

■ Oncology
 ■ Immunology
 ■ Neuroscience
 ■ Hematology
 ■ Ophthalmology
 ■ Other Diseases

Development Pipeline [Main table] (as of January 29, 2026)

Development code Origin	Generic name Product name	Indication # Additional indication (Combination drug)	Country/region	Projected submission	Mode of Action Modality (Dosage form)	Partner
Filed						
■ AF802/RG7853 in-house	alectinib Alecensa	ALK fusion / rearrangement gene-positive unresectable advanced or recurrent solid tumors #	Japan	June 2025	ALK inhibitor Small molecule (oral)	—
■ RG7446 Roche	atezolizumab Tecentriq	Adjuvant therapy for MRD- positive bladder cancer #	Japan	January 2026	Engineered anti-PD-L1 monoclonal antibody Antibody (IV)	—
■ RG7828 Roche	mosunetuzumab Lunsumio	Relapsed or refractory aggressive B-cell non- Hodgkin's lymphoma # (Polivy) #	Japan	May 2025	Anti-CD20/CD3 bispecific antibody Antibody (SC)	Roche
■ RG435 Roche	bevacizumab Avastin	Neurofibromatosis type 2 (NF2) #	Japan	August 2025	Anti-VEGF humanized monoclonal antibody Antibody (IV)	—
Phase III						
■ AF802/RG7853 in-house	alectinib Alecensa	Maintenance treatment of NSCLC (stage III) after chemoradiotherapy #	Global	—	ALK inhibitor Small molecule (oral)	Roche
■ RG7446 Roche	atezolizumab Tecentriq	HCC (intermediate stage) # (Avastin) #	Japan	2027	Engineered anti-PD-L1 monoclonal antibody Antibody (IV)	Roche
		HCC (2nd Line) # (lenvatinib or sorafenib)	Japan	—		Roche
■ RG6171 Roche	giredestrant —	Breast cancer (adjuvant)	Japan	2026	SERD (Selective Estrogen Receptor Degradar) Small molecule (Oral)	Roche
		Breast cancer (1st Line) (palbociclib)	Japan	2026		Roche
		Breast cancer (1st Line-3rd Line) (everolimus)	Japan	2026		Roche

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■ RG7828 Roche	mosunetuzumab Lunsumio	Follicular lymphoma (2nd Line) # (lenalidomide)	Japan	2026	Anti-CD20/CD3 bispecific antibody Antibody (IV)	Roche
		Previously untreated follicular lymphoma #	Japan	2028 and beyond		Roche
■ RG6026 Roche	glofitamab —	Previously untreated large B- cell lymphoma (Polivy)	Japan	2028 and beyond	Anti-CD20/CD3 bispecific antibody Antibody (IV)	Roche
■ RG6330 Roche	divarasib —	NSCLC (2nd Line)	Japan	2027	KRAS G12C inhibitor Small molecule (Oral)	Roche
		NSCLC (1st Line)	Japan	2028 and beyond		
■ RG7159 GlycArt Biotechnology	obinutuzumab Gazyva	Lupus nephritis #	Japan	2026	Glycoengineered type II anti-CD20 monoclonal Antibody Antibody (IV)	Nippon shinyaku
		Pediatric nephrotic syndrome #	Japan	2026		Nippon shinyaku
		Extra renal lupus #	Japan	2027		Nippon shinyaku
■ RG6299/ASO factor B Ionis Pharmaceuticals	sefaxersen —	IgA nephropathy	Japan	2028 and beyond	Antisense oligonucleotide targeting <i>complement</i> <i>factor B</i> mRNA Nucleic acid (SC)	Roche
■ RG6631 Roche	afimkibart —	Ulcerative colitis	Japan	2027	Anti-TL1A antibody Antibody (-)	Roche
		Crohn's disease	Japan	2028 and beyond		
■ — Bristol Myers Squibb	sparsentan —	IgA nephropathy	Japan	2026	Dual endothelin /angiotensin II receptor antagonist Small molecule (Oral)	—

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■ SA237/RG6168 in-house	satralizumab Enspryng	Myelin oligodendrocyte glycoprotein antibody- associated disease (MOGAD) #	Global	2026	pH-dependent binding humanized anti-IL-6 receptor monoclonal antibody Antibody (SC)	Roche
		Autoimmune encephalitis (AIE) #	Global	2027		Roche
■ RG6102 MorphoSys	trontinemab —	Alzheimer's disease	Japan	2028 and beyond	Anti-amyloid beta/TfR1 fusion protein Antibody (IV)	Roche
■ RG6356/SRP-9001 Sarepta	delandistrogene moxeparvovec Elevydis	Duchenne muscular dystrophy (DMD) (non-ambulatory) #	Japan	2028 and beyond	Microdystrophin gene therapy Gene therapy (IV)	Sarepta*
■ SKY59/RG6107 in-house	crovalimab PiaSky	Atypical hemolytic uremic syndrome (aHUS) #	Global	2026	Anti-C5 recycling antibody Antibody (SC)	Roche
■ ACE910/RG6013 In-house	emicizumab Hemlibra	Type 3 von Willebrand disease #	Global	2027	Anti-coagulation factor IXa/X humanized bispecific monoclonal antibody Antibody (SC)	Roche
■ SA237/RG6168 in-house	satralizumab Enspryng	Thyroid eye disease (TED) #	Global	2026	pH-dependent binding humanized anti-IL-6 receptor monoclonal antibody Antibody (SC)	Roche
■ RG6179 Roche	vamikibart —	Noninfectious uveitic macular edema (UME)	Japan	2026	Anti-IL-6 monoclonal antibody Antibody (vitreous injection)	Roche
■ RG7716 Roche	faricimab Vabysmo	Non-proliferative diabetic retinopathy (NPDR) #	Japan	2028 and beyond	Anti-VEGF/Anti-Ang-2 bispecific antibody Antibody (vitreous injection)	—

Development code Origin		Generic name Product name	Indication # Additional indication (Combination drug)	Country/region	Projected submission	Mode of Action Modality (Dosage form)	Partner
	RG6615 Alnylam Pharmaceuticals	zilebesiran —	Hypertension	Japan	2028 and beyond	RNAi therapeutic targeting angiotensinogen (AGT) RNAi (SC)	Alnylam Pharmace uticals
Phase II/III							
	GYM329/ RG6237 in-house	emugrobart —	Spinal muscular atrophy (SMA) (Evrysdi)	Global	2028 and beyond	Anti-latent myostatin sweeping antibody Antibody (SC)	Roche
Phase II							
	RG6026 Roche	glofitamab —	Relapsed or refractory diffuse large B-cell lymphoma	Japan	2026	Anti-CD20/CD3 bispecific antibody Antibody (IV)	—
			Relapsed or refractory mantle cell lymphoma	Japan	2028 and beyond		—
	GYM329/ RG6237 in-house	emugrobart —	Facioscapulohumeral muscular dystrophy (FSHD)	Global	2028 and beyond	Anti-latent myostatin sweeping antibody Antibody (SC)	Roche
	SA237/RG6168 in-house	satralizumab Enspryng	Duchenne muscular dystrophy (DMD) #	Global	2028 and beyond	pH-dependent binding humanized anti-IL-6 receptor monoclonal antibody Antibody (SC)	Roche
	RG6042 Ionis Pharmaceuticals	tominersen —	Huntington's disease	Japan	—	Antisense oligonucleotide targeting <i>HTT</i> mRNA Nucleic acid (IV)	Roche
	GYM329/RG6237 In-house	emugrobart —	Obesity	Global	2028 and beyond	Anti-latent myostatin sweeping antibody Antibody (SC)	Roche
Phase I/II							

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	RG6114 Roche	inavolisib —	PIK3CA-mutated breast cancer (palbociclib + fulvestrant)	Japan	—	PI3Kα inhibitor Small molecule (Oral)	Roche
	NXT007/ RG6512 in-house	— —	Hemophilia A	Global	2028 and beyond	Anti-coagulation factor IXa/X bispecific antibody Antibody (SC)	Roche
	RG6321 Roche	ranibizumab (Port Delivery Platform with ranibizumab) —	Neovascular age-related macular degeneration	Japan	2026	Humanized anti-VEGF monoclonal antibody Fragment Fab	—
			Diabetic macular edema	Japan	2026	Antibody (injection via implant)	—
Phase I							
	GC33 in-house	codrituzumab —	HCC	Global	—	Anti-Glypican-3 humanized monoclonal antibody Antibody (IV)	—
	ALPS12 in-house	clesitamig —	Solid tumors	Global	—	Anti-DLL3/CD3/CD137 trispecific antibody Antibody (IV)	—
	ROSE12 in-house	— —	Solid tumors	Global	—	Anti-CTLA-4 Switch antibody Antibody (IV)	—
	MINT91 in-house	— —	Solid tumors	Global	—	- Small molecule (Oral)	—
	AUBE00 In-house	— —	Solid tumors	Global	—	Pan-KRAS inhibitor Macrocyclic peptide (Oral)	—
	RG7421 Exelixis	cobimetinib —	Solid tumors	Japan	—	MEK inhibitor Small molecule (Oral)	—
	RG6160 Roche	cevostamab —	Relapsed or refractory multiple myeloma	Japan	—	Anti-FcRH5/CD3 bispecific antibody Antibody (IV)	—

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■ DONQ52 in-house	— —	Celiac disease	Global	—	Anti-HLA-DQ2.5/gluten peptides multispecific antibody Antibody (SC)	—
■ RAY121 in-house	— —	Autoimmune disease	Global	—	Anti-C1s recycling antibody Antibody (SC)	—
■ RG7935 Prothena	prasinezumab —	Parkinson's disease	Japan	—	Anti- α -synuclein monoclonal antibody Antibody (IV)	—
■ REVN24 in-house	— —	Acute diseases	Global	—	— Small molecule (IV)	—
■ RAY121 in-house	— —	—	Global	—	Anti-C1s recycling antibody Antibody (-)	—
Development Discontinued						
Development code Origin	Generic name Product name	Indication # Additional indication (Combination drug)	Country/region	Development stage	Mode of Action Modality (Dosage form)	Partner
■ RG7446 Roche	atezolizumab Tecentriq	NSCLC (perioperative) #	Japan	Phase III	Engineered anti-PD-L1 monoclonal antibody Antibody (IV)	Roche
■ BRY10 In-house	— —	Chronic diseases	Global	Phase I	— Antibody (SC)	—

In principle, completion of first dose is regarded as pipeline entry into each phase of clinical studies.

* Sarepta manages the global study including Japan

Changes from the last announcement on October 24, 2025

Oncology

- RG7446 Filed (Unresectable thymic carcinoma) → Approved
- RG7446 Phase III (Muscle-invasive bladder cancer (adjuvant))
→ Filed (Adjuvant therapy for MRD (molecular residual disease)-positive bladder cancer)
- RG7446 Phase III (NSCLC (perioperative)) → Development discontinued
- RG6330 Phase I/II (NSCLC (1st Line)) → Phase III

Immunology

- sparsentan Domestic Phase III (IgA nephropathy: development started)

Neuroscience

- RG6102 Phase I/II (Alzheimer's disease) → Phase III

Other Diseases

- RG6615 Phase I/II (Hypertension) → Phase III
- BRY10 Phase I (Chronic diseases) → Development discontinued

R&D Activities

For the changes during the FY2025 (January 1 – December 31), please refer to page 5 of “CONSOLIDATED FINANCIAL STATEMENTS (IFRS) (Non-Audited) (for the fiscal year 2025).”

Changes from January 1, 2026 to January 29, 2026 are as follows:

Oncology

- We filed for an antineoplastic agent / humanized anti-PD-L1 monoclonal antibody RG7446 (Product name: Tecentriq) for an additional indication of adjuvant therapy for MRD-positive bladder cancer in January 2026.
- We started global Phase III study for a KRAS G12C inhibitor RG6330 for the treatment of NSCLC (1st Line) in January 2026.

Development Pipeline [Attached table] (Major Chugai originated developments licensed out to 3rd parties excluding Roche)

Development code licensee/In-house	Generic name Product name	Indication # Additional Indication (combination)	Stage Country/region	Mode of Action Modality (Dosage form)	Licensee (Granted rights)
VS-6766/CKI27	avutometinib AVMAPKI	KRAS-mutated recurrent LGSOC (defactinib)	Phase III Overseas, U.S.	RAF/MEK clamp Small molecule (Oral)	Verastem Oncology (exclusive global license for the manufacturing, development and marketing)
			Phase II Japan		
		mPDAC (1st line) (defactinib+chemotherapy)	Phase I/II U.S.		
— /CIM331	nemolizumab NEMLUVIO	Chronic pruritus of unknown origin (CPUO)	Phase II Overseas	Anti-IL-31 receptor A humanized monoclonal antibody Antibody (SC)	Galderma (exclusive global license for the development and marketing excluding Japan)
LY3502970/OWL833	orforglipron —	Type 2 diabetes	Phase III Global	Oral non-peptidic GLP-1 receptor agonist Small molecule (Oral)	Eli Lilly and Company (worldwide development and commercialization rights)
		Obesity	Phase III Global Filed U.S.		
		Obstructive sleep apnea	Phase III Global		
		Hypertension	Phase III Global		
		Osteoarthritis	Phase III Global		
		Stress urinary incontinence	Phase III Global		
		Investigation of the effect of orforglipron on the incidence of major adverse cardiovascular events*	Phase III Global		

		Peripheral arterial disease	Phase III Global		
AP306/EOS789	— —	Hyperphosphatemia	Phase II China	Oral inhibitor of phosphate transporters Small molecule (Oral)	Alebund (exclusive global license for the manufacturing, development and marketing)

*In Participants with established atherosclerotic cardiovascular disease and/or chronic kidney disease

Progress made in R&D activities of major Chugai originated developments licensed out to 3rd party excluding Roche during the period from January 1, 2025 to January 29, 2026 was as follows.

- In Europe, Galderma obtained regulatory approval for the anti-IL-31 receptor A humanized monoclonal antibody CIM331 (Product name in Europe: NEMLUVIO (nemolizumab)) for moderate-to-severe atopic dermatitis and prurigo nodularis in February 2025. It started a global Phase II study for the treatment of chronic pruritus of unknown origin (CPUO) in Q4 2025.
- In the U.S., Verastem obtained regulatory approval under the accelerated approval pathway for the RAF/MEK clamp VS-6766/CKI27 (Product name in the U.S.: AVMAPKI) for *KRAS*-mutated recurrent LGSOC (combination with defactinib) in May 2025. It decided to discontinue the development for advanced *KRAS G12C*-mutated non-small cell lung cancer in December 2025.
- Eli Lilly and Company started a global Phase III study for the oral non-peptidic GLP-1 receptor agonist LY3502970/OWL833 (orforglipron) for the treatment of hypertension in Q3 2025, for the treatment of osteoarthritis, stress urinary incontinence, Investigation of the effect of orforglipron on the incidence of major adverse cardiovascular events in Q4 2025, and for the treatment of peripheral arterial disease in Q1 2026, respectively. In addition, it has filed for approval to the U.S. Food and Drug Administration for the treatment of obesity in Q4 2025.
- PI3K Class I inhibitor "PA799" was granted exclusive worldwide rights for manufacturing, development, and commercialization to Menarini Group in November 2016, but all licensing rights were returned to Chugai in June 2025.

Response to Requests from the MHLW Review Committee on Unapproved Drugs and Indications with High Medical Needs (As of January 29, 2026)

Development Request	Product	Indication	Development Status
Fourth development request	Xeloda*	Neuroendocrine tumor	Submitted company opinion and waiting for evaluation by committee
	Avastin	Cerebral edema induced by radiation necrosis	Submitted company opinion and waiting for evaluation by committee
	Mircera	Anemia associated with chronic kidney disease (CKD) in pediatric patients 3 months of age and older	Submitted company opinion and waiting for evaluation by committee

*Transferred the marketing authorization holder to CHEPLAPHARM K.K. as of February 1, 2024

Major Clinical Trials

Project	Expected indication	Study design	Study name	Stage	CT information
Oncology					
AF802/RG7853 (Alecensa)	Maintenance treatment of NSCLC (stage III) after chemoradiotherapy	ALK fusion-positive: Alecensa vs. durvalumab	HORIZON01	Phase III	NCT05170204
RG7446 (Tecentriq)	HCC (intermediate stage)	Tecentriq + Avastin + TACE vs. TACE	TALENTACE	Phase III	NCT04712643
	HCC [2nd line]	Tecentriq + lenvatinib or sorafenib vs. lenvatinib or sorafenib	IMbrave251	Phase III	NCT04770896
RG6171/SERD (giredestrant)	Breast cancer (adjuvant)	HR positive: RG6171 vs. endocrine therapy	lidERA	Phase III	NCT04961996
	Breast cancer [1st line]	HR positive: RG6171 + palbociclib vs. letrozole + palbociclib	persevERA	Phase III	NCT04546009
	Breast cancer [1st line-3rd line]	HR positive: RG6171 + everolimus vs. endocrine therapy+ everolimus	evERA	Phase III	NCT05306340
RG7828 (Lunsumio)	Follicular lymphoma [2nd line]	Lunsumio + lenalidomide vs. Rituxan + lenalidomide	CELESTIMO	Phase III	NCT04712097
	Relapsed or refractory aggressive B-cell non-Hodgkin's lymphoma	Lunsumio + Polivy vs. Rituxan + chemotherapy	SUNMO	Phase III	NCT05171647
	Previously untreated follicular lymphoma	Lunsumio + lenalidomide vs. Rituxan + chemotherapy	—	Phase III (domestic)	JRCT2011240017
RG6026 (glofitamab)	Previously untreated large B-cell lymphoma	RG6026 + Polivy + Rituxan + chemotherapy vs. Polivy + Rituxan + chemotherapy	SKYGLO	Phase III	NCT06047080
	Relapsed or refractory diffuse large B-cell lymphoma	RG6026 + gemcitabine + oxaliplatin (GemOx) (single arm)	—	Phase II (domestic)	JRCT2051250036
	Relapsed or refractory mantle cell lymphoma	RG6026 (single arm)			

Project	Expected indication	Study design	Study name	Stage	CT information
RG6330 (divarasib)	NSCLC [2nd line]	RG6330 vs. sotorasib or adagrasib	Krascendo 1	Phase III	NCT06497556
	NSCLC [1st line]	RG6330 + pembrolizumab vs. pembrolizumab +chemotherapy	Krascendo 2	Phase III	NCT06793215
RG6114 (inavolisib)	<i>PIK3CA</i> -mutated breast cancer	RG6114 + palbociclib + fulvestrant (single arm)	—	Phase I/II (domestic)	JRCT2031250161
Immunology					
RG7159 (Gazyva)	Lupus nephritis	Standard of care treatment ± Gazyva	—	Phase III (domestic)	JRCT2011210059
	Pediatric nephrotic syndrome	Gazyva vs. MMF	INShore	Phase III	NCT05627557
	Extra renal lupus	Gazyva vs. Placebo	—	Phase III (domestic)	JRCT2071230031
RG6299 (sefaxersen)	IgA nephropathy	RG6299 vs. Placebo	IMAGINATION	Phase III	NCT05797610
RG6631 (afimkibart)	Ulcerative colitis	RG6631 vs. Placebo	Ametrine-1	Phase III	NCT06589986
	Crohn's disease	RG6631 vs. Placebo	SIBERITE-1	Phase III	NCT06819878
sparsentan	IgA nephropathy	sparsentan (single arm)	—	Phase III (domestic)	JRCT2051240070
Neuroscience					
SA237/RG6168 (Enspryng)	Myelin oligodendrocyte glycoprotein antibody-associated disease (MOGAD)	Enspryng vs. Placebo	METEOROID	Phase III	NCT05271409
	Autoimmune encephalitis (AIE)	Enspryng vs. Placebo	CIELO	Phase III	NCT05503264
GYM329/RG6237 (emugrobarf)	Spinal muscular atrophy (SMA)	Evrysdi ± GYM329	MANATEE	Phase II/III	NCT05115110
RG6356/SRP-9001 (delandistrogene moxeparvovec)	Duchenne muscular dystrophy (DMD) (non-ambulatory)	RG6356 vs. Placebo	ENVISION	Phase III	NCT05881408
RG6102 (trontinemab)	Alzheimer's disease	RG6102 vs. Placebo	TRONTIER 1/ TRONTIER 2	Phase III	NCT07169578 NCT07170150
Hematology					

Project	Expected indication	Study design	Study name	Stage	CT information
SKY59/RG6107 (PiaSky)	Atypical hemolytic uremic syndrome (aHUS)	PiaSky (single arm)	COMMUTE-a	Phase III	NCT04861259
		PiaSky (single arm)	COMMUTE-p	Phase III	NCT04958265
ACE910/RG6013 (Hemlibra)	Type 3 von Willebrand disease	Hemlibra vs. On-demand therapy (standard of care treatment)	WILL-EMI	Phase III	NCT06998524
Ophthalmology					
SA237/RG6168 (Enspryng)	Thyroid eye disease (TED)	Enspryng vs. Placebo	SatraGO-1/ SatraGO-2	Phase III	NCT05987423 NCT06106828
RG6179 (vamikibart)	Noninfectious uveitic macular edema	RG6179 vs. sham	Sandcat	Phase III	NCT05642325
RG7716 (Vabysmo)	Non-proliferative diabetic retinopathy	RG7716 vs. sham	AZUSA	Phase III (domestic)	jRCT2071250009
RG6321 (ranibizumab (Port Delivery Platform with ranibizumab))	Neovascular age-related macular degeneration / Diabetic macular edema	RG6321 (single arm)	TEIEN	Phase I/II (domestic)	jRCT2071210073
Other diseases					
RG6615 (zilebesiran)	Hypertension	RG6615 vs. Placebo	ZENITH	Phase III	NCT07181109

Clinical Trials of In-House Developed Projects

Excluding projects listed in Major Clinical Trials among the projects listed in the development pipeline. Only clinical trials led by Chugai or Roche are listed.

Project	Expected indication	Stage	Enrollment* as of December 31, 2025	Study start	CT information
Oncology					
GC33	HCC	Phase I	27	November, 2008	NCT00746317
		Phase I	42	October, 2009	NCT00976170
		Phase I (domestic)	18	October, 2010	jRCT2080221218
		Phase II	185	May, 2012	NCT01507168
		Phase I	27	August, 2016	jRCT2080223270

Project	Expected indication	Stage	Enrollment* as of December 31, 2025	Study start	CT information
ALPS12 (clesitamig)	Solid tumors	Phase I	122	October, 2025	NCT07107490
ROSE12	Solid tumors	Phase Ia/Ib	219	June, 2023	NCT05907980
MINT91	Solid tumors	Phase I	122	April, 2025	jRCT2031240713
AUBE00	Solid tumors	Phase I	100	June, 2025	jRCT2031250094
Immunology					
DONQ52	Celiac disease	Phase Ia/Ib	56	September, 2022	NCT05425446
		Phase Ic	63	July, 2024	ACTRN12624000316505
RAY121	Autoimmune disease	Phase Ib	144	August, 2024	NCT06723106
Neuroscience					
GYM329/RG6237 (emugrobart)	Facioscapulohumeral muscular dystrophy (FSHD)	Phase II	51	March, 2023	NCT05548556
SA237/RG6168 (Enspryng)	Duchenne muscular dystrophy (DMD)	Phase II	50	April, 2025	NCT06450639
Hematology					
NXT007/RG6512	Hemophilia A	Phase I/II (domestic)	124	August, 2019	jRCT2080224835
		Phase I (domestic) (only healthy adults)	30	May, 2022	jRCT2031220050
		Phase I/II	60	October, 2023	NCT05987449
Other diseases					
REVN24	Acute diseases	Phase I (domestic) (only healthy adults)	210	October, 2023	jRCT2071230074
RAY121	—	Phase I (only healthy adults)	36	March, 2025	2024-515151-38-00
GYM329/RG6237 (emugrobart)	Obesity	Phase II	285	May, 2025	NCT06965413

* The number of enrollments is listed based on public information and generally refers to estimations or actual results.

FoundationOne CDx Cancer Genomic Profile: companion diagnostic indications (as of January 29, 2026)

Alterations	Cancer type	Relevant drugs
Activating <i>EGFR</i> alterations	NSCLC	afatinib maleate, erlotinib, gefitinib, osimertinib mesilate, dacomitinib hydrate
<i>EGFR</i> exon 20 T790M alteration		osimertinib mesilate
<i>ALK</i> fusion genes		alectinib, crizotinib, ceritinib, brigatinib
<i>ROS1</i> fusion genes		entrectinib
<i>MET</i> exon 14 skipping alterations		capmatinib hydrochloride hydrate
<i>BRAF</i> V600E and V600K alterations	Malignant melanoma	dabrafenib mesilate, trametinib dimethyl sulfoxide, vemurafenib, encorafenib, binimetinib
<u><i>BRAF</i> V600 alterations and <i>BRAF</i> fusion genes</u>	<u>Glioma</u>	<u>tovorafenib</u>
<i>ERBB2</i> copy number alterations (HER2 gene amplification positive)	Breast cancer	trastuzumab
<i>AKT1</i> alterations		capivasertib
<i>PIK3CA</i> alterations		
<i>PTEN</i> alterations		
<i>KRAS/NRAS</i> wildtype	Colorectal cancer	cetuximab (genetical recombination), panitumumab (genetical recombination)
Microsatellite instability-high		nivolumab (genetical recombination)
Microsatellite instability-high	Solid tumors	pembrolizumab (genetical recombination)
Tumor mutational burden-high		pembrolizumab (genetical recombination)
<i>NTRK1/2/3</i> fusion genes		entrectinib, larotrectinib sulfate, repotrectinib
<i>RET</i> fusion genes		seelpercatinib
<u><i>ALK</i> fusion genes</u>		<u>alectinib</u>
<i>BRCA1/2</i> alterations	Ovarian cancer	olaparib
<i>BRCA1/2</i> alterations	Prostate cancer	olaparib, talazoparib tosilate
<i>FGFR2</i> fusion genes	Biliary tract cancer	pemigatinib

* Underlined are the companion diagnostic features and relevant drugs currently under application for regulatory approval.

FoundationOne Liquid CDx Cancer Genomic Profile: companion diagnostic indications (as of January 29, 2026)

Alterations	Cancer type	Relevant drugs
Activating <i>EGFR</i> alterations	NSCLC	afatinib maleate, erlotinib, gefitinib, osimertinib mesilate
<i>EGFR</i> exon 20 T790M alteration		osimertinib mesilate
<i>ALK</i> fusion genes		alectinib, crizotinib, ceritinib
<i>ROS1</i> fusion genes		entrectinib
<i>MET</i> exon 14 skipping alterations		capmatinib hydrochloride hydrate
<i>NTRK1/2/3</i> fusion genes	Solid tumors	entrectinib
<i>BRCA1/2</i> alterations	Prostate cancer	olaparib