
TAK-280	Redirected immunity	Solid tumors	Phase 1/2	B7-H3 x CD3 targeting COBRA (COnditional Bispecific Redirected Activation) T cell engager immunotherapy
TAK-500	Innate immunity enhancer	Solid tumors	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