

Kyowa Kirin Co., Ltd.

Appendix to the Consolidated Financial Summary (IFRS) Fiscal 2025 Semi Annual

(January 1, 2025 - June 30, 2025)

⁻ These materials were made as a supplement to the Kessan Tanshin (Consolidated Financial Summary, IFRS), disclosed at the Tokyo Stock Exchange on July 31, 2025 for the first six months of Fiscal 2025, from January 1, 2025 to June 30, 2025.

⁻ This document is an English translation of the Japanese-language original.

⁻ The statements, including earnings forecasts, contained in these materials are based on the information currently available to the Company and on certain assumptions deemed to be reasonable by management. As such, they do not constitute guarantees by the Company of future performance. Actual results may differ materially from these projections for a wide variety of reasons.

⁻ Figures presented in these materials have been rounded to the nearest tenth.

⁻ Figures inside parenthesis presented in these materials indicate negative values.

⁻ Change amount in these materials presents change amount compared to the same period of the previous fiscal year.



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The average exchange rates for each period were as follows:

Unit: Yen

		FY 2024	4 results	FY 2025	ī results	FY2025 forecasts	
	Jan - Mar	Jan - Jun	Jan - Sep	Jan - Dec	Jan - Mar	Jan - Jun	Jan - Dec
USD	147	151	151	151	154	150	145
GBP	187	191	193	193	193	193	190
EUR	160	163	164	164	161	162	160

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I. Consolidated Financial Results

1. Trends in consolidated profit

<accumulative></accumulative>	The "★" symbol indicates financial KPIs (numerical guidance) that were set a FY 2024 results			et as targets in the	FY 2025		33 T Idil.	Unit: I	FY2021-2025		
	Jan - Mar	Jan - Jun	Jan - Sep	Jan - Dec	Jan - Mar	Jan - Jun	Change amount	Rate of change	Jan - Dec	Progress	Medium Term Business Plan Financial KPIs
Revenue	105.6	233.0	362.8	495.6	104.7	230.7	(2.3)	(1)%	478.0	48%	
★ CAGR (compared to FY 2020)	-	-	-	11.7%	-	-	-	-	8.5%	-	10% or higher
Cost of sales	(25.6)	(59.5)	(94.0)	(132.6)	(24.6)	(61.9)	(2.4)	4%	(126.0)	49%	
Gross profit	80.0	173.5	268.8	362.9	80.1	168.8	(4.7)	(3)%	352.0	48%	
Gross profit to revenue ratio	75.8%	74.5%	74.1%	73.2%	76.5%	73.2%	-	-	73.6%	-	
Selling, general and administrative expenses	(40.2)	(83.2)	(123.6)	(167.5)	(42.0)	(82.8)	0.4	(0)%	(166.0)	50%	
Research and development expenses	(23.3)	(49.2)	(74.3)	(103.5)	(28.6)	(52.5)	(3.3)	7%	(107.0)	49%	
★ R&D expense ratio	22.1%	21.1%	20.5%	20.9%	27.3%	22.8%	-	-	22.4%	-	Target of 18-20%
Share of profit (loss) of investments accounted for using equity method	0.9	3.1	3.5	3.5	(0.9)	1.5	(1.6)	(50)%	1.0	155%	
Core operating profit	17.4	44.1	74.4	95.4	8.6	35.0	(9.1)	(21)%	80.0	44%	
★ Core operating profit ratio	16.5%	18.9%	20.5%	19.3%	8.2%	15.2%	-	-	16.7%	-	25% or higher
Other income	2.6	4.4	13.3	13.1	0.4	0.6	(3.8)	(87)%			
Other expenses	(2.8)	(4.7)	(16.9)	(19.3)	(1.6)	(12.9)	(8.2)	176%			
Finance income (costs)	0.8	2.6	0.8	(5.8)	0.4	(0.7)	(3.4)	(128)%			
Profit before tax	18.1	46.5	71.6	83.5	7.9	22.0	(24.5)	(53)%	74.0	30%	
Income tax expense	(3.5)	(8.7)	(15.7)	(23.6)	(1.7)	(5.7)	3.1	(35)%	(17.0)	33%	
Ratio of income tax burden	19.2%	18.8%	21.9%	28.3%	21.5%	25.7%	-	-	23.0%	-	
Profit	14.6	37.8	55.9	59.9	6.2	16.3	(21.5)	(57)%	57.0	29%	
Profit to revenue ratio	13.9%	16.2%	15.4%	12.1%	5.9%	7.1%	-	-	11.9%	-	
EPS (¥/share)	27.26	70.76	105.20	113.06	11.78	31.18	(39.58)	-	108.91	-	
Core EPS (¥/share)*1	27.46	71.16	110.52	121.44	13.57	48.63	(22.53)	-	119.23	-	
Annual dividend (¥/share) ★ Dividend payout ratio (%) ^{*2}				58.00 47.76					60.00 50.32	-	Target of 40%
★ ROE (%)				7.10					6.60	-	10% or higher

^{*1} Core EPS is calculated as an indicator showing recurring profitability by dividing core profit (determined by subtracting "other income," "other expenses" and the related "income tax expense" from "profit") by the average number of shares during the period.

^{*2} Dividend payout ratio is shown based on core EPS.

<quarterly></quarterly>							Unit: E	Billions of yen
		FY 2024	results			FY 2025	5 results	
	Jan - Mar	Apr - Jun	Jul - Sep	Oct - Dec	Jan - Mar	Apr - Jun	Change amount	Rate of change
Revenue	105.6	127.4	129.8	132.8	104.7	125.9	(1.5)	(1)%
Cost of sales	(25.6)	(33.9)	(34.5)	(38.6)	(24.6)	(37.3)	(3.4)	10%
Gross profit	80.0	93.5	95.3	94.2	80.1	88.7	(4.9)	(5)%
Gross profit to revenue ratio	75.8%	73.4%	73.4%	70.9%	76.5%	70.4%	-	-
Selling, general and administrative expenses	(40.2)	(43.1)	(40.4)	(43.9)	(42.0)	(40.8)	2.3	(5)%
Research and development expenses	(23.3)	(25.9)	(25.0)	(29.3)	(28.6)	(23.9)	2.0	(8)%
★ R&D expense ratio	22.1%	20.4%	19.3%	22.0%	27.3%	19.0%	-	-
Share of profit (loss) of investments accounted for using equity method	0.9	2.2	0.4	0.0	(0.9)	2.5	0.3	12%
Core operating profit	17.4	26.7	30.3	21.0	8.6	26.4	(0.4)	(1)%
★ Core operating profit ratio	16.5%	21.0%	23.3%	15.8%	8.2%	21.0%	-	-
Other income	2.6	1.8	8.9	(0.2)	0.4	0.2	(1.6)	(90)%
Other expenses	(2.8)	(1.9)	(12.2)	(2.4)	(1.6)	(11.3)	(9.4)	497%
Finance income (costs)	0.8	1.8	(1.9)	(6.5)	0.4	(1.2)	(3.0)	(164)%
Profit before tax	18.1	28.4	25.1	11.9	7.9	14.1	(14.3)	(50)%
Income tax expense	(3.5)	(5.3)	(6.9)	(7.9)	(1.7)	(4.0)	1.3	(25)%
Profit	14.6	23.1	18.1	4.0	6.2	10.2	(13.0)	(56)%
Profit to revenue ratio	13.9%	18.2%	14.0%	3.0%	5.9%	8.1%	-	-



2. Revenue by regional control function

<Accumulative> Unit: Billions of yen FY 2024 results FY 2025 results FY 2025 forecasts Change amount Jan - Mar Jan - Jun Jan - Sep Jan - De Jan - Mar Jan - Jun Jan - Dec Progress 31.6 65.3 134.7 58.4 (6.9)48% Japan 98.0 27.2 121.8 32.3 North America 79.9 120.1 174 4 35.5 88.4 8.5 191.0 46% EMEA 16.7 36.9 65.7 84.9 19.7 37.0 0.1 73.7 50% 25.0 50.9 78.9 101.5 22.3 46.9 (4.0)91.5 51% Total consolidated revenue 105.6 233.0 362.8 495.6 104.7 230.7 (2.3)478.0 48% 34.3 68.6 100.9 141.2 28.3 63.1 (5.5) 130.0 48% Japan (location of customer) Overseas (location of customer) 71.3 164.4 261.9 354.4 76.4 167.6 3.2 348.0 48% Overseas ratio 68% 72% 72% 73% 73% 73% 71%

<Quarterly> Unit: Billions of yen

		FY 2024	4 results		FY 2025 results			
	Jan - Mar	Apr - Jun	Jul - Sep	Oct - Dec	Jan - Mar	Apr- Jun	Change amount	
Japan	31.6	33.7	32.8	36.6	27.2	31.2	(2.5)	
North America	32.3	47.6	40.2	54.3	35.5	52.8	5.2	
EMEA	16.7	20.1	28.8	19.2	19.7	17.3	(2.8)	
Others	25.0	26.0	28.0	22.6	22.3	24.5	(1.4)	
Total consolidated revenue	105.6	127.4	129.8	132.8	104.7	125.9	(1.5)	

3. Capital expenditures and intangible assets investment, depreciation and amortization

3. Capital experiolities and intangible assets investi	nent, depre	ciation and	arriortizatio	/11		Unit:	Billions of yen	
		FY 202	24 results		FY 202	FY 2025 results		
	Jan - Mar	Jan - Jun	Jan - Sep	Jan - Dec	Jan - Mar	Jan - Jun	Jan - Dec	
Capital expenditures (property, plant and equipment)	4.9	12.3	19.0	29.5	8.2	14.9	34.5	
Intangible assets investment	17.7	21.9	25.5	79.3	1.3	9.9	48.5	
Total	22.5	34.2	44.6	108.7	9.4	24.9	83.0	
Depreciation (property, plant and equipment)	3.7	7.4	11.0	14.8	3.8	7.6	16.5	
Amortization (intangible assets)	2.0	4.7	7.8	10.0	2.3	4.7	9.0	
Total	5.6	12.1	18.8	24.8	6.1	12.3	25.5	

^{*} Acquisitions of right-of-use assets are not included.

4. Number of employees by regional control function

		FY 20	24 results			FY 2025 results			
	As of March 31					As of June 30	Change amount		
Japan	4,090	4,103	4,093	4,020	March 31 3,898	3,890	(213)		
North America	648	664	664	668	709	678	14		
EMEA	543	548	553	547	538	540	(8)		
Others	873	866	877	434	375	381	(485)		
Total	6,154	6,181	6,186	5,669	5,520	5,489	(692)		

 $^{^{\}star}$ Others consists of number of employees of APAC subsidiaries, Orchard Therapeutics, etc.

II. Consolidated Statement of Cash Flows

ii. Consolidated Ctatement of Cash Flows						Unit:	Billions of yen
		FY 202	24 results		FY 2025 results		
	Jan - Mar	Jan - Jun	Jan - Sep	Jan - Dec	Jan - Mar	Jan - Jun	Change amount
Cash flows from operating activities	19.2	46.9	69.6	67.9	7.4	39.8	(7.0)
Cash flows from investing activities	(50.3)	(80.5)	(95.8)	(142.4)	(21.5)	(34.7)	45.8
Cash flows from financing activities	(41.3)	(63.2)	(83.7)	(84.7)	(16.5)	(17.2)	46.0
Effect of exchange rate changes on cash and cash equivalents	2.5	4.9	3.1	0.8	0.3	2.0	(2.9)
Net increase (decrease) in cash and cash equivalents	(70.0)	(91.9)	(106.8)	(158.4)	(30.3)	(10.1)	81.9
Cash and cash equivalents at beginning of period	403.1	403.1	403.1	403.1	244.7	244.7	(158.4)
Cash and cash equivalents at end of period	333.1	311.1	296.3	244.7	214.4	234.6	(76.5)

Revenue by regional control function is classified based on consolidated revenue from products of regional control functions in the One Kyowa Kirin (OKK) matrix global management structure, which

combines a regional organization, a functional organization, and a product organization (product franchises). * EMEA consists of Europe, the Middle East, Africa, etc.

[.] Others consists of revenue from technology out-licensing, revenue from APAC, hematopoietic stem cell gene therapy (revenue from Orchard Therapeutics), original equipment manufacturing, etc.

Revenue that was classified under "APAC" in 2024, will be included in "Others" starting from 2025 due to the business reoraganization in the APAC region.

Number of employees that were classified under "APAC" in 2024, will be included in "Others" starting from 2025 due to the business reoraganization in the APAC region.



III. Revenue from Main Products

<Accumulative> Unit: Billions of yen FY 2024 results FY 2025 results FY 2025 forecasts Change Jan - Jun Jan - Sep Jan - Dec Jan - Dec Progress Jan - Ma Jan - Jun 208.7 205.2 48% Products*1 93.4 327.9 446.8 91.8 (3.5)425.7 Crysvita 37.8 90.9 134.9 196.6 42.4 99.8 210.2 47% 8.9 Poteligeo 8.6 19.1 29.1 39.9 9.8 21.6 2.5 45.4 48% Libmeldy/Lenmeldy 1.4 2.1 4.4 3.0 6.9 64% 1.1 2.2 3.3 Nourianz 1.6 3.5 6.2 8.8 2.0 4.6 1.1 8.2 56% **PHOZEVEL** 0.6 1.7 2.9 3.7 2.0 8.9 41% 4.7 1.5 12.7 Duvroq 2.5 5.7 8.9 3.0 6.9 1.2 15.5 44% (0.5) 0.7 1.4 0.5 44% Nesp 2.0 2.6 0.9 2.0 Darbepoetin Alfa Injection Syringe [KKF] 2.8 5.6 8.5 11.6 2.3 4.9 (0.7)9.6 51% G-Lasta 5.8 10.5 15.3 20.5 4.3 9.1 (1.4)17.0 53% 50% 3.0 6.5 13.9 7.3 14.6 Romiplate 9.9 3.4 8.0 5.4 Orkedia 2.2 4.9 7.5 10.4 2.4 0.5 10.7 50% Rituximab BS [KHK] 1.9 3.8 5.7 7.8 1.5 3.2 (0.5)6.0 54% Nouriast 1.5 3.4 5.1 6.9 1.4 3.1 (0.3)6.5 47% HARUROPI 1.0 2.2 3.3 4.8 45% 4.6 1.0 2.1 (0.0)Dovobet *2 1.8 5.8 39 79 (7.9)34.9 Technology out-licensing*3 12.1 24.3 48.8 13.0 25.5 1.2 523 49% Benralizumab royalty*4 6.4 14.4 15.7 21.6 31.4 7.4 1.3 Total 105.6 233.0 362.8 495.6 104.7 230.7 (2.3)48%

^{*1} For revenue for Japan, the figures shown are those before the deduction of discounts and other items, and for revenue for overseas, the figures shown are those after the deduction of discounts and other items and include the impact of exchange rates.

^{*2} Due to the termination of the distribution and co-promotion agreement with LEO Pharma K.K. for Dovobet, sales by the Company ended on December 31, 2024.

^{*3} Revenue listed as "Technology out-licensing" represents the upfront income, milestone revenue and running royalty income that are obtained based on licensing agreements recognizing the granting to third parties the rights for development, manufacturing and sales of the Group's pipeline compounds or the use of technology, etc.

^{*4} Benralizumab royalty only refers to the royalty on sales of Fasenra by AstraZeneca (including the Company's own estimates).



III. Revenue from Main Products

<Quarterly> Unit: Billions of yen

Quarterly					U	IIIL. DIIIIOI	3 OI YEII	
	FY 2024 results							
	Jan - Mar	Apr - Jun	Jul - Sep	Oct - Dec	Jan - Mar	Apr - Jun	Change amount	
Products*1	93.4	115.3	119.2	118.9	91.8	113.4	(1.9)	
Crysvita	37.8	53.0	44.0	61.7	42.4	57.3	4.3	
Poteligeo	8.6	10.4	10.0	10.8	9.8	11.8	1.4	
Libmeldy/Lenmeldy	1.1	0.3	0.7	1.1	2.1	2.3	2.0	
Nourianz	1.6	2.0	2.6	2.6	2.0	2.6	0.6	
PHOZEVEL	0.6	1.1	1.2	1.8	1.5	2.1	1.1	
Duvroq	2.5	3.2	3.3	3.8	3.0	3.8	0.6	
Nesp	0.7	0.7	0.6	0.7	0.5	0.4	(0.3)	
Darbepoetin Alfa Injection Syringe [KKF]	2.8	2.8	2.9	3.1	2.3	2.6	(0.2)	
G-Lasta	5.8	4.7	4.8	5.2	4.3	4.8	0.1	
Romiplate	3.0	3.4	3.5	4.0	3.4	3.9	0.4	
Orkedia	2.2	2.7	2.5	3.0	2.4	2.9	0.2	
Rituximab BS [KHK]	1.9	1.9	1.9	2.1	1.5	1.7	(0.2)	
Nouriast	1.5	1.9	1.7	1.9	1.4	1.7	(0.2)	
HARUROPI	1.0	1.2	1.1	1.3	1.0	1.2	(0.0)	
Dovobet ^{*2}	1.8	2.1	1.8	2.1	1	-	(2.1)	
Technology out-licensing*3	12.1	12.2	10.7	13.8	13.0	12.5	0.4	
Benralizumab royalty*4	6.4	8.0	7.1	9.9	7.4	8.3	0.3	
Total	105.6	127.4	129.8	132.8	104.7	125.9	(1.5)	

^{*1} For revenue for Japan, the figures shown are those before the deduction of discounts and other items, and for revenue for overseas, the figures shown are those after the deduction of discounts and other items and include the impact of exchange rates.

^{*2} Due to the termination of the distribution and co-promotion agreement with LEO Pharma K.K. for Dovobet, sales by the Company ended on December 31, 2024.

 ² Due to the termination of the distribution and co-promotion agreement with LEO Pharma K.K. for Dovobet, sales by the Company ended on Decembe
 3 Revenue listed as "Technology out-licensing" represents the upfront income, milestone revenue and running royalty income that are obtained based on licensing agreements recognizing the granting to third parties the rights for development, manufacturing and sales of the Group's pipeline compounds or the use of technology, etc.
 4 Benralizumab royalty only refers to the royalty on sales of Fasenra by AstraZeneca (including the Company's own estimates).



III. Revenue from Main Products Revenue by location

<Accumulative> Unit: Billions of yen

			FY 202	24 results		F	Y 2025 resu	ılts	FY 2025 forecasts		
	Product name	Jan - Mar	Jan - Jun	Jan - Sep	Jan - Dec	Jan - Mar	Jan - Jun	Change amount	Jan - Dec	Progress	
Cry	vsvita	37.8	90.9	134.9	196.6	42.4	99.8	8.9	210.2	47%	
	Japan	2.5	5.4	8.2	11.7	2.8	6.1	0.7	13.1	47%	
	North America	22.8	58.7	87.2	130.0	24.1	64.0	5.3			
	[Millions of USD]	155	388	575	860	157	430	41			
	EMEA	11.9	25.4	37.1	51.5	14.8	28.0	2.5	197.1	48%	
	[Millions of GBP]	64	133	193	267	77	145	12			
	Others	0.6	1.3	2.3	3.3	0.8	1.7	0.4			
Pot	teligeo	8.6	19.0	29.0	39.9	9.8	21.6	2.6	45.4	48%	
	Japan	0.4	1.0	1.4	1.8	0.3	0.7	(0.3)	1.9	37%	
	North America	6.3	14.1	21.6	29.7	6.9	16.0	1.9	34.1	47%	
	[Millions of USD]	43	93	143	197	45	107	14	235	46%	
	EMEA	1.9	3.9	6.0	8.2	2.6	4.9	0.9	9.2	53%	
	[Millions of GBP]	10	21	31	43	13	25	5	48	52%	
	Others	0.0	0.1	0.1	0.1	0.0	0.1	(0.0)	0.3	23%	
Lib	meldy/Lenmeldy	1.1	1.4	2.2	3.3	2.1	4.4	3.0	6.9	64%	
	US	-	-	-	-	1.1	1.6	1.6			
	EMEA	1.1	1.4	2.2	3.3	1.0	2.8	1.4			

Unit: Billions of yen <Quarterly>

Draduat nama		FY 202	24 results		FY 2025 results				
Product name	Jan - Mar	Apr - Jun	Jul - Sep	Oct - Dec	Jan - Mar	Apr - Jun	Change amount		
Crysvita	37.8	53.0	44.0	61.7	42.4	57.3	4.3		
Japan	2.5	2.9	2.8	3.6	2.8	3.3	0.4		
North America	22.8	35.9	28.6	42.8	24.1	39.9	4.0		
[Millions of USD]	155	233	187	285	157	273	40		
EMEA	11.9	13.5	11.7	14.4	14.8	13.2	(0.3)		
[Millions of GBP]	64	70	59	74	77	68	(1)		
Others	0.6	0.7	0.9	1.0	0.8	1.0	0.2		
Poteligeo	8.6	10.4	10.0	*1 <u>10.9</u>	9.8	11.8	1.5		
Japan	0.4	0.5	0.4	0.5	0.3	0.4	(0.1)		
North America	6.3	7.8	7.5	8.1	6.9	9.1	1.3		
[Millions of USD]	43	51	49	54	45	62	11		
EMEA	1.9	2.0	2.0	2.3	2.6	2.3	0.3		
[Millions of GBP]	10	10	10	12	13	12	1		
Others	0.0	0.1	0.0	0.0	0.0	0.0	(0.0)		
Libmeldy/Lenmeldy	1.1	0.3	0.7	1.1	2.1	2.3	2.0		
US	-	-	-	-	1.1	0.5	0.5		
EMEA	1.1	0.3	0.7	1.1	1.0	1.8	1.5		

^{*1} The underlined portion indicates that the amount has been changed from the amount [10.8] presented in "Appendix to the Consolidated Financial Summary (IFRS) Fiscal 2025 First Quarter".

* Revenue is classified based on consolidated revenue from regional control functions.

^{*} The revenue, generated in various currencies inside each corporate region, is converted and aggregated in USD for North America and in GBP for EMEA.

^{*} Revenue that was classified under "APAC" in 2024, will be included in "Others" starting from 2025 due to the business reoraganization in the APAC region.



IV. R&D pipeline







Updated since Dec. 31, 2024 Updated since Mar. 31, 2025

As of Jun. 30, 2025

	Code Name				Stage		As of Jun. 30, 202
	Generic Name	Mechanism of Action	Indication	DL.		DLTTT	[In-House or Licensed] Remarks
¥	Formulation KK8123 Injection	Anti-FGF23 Fully Human Antibody	X-linked Hypophosphatemia	PhI	PhII	PhIII	[In-House] Clinical study is being conducted in NA and EU as a global product
*	KK8398 infigratinib Oral	FGFR 3 Inhibitor	Achondroplasia				[QED Therapeutics] Preparation underway for Ph III in JP
			Acute Myeloid Leukemia (AML) (Monotherapy)		-		[Kura Oncology] The detailed results presented at ASCO in June 2025 Adult Relapsed or Refractory AML with a NPM1 Mutation KOMET-001
			Acute Lymphoblastic Leukemia (ALL) (Monotherapy)				Clinical study is being conducted in NA and EU as a global product KMT2A-rearranged ALL KOMET-001
*	ziftomenib ※	Menin Inhibitor	Acute Myeloid Leukemia (AML) (Monotherapy)				Clinical study is being conducted in NA and EU as a global product Non-NPM1-mutant AML/Non-KMT2A-rearranged AML KOMET-001
	Oral	Henri Hinoto					Clinical study is being conducted in NA as a global product NPM1-mutant AML/KMT2A-rearranged AML Combinations with venetoclax + azacitidine, and cytarabine + daunorubicin KOMET-007
			Acute Myeloid Leukemia (AML) (Combination)				Clinical study is being conducted in NA and EU as a global product NPM1-mutant AML/KMT2A-rearranged AML Combinations with gilteritinib, FLAG-IDA, LDAC KOMET-008
							Preparation underway for Ph III as a global product NPM1-mutant AML/KMT2A-rearranged AML Combinations with venetoclax + azacitidine, and cytarabine + daunorubicin KOMET-017
¥	KK2845	Anti-TIM-3 ADC	Acute Myeloid Leukemia (AML)				[In-House] Antibody-Drug Conjugate Clinical study is being conducted in JP as a global product
8	OTL-203	Hematopoietic Stem Cell (HSC) Gene Therapy	MPS-IH (Hurler Syndrome)				[In-House] Rare Pediatric Disease (RPD) and Fast Track designations (FDA) Priority Medicines (PRIME) designation (EMA) Area of clinical study: NA and EU
8	OTL-201	Hematopoietic Stem Cell (HSC) Gene Therapy	MPS-IIIA (Sanfilippo Syndrome type A)		Ph I / Ph II		[In-House] Rare Pediatric Disease (RPD) designation (FDA) Preparation underway for registrational study (equivalent to PhⅢ study)
			Moderate to Severe Atopic Dermatitis				[In-House] POTELLIGENT Human monoclonal antibody production technology Collaboration agreement with Amgen for the development of rocatinlimab in all the countries exce for Japan Clinical study is being conducted in JP, NA, EU, UK,
¥	KHK4083/AMG 451 rocatinlimab Injection	Anti-OX40 Antibody					Middle East, Asia, Oceania, and other regions as a global product Clinical study is being conducted in JP, NA, EU, Asia,
			Prurigo Nodularis Moderate to Severe				and Oceania as a global product Clinical study is being conducted in JP, NA, EU, Asia,
	KIIKAOF4		Asthma Diabetic Macular Edema				and Oceania as a global product [In-House] Clinical study is being conducted in JP, NA, Asia, and
來	KHK4951 tivozanib Ophthalmic	VEGF Receptor Tyrosine Kinase Inhibitor	Neovascular Age-Related Macular Degeneration				Oceania as a global product Clinical study is being conducted in JP, NA, Asia, and Oceania as a global product
¥	KK2260 Injection	EGFR-TfR1Bispecific Antibody	Advanced or Metastatic Solid Tumors				[In-House] REGULGENT Fully human antibody production technology Clinical study is being conducted in JP, and a clinical study is prepared under way for PhI in NA as a global



IV. R&D pipeline

Code Name Generic Name Formulation		Mechanism of Action	Indication	Stage			[In-House or Licensed]				
				PhI	PhII	PhIII	Remarks				
¥	KK2269 Injection	EpCAM-CD40Bispecific Antibody	Advanced or Metastatic Solid Tumors				[In-House] REGULGENT Fully human antibody production technology Clinical study is being conducted in JP and NA as a global product				
W	KK4277 Injection	Anti-PTPRS Humanized Antibody	Systemic Lupus Erythematosus/Cutaneous Lupus Erythematosus				[SBI Biotech] POTELLIGENT Clinical study is being conducted in JP and Asia				
W	KK3910 Injection		Essential Hypertension	~			[In-House] Clinical study is being conducted in JP as a global product				

 $[\]begin{tabular}{ll} \hline \& For detailed information on ziftomenib's development status, please refer to Kura Oncology's website. https://kuraoncology.com/ \end{tabular}$

Major Applications and Approvals

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	Code Name, Generic Name, Product Name	Indication	Application/Under Review	Countries/Regions Received Approval in 2025				
	ziftomenib	Adult Relapsed or Refractory (R/R) Acute Myeloid Leukemia (AML) with a Nucelophosmin1 (NPM1) Mutation	US	-				