ADVANCING TO GLOBAL LAUNCHES & CONTINUED PIPELINE PROGRESS

CORPORATE PRESENTATION

September 2023

Nasdaq/AIM:HCM | HKEX:13





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A global science-focused biopharma



Fully integrated R&D and commercialization platform



Global novel drug discovery & manufacturing operations

20+ years novel drug discovery – **13 clinical-stage innovative NMEs**^[1] discovered in-house New flagship factory expected to come online in 2023/4 to expand capacity by 5x Listed on the LSE (HCM), NASDAQ (HCM), and HKEX (13)



- China, U.S., EU & Japan clinical capabilities
- First 3 novel oncology medicines approved



Commercial teams in China

- Oncology commercial team covering >3,000 hospitals in China
- Commercial partnering outside of China

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The path to a sustainable business...

HUTCHMED medium-term & long-term strategy

AMBITION

to mature into a profitable biopharma from an emerging growth co

VISION UNCHANGED:

discovering, developing & bringing new innovative medicines to patients worldwide

2022

2nd sNDA-enabling Phase III



Peak year for cash burn

Target 2025 to be self-sustaining

6-7 products potentially launched in China



Growth & operating leverage

HUTCHME

Blockbuster in-market sales
Accelerating China growth
Royalties from ex-China sales



2023 - 2024

Reduction in R&D costs

Continued revenue growth from new indications / regions

Global commercialization through partnerships

H1 2023: strong execution on strategic direction





STRATEGIC DELIVERY

- ✓ Delivering revenue growth & executing long-term growth strategy
- ✓ **Delivering global partnerships**, to reach more patients than ever: Takeda licensing closed
- ✓ **Delivering cash conservation &**cost optimization: \$856m cash
 & 20% lower R&D costs vs H1'22



PRODUCTS & PIPELINE PROGRESS



LATE STAGE

- ✓ Fruq US & EU NDA/MAA accepted (CRC)
- ✓ Fruq China sNDA accepted (GC)
- ✓ Savo & '453 initiated new registration studies (GC, IHCC)
- Savo SAVANNAH enrollment to finish H2 2023 (2L NSCLC)

2ND WAVE

- ✓ Sovle ESLIM Ph III readout Aug 2023 (ITP)
- Amdiz reg Ph II H2'23 readout (FL)
- Taz bridging study to fully enroll H2'23 (FL)



CHINA COMMERCIAL DELIVERY



- ✓ All 3 medicines on NRDL
- ✓ Oncology/Immunology prod. rev. +26% (+35% CER) on track to meet guidance
- ✓ Combined in-market sales +16% (+25% CER) for ELUNATE®, SULANDA® & ORPATHYS®

Agenda



1

Financial review & outlook

Underpinned by strong financial & strategic fundamentals

2

Commercial delivery

Novel oncology products continue to bring growth

3

Manufacturing update

New facility to enable more control and cost savings

4

Late-stage pipeline

15+ potential NDAs & sNDAs in the next 3 years

5

Our strategy

Revenue growth & strategic actions on path to profitability



Well-financed position

HUTCHMED

On path to sustainable business

Condensed Consolidated Balance Sheets

(in US\$ millions)	Jun 30, 2023	Dec 31, 2022
Assets		
Cash, cash equivalents & short-term investments	856.2	631.0
Accounts receivable	129.2	98.0
Other current assets	105.1	110.9
Property, plant and equipment	96.8	75.9
Investments in equity investees	37.7	73.8
Other non-current assets	72.5	39.8
Total assets	1,297.5	1,029.4
Liabilities and shareholders' equity		
Accounts payable	54.6	71.1
Other payables, accruals and advance receipts	227.2	264.6
Deferred revenue	149.5	13.5
Bank borrowings [1]	40.1	18.1
Other liabilities	26.1	25.2
Total liabilities	497.5	392.5
Company's shareholders' equity	782.0	610.4
Non-controlling interests	18.0	26.5
Total liabilities and shareholders' equity	1,297.5	1,029.4

As of June 30, 2023

Cash Resources

- \$856m cash / cash eq. / ST inv. [2]
- \$65m unutilized banking facilities

Borrowing

\$40m in bank borrowings

Others

\$44m additional cash at SHPL JV

Impact of Takeda transaction

- \$400m non-refundable upfront payment was received in April 2023
- \$259m recognized on *Consolidated Statement of Operations* as *Revenue*, and the remainder was recorded in *Deferred revenue* as of June 30, 2023





Oncology sales growth & Other Ventures income

Oncology consolidated revenues FY2023 guidance unchanged: \$450-\$550 million

(including recognizing ~\$280m of the Takeda upfront payment)

Condensed Consolidated Statements of Operations

(Unaudited, in US\$ millions, except share and per share data)	Six months end	ed June 30	
	2023	2022	
Revenues:			
Oncology/Immunology – Marketed Products	80.1	63.5	
Oncology/Immunology – R&D	279.1	27.6	
Oncology/Immunology consolidated revenues	359.2	91.1	
Other Ventures	173.7	110.9	
Total revenues	532.9	202.0	
Operating expenses:			
Costs of revenues	(208.3)	(137.3)	
R&D expenses	(144.6)	(181.7)	
Selling & general admin. expenses	(68.3)	(79.8)	
Total operating expenses	(421.2)	(398.8)	
	111.7	(196.8)	
Other income/(expense), net	25.4	(3.8)	
Income/(loss) before income taxes & equity in earnings			
of equity investees	137.1	(200.6)	
Income tax (expense)/benefit	(2.7)	4.2	
Equity in earnings of equity investees, net of tax	35.1	33.5	
Net income/(loss)	169.5	(162.9)	
Less: Net (income)/loss attrib. to non-controlling interests		0.0	
Net income/(loss) attributable to HUTCHMED	168.6	(162.9)	
Per ordinary share (basic)	0.20	(0.19)	
Per ordinary share (diluted)	0.19	(0.19,	
Per ADS share (basic)	1.00	(0.96)	
Per ADS share (diluted)	0.97	(0.96)	

Total Consolidated Revenues up 164% (173% CER) to \$533m

- Oncology product rev. up 26% (35% CER) to \$80m (H1'22: \$64m)
- R&D revenues of \$279m included recognition of \$259m in Takeda upfront payment (out of \$400m)

Control over operating expenses

- R&D: supporting 15+ registration enabling programs
 - As a result of strategic prioritization of our pipeline
 - Ex-China decreased to \$56m (H1'22: \$84m)
- SG&A expenses decline primarily reflected the restructuring of the ex-China commercial infrastructure in late 2022

Benefitting from our 50% share of SHPL JV's growth

- **Net revenues up 11% (19% CER)** to \$235m (H1'22: \$212m)
- Net income attributable to HUTCHMED from equity investees up 5% (12% CER) to \$35m (H1'22: \$34m)

RMB declined 7% in H1 2023 vs USD

Continuing growth of oncology product sales









(US\$ in millions)	H1 2023	H1 2022	%Δ (CER)	H1 2023	H1 2022	%Δ (CER)
	In-	market Sa	ales ^[1]	Consolidated Revenues [
ELUNATE® (fruquintinib)	\$56.3	\$50.4	+12% (+20%)	\$42.0	\$36.0	+16% (+25%)
SULANDA® (surufatinib)	\$22.6	\$13.6	+66% (+79%)	\$22.6	\$13.6	+66% (+79%)
ORPATHYS® (savolitinib)	\$22.0	\$23.3	-5% (+2%)	\$15.1	\$13.8	+10% (+17%)
TAZVERIK® (tazemetostat)	\$0.4	\$0.1	+560% (+583%)	\$0.4	\$0.1	+560% (+583%)
Product Sales ^[2]	\$101.3	\$87.4	+16% (+25%)	\$80.1	\$63.5	+26% (+35%)
Other R&D Service income				\$20.4	\$12.6	+62% (+66%)
Upfront & Milestone payment				\$258.7	\$15.0	
Total				\$359.2	\$91.1	+294%(+301%)

^[1] For ELUNATE® and ORPATHYS®, represents total sales to third parties as provided by Lilly and AstraZeneca, respectively; and their sales to other third parties as invoiced by HUTCHMED.

^[2] For ELUNATE®, represents manufacturing fees, commercial service fees and royalties paid by Lilly, to HUTCHMED, and sales to other third parties invoiced by HUTCHMED; for ORPATHYS® represents manufacturing fees and royalties paid by AstraZeneca and sales to other third parties invoiced by HUTCHMED; for SULANDA® and TAZVERIK®, represents the Company's sales of the products to third parties.

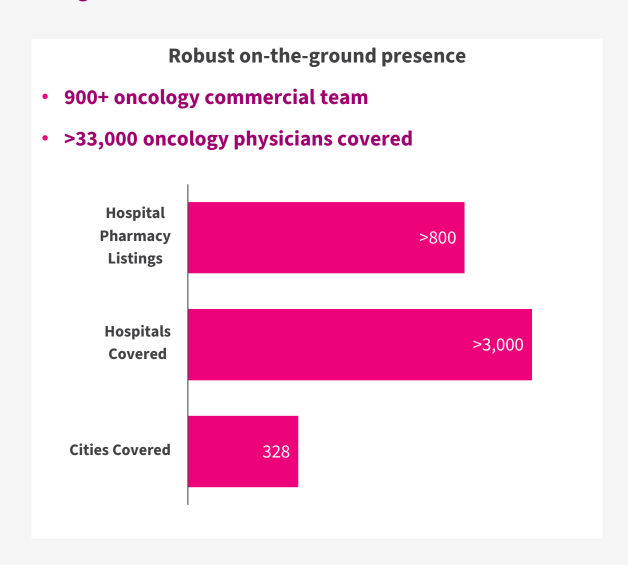
Commercial environment



Policies benefitting innovative medicines access + commercial organization at scale

Favorable operating environment

- China is the second largest country by pharmaceutical spending [1]
- Policies promoting access to innovative medicines
 - Life sciences is one of the key strategic priority sectors to provide a better quality of life
 - Simplification of NRDL renewals (Jul 2023)
 - CDE guideline to accelerate review of innovative marketing applications with breakthrough designations (Apr 2023)
 - Guidance on development of oncology medicines, encouraging innovations that have real clinical value (Dec 2021)
- Emergence of commercial health insurance

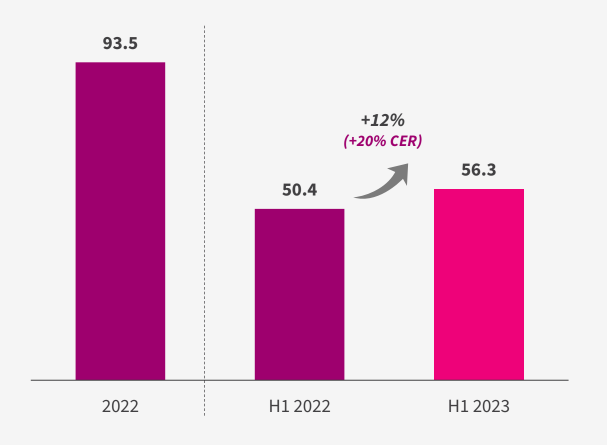


ELUNATE® (fruquintinib) remains market leader in 3L CRC





In-market sales (US\$ millions)



Underlying demand remains strong

- **~17,000 est. new patients** treated, up ~20% vs H1 2022
- COVID-related impact early in the year has resolved
- Improving access to diagnosis & treatment in Tier 3 & 4 cities

Strong competitive position

- Inclusion in CSCO & CACA CRC Guidelines^[1]
- Inclusion in Pan-Asian mCRC Clinical Practice Guidelines
- Maintaining leadership in patient share in 3L CRC

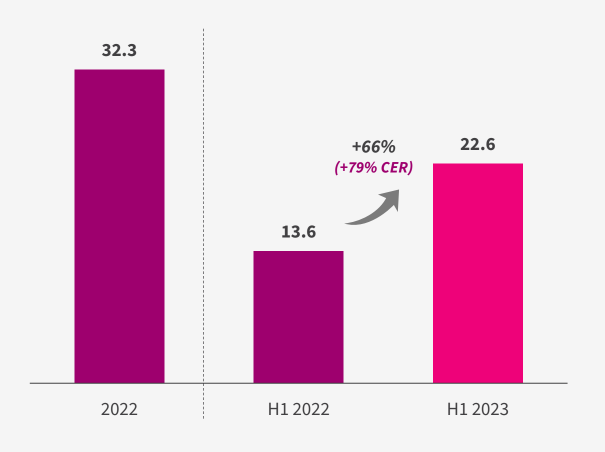
(IQVIA ¹³)	Q4-18	Q4-19	Q4-20	Q4-21	Q4-22	Q2-23
ELUNATE ®	2%	25%	33%	39%	44%	47%
STIVARGA®	29%	32%	35%	34%	29%	26%

SULANDA (surufatinib) increasing patient access & duration of treatment





In-market sales (US\$ millions)



Continued benefit of NRDL inclusion

- **~12,000 est. new patients** treated, vs ~7,500 in H1 2022
- Increase in duration of treatment
- Additional hospital listings (+19%)

Maintaining market share position

- Included in CSCO & CACA NENs Guidelines^[1] and China GEP NETs Expert Consensus
- Inclusion in guidelines outside of NENs/NETs
- Ranked the 2nd brand in NET market since Q3 2022,
 surpassed Sutent[®] & Afinitor[®] (IQVIA^[2])

Q1 2023	SANDOSTATIN®	SULANDA®	SUTENT®	AFINITOR®	Other
Pt share	36%	17%	13%	11%	23%

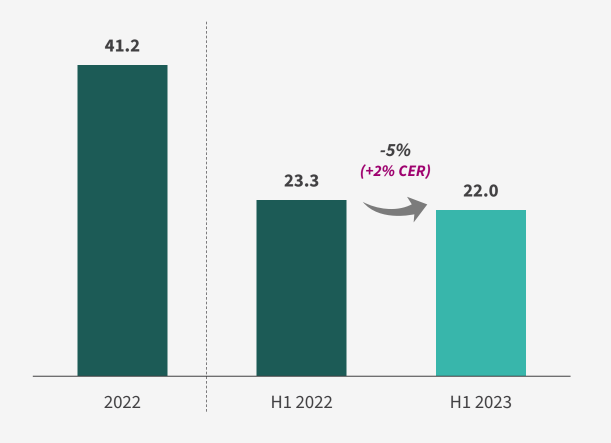
ORPATHYS® (savolitinib) first-in-class MET inhibitor



NRDL inclusion has expanded patient access significantly



In-market sales (US\$ millions)



NRDL inclusion from March 1, 2023

- Sales flat due to delayed start of NRDL plus a ~38% price reduction
- However, volume up +84% in Q2'23 vs. Q2'22 aided by NRDL inclusion
- Approved in Macau in March 2023

Inclusion in key treatment guidelines

- NHC, CSCO, CACA, CMA, CTONG [1]
- MET diagnostic testing is now recommended as SOC for late-stage NSCLC

AZ a strong China commercial partner

• Top lung cancer franchise synergies

New Shanghai manufacturing facility ready

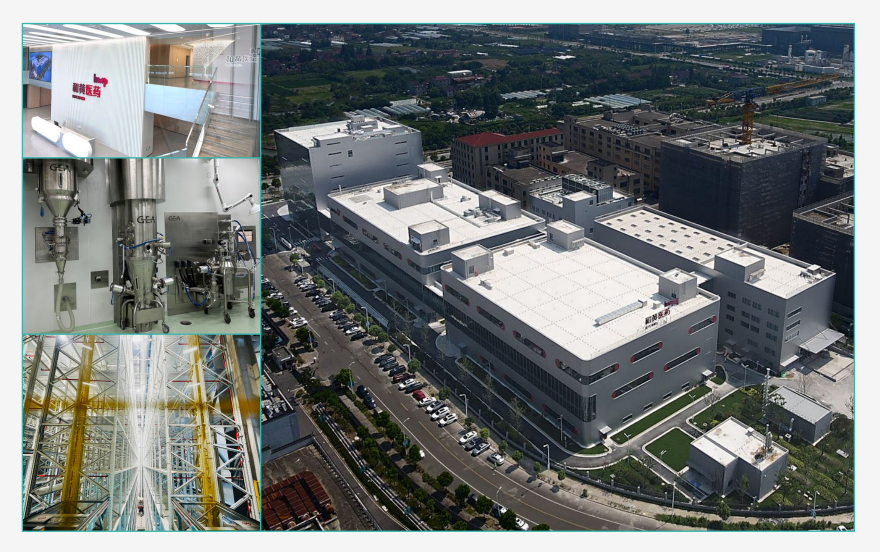


Shanghai facility construction is completed

- Expands capacity >5x
- Qualification of facility & equipment in 2023
- Clinical supplies manufacturing in 2023
- Commercial supplies manufacturing around 2025

Solar panels installation in 2023

In line with sustainability initiatives



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HUTCHMED registration/potential registration studies

15+ programs for seven drug candidates supporting potential near-term NDA filings

Drug	Study	Target Disease	Region	Design (N, arms, 1° endpoint)	Status	Est. (s)NDA filing if positive
FRUQ	FRESCO-2	3L+ colorectal cancer	Global	~690, treatment vs. BSC, OS	US Priority Review granted, EMA validated, Japan filing in 2023	US PDUFA Nov 30
FRUQ	FRUTIGA	2L GC, combo with chemo	China	~700, combo vs. chemo, OS & PFS	sNDA in China accepted April 2023	Review ongoing
SOVLE	ESLIM-01	2L immune thrombocytopenia	China	~180, 2 arms (placebo), DRR	Positive topline Aug '23	End of 2023
AMDIZ	3L FL	3L follicular lymphoma	China	~100, 1 arm, ORR	LPI Feb '23	End of 2023
SAVO*	Confirmatory	NSCLC, MET Exon 14 alteration	China	~160, 1 arm, ORR	LPI H1 '23	2024
FRUQ	2L EMC	2L EMC, combo with PD-1	China	~130, 1 arm, ORR	LPI July '23, China BTD	2024
AMDIZ	2L MZL	2L marginal zone lymphoma	China	~80, 1 arm, ORR	FPI Apr '21	2024
TAZ^	Bridging	3L follicular lymphoma	China	~40, 2 arms (EZH2+ or wt), ORR	FPI Jul '22	2024
SAVO*	SACHI	2L EGFR TKI refractory NSCLC, MET+	China	~250, combo vs. chemo, PFS	FPI Nov '21	2024
SAVO*	SAVANNAH	2/3L Tagrisso® refractory NSCLC, MET+	Global	New cohort for pot. AA	FPI Jan '19 Re-opened in Sept 2022	2024
SURU	SURTORI-01	2L NEC, combo with PD-1	China	~190, combo vs. chemo, OS	FPI Sep '21	2024
SAVO*	GASTRIC	Ref. GC, MET amplified	China	~60, 1 arm, ORR	FPI Jul '21 Reg. cohort opened Mar 2023	2025
FRUQ	2L RCC	2L RCC, combo with PD-1	China	~260, 2 arms, PFS	FPI Oct '22	2025
SOVLE	wAIHA	2L wAIHA	China	~110, 2 arms (placebo), Hb response	FPI Sep '22	2025
SAVO*	SANOVO	1L EGFRm+ NSCLC, MET+	China	~320, combo vs. Tagrisso, PFS	FPI Sep '21	2026
SAVO*	SAMETA	MET driven PRCC, combo with PD-L1	Global	~200, 3 arms combo vs. monos, PFS	FPI Oct '21	2026
SAVO*	SAFFRON	2/3L Tagrisso® refractory NSCLC, MET+	Global	~320, combo vs. chemo, PFS	FPI Aug '22	2026
453	IHCC, FGFR2	IHCC, FGFR2 fusion	China	~90, 1 arm, ORR	FPI Sept '20 Reg. cohort opened Mar 2023	2026

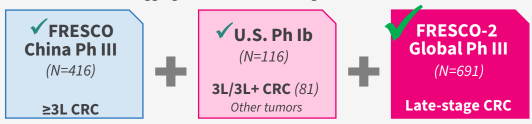


Fruquintinib global filings based on FRESCO-2 & FRESCO

US NDA PDUFA date Nov 30, 2023; MAA validated in June 2023; filing in Japan planned for 2023

Fruquintinib – Basis for global filings

Aggregation of China, U.S. & global studies

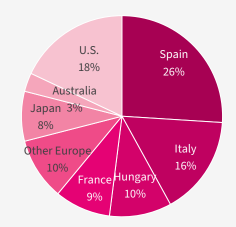


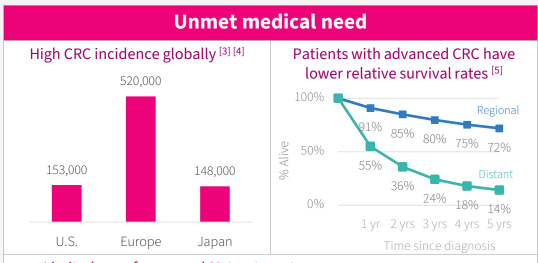
Consistency of effect across late-stage settings enriches the continuum of care

0					
	FRES	CO [1]	FRESCO-2 ^[2]		
	Fruq	Pbo	Fruq	Pbo	
mOS, mo.	9.3	6.6	7.4	4.8	
[95% CI]	[8.2-10.5]	[5.9-8.1]	[6.7-8.2]	[4.0-5.8]	
HR	0.	65	0.	66	
(95% CI, p-value)	(0.51-0.83	8, p<0.001)	(0.55-0.80, p<0.001)		
mPFS, mo.	3.7	1.8	3.7	1.8	
[95% CI]	[3.7-4.6]	[1.8-1.8]	[3.5-3.8]	[1.8-1.9]	
HR	0.26		0.	32	
(95% CI, p-value)	(0.21-0.34	!, p<0.001)	(0.27-0.39, p<0.001)		
	DCO: Janua	anv 17 2017	DCO: lun	0.24.2022	

FRESCO-2 MRCT

691 pts | 14 countries





Limited use of approved 3L treatments

- Regorafenib (approved Q3 2012)
- TAS-102 (approved Q3 2015) +/- bevacizumab
- Chemotherapy, anti-VEGF & anti-EGFR agents used across all lines

Newer treatment options focus on discrete actionable mutations

- ~10% BRAF mutation [6]
- ~15% MSI-H or dMMR ^[7] ^[8]
- 3-5% HER2 alterations [9]

[1] Li J, et al. Effect of Fruquintinib vs Placebo on Overall Survival in Patients With Previously Treated Metastatic Colorectal Cancer: The FRESCO Randomized Clinical Trial. JAMA. 2018;319(24):2486-2496. doi:10.1001/jama.2018.7855; [2] Dasari A, et al. Fruquintinib versus placebo in patients with refractory metastatic colorectal cancer (FRESCO-2): an international, multicentre, randomised, double-blind, phase 3 study. Lancet. 2023;402(10395):41-53. doi:10.1016/S0140-6736(23)00772-9; [3] International Agency for Research on Cancer; [4] D'Haene N, et al. Clinical application of targeted next-generation sequencing for colorectal cancer patients: a multicentric Belgian experience. Oncotarget. 2018;9(29):20761-20768. Published 2018 Apr 17. doi:10.18632/oncotarget.25099; [7] Venderbosch S, et al. Mismatch repair status and BRAF mutation status in metastatic colorectal cancer patients: a pooled analysis of the CAIRO, CAIRO2, COIN, and FOCUS studies. Clin Cancer Res. 2014;20(20):5322-5330. doi:10.1158/1078-0432.CCR-14-0332; [8] André T, et al. Pembrolizumab in Microsatellity-High Advanced Colorectal Cancer. N Engl J Med. 2020;383(23):2207-2218. doi:10.1056/NEJMoa2017699; [9] Ahcene Djaballah S, et al. HER2 in Colorectal Cancer: The Long and Winding Road From Negative Predictive Factor to Positive Actionable Target. Am Soc Clin Oncol Educ Book. 2021;42:1-14. doi:10.1200/EDBK 351354.

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Fruquintinib partnership with Takeda progressing well

Working closely together and preparing to transfer regulatory sponsorship

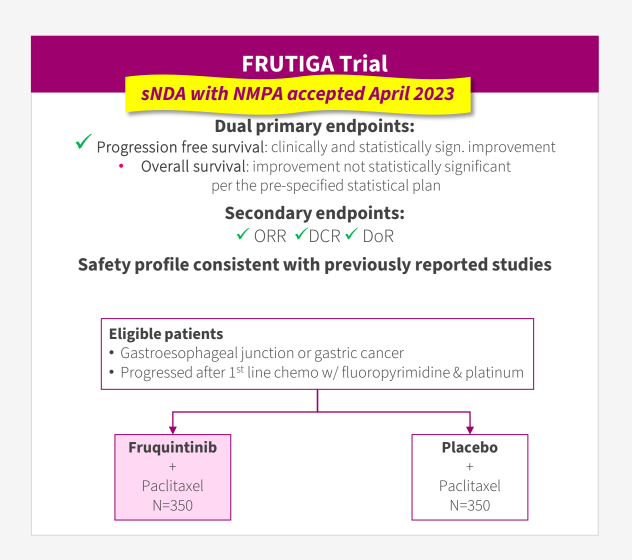
Financials & governance	✓ US\$400m upfront receivedJoint team established and started collaboration	
Regulatory Filings	 ✓ Completed U.S. NDA rolling submission in March 2023; PDUFA Nov 30, 2023 ✓ MAA submission in Europe validated June 2023 • Submit JNDA to the Japan PMDA in 2023 	
Commercial Launch	Takeda initiating launch readiness in advance of PDUFA date	4
Further Clinical Development (LCM)	 Joint team discussions and advisory boards held to discuss LCM strategy HUTCHMED ongoing programs in China may inform decisions 	

Fruquintinib 2L gastric: sNDA accepted April 2023



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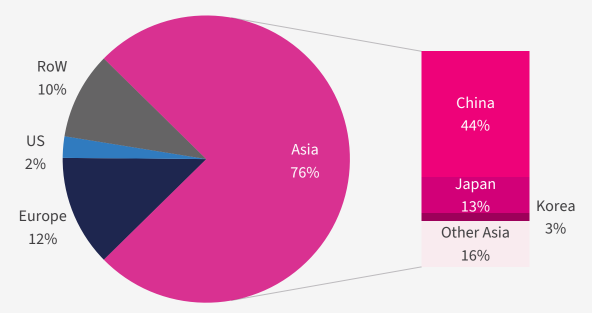
FRUTIGA combo study with paclitaxel in 2L gastric cancer to expand patients served



5th MOST COMMONLY DIAGNOSED CANCER WORLDWIDE DISPROPORTIONATELY AFFECTS ASIA

- 1.09 million new patients globally
- China, Japan & Korea account for ~60% of newly diagnosed
- 478,500 diagnosed in China every year

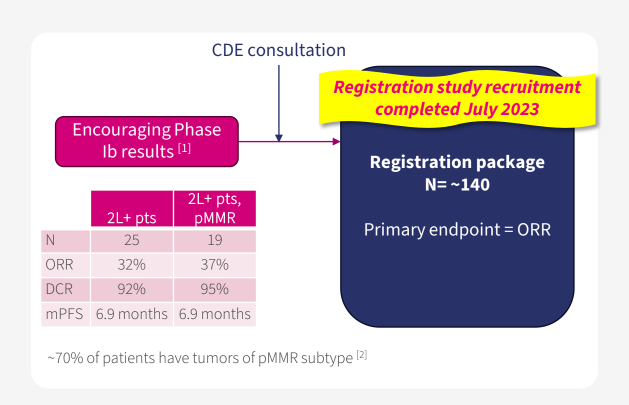
Annual incidence of gastric cancer by geography



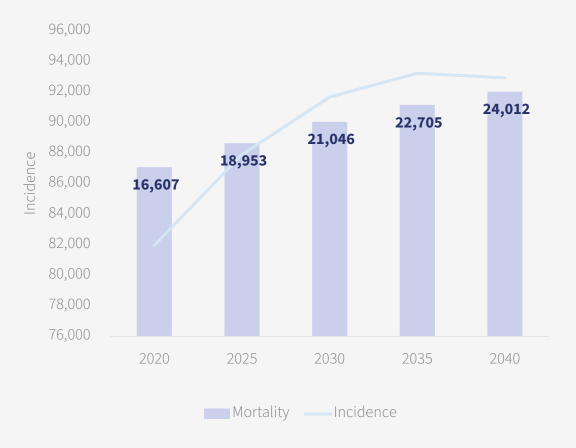
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Fruquintinib endometrial cancer: Lead ICI combo in China

Breakthrough Therapy Designation in China for pMMR subtype



Medical need: Mortality from EMC projected to grow in China [3]



^[1] CSCO 2021

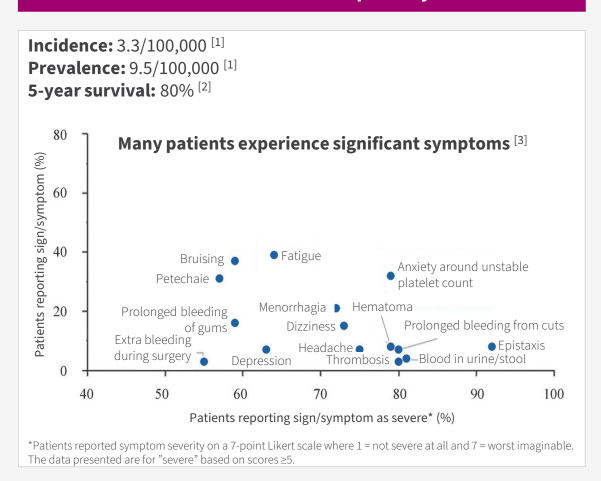
^[2] National Comprehensive Network Clinical Practice Guidelines in Oncology for Uterine Neoplasms. Version 2.2023 - April 28, 2023.

Primary ITP landscape

HUTCHMED

Significant burden of disease remains

Unmet medical need in primary ITP



Treatments for chronic ITP [4]

Agent	Response (1x PLT ≥50×10 ⁹ /L)	Durable response	Use of rescue medication	2022 Revenues for all indications ^[5]
TPO-RA treatmer	nt increases plate	let production		
PROMACTA® (eltrombopag) ^[6]	59-70% (6 weeks) ^[7]	60% (6/8 visits, weeks 18-26) ^[8]	18% (vs 40%) ^[8]	\$2.1 billion (approved for ITP + severe aplastic anemia)
NPLATE® (romiplostim) ^[6]	79-88% (24 weeks)	38-61% (6/8 visits, weeks 16-24)	20-26% (vs 57-62%)	\$1.3 billion (approved for ITP + radiation sickness)
Hetrombopag ^[9]	59-64% (8 weeks)	Median maximum continuous durations of response: 64 days	10-13% (vs 38%)	n/a (approved for ITP + severe aplastic anemia)
DOPTELET® (avatrombopag) ^[6]	66% at day 8	Median cumulative nu response without res 12 wee	\$245 million / \$107 million to China distributor Fosun (approved for ITP + chronic liver disease)	
Treatments to de	crease platelet d	estruction		
RITUXAN® (rituximab) ^[4]	~60% (4 weeks of tx)	20-25%	n/a	Not FDA approved for ITP
TAVALISSE® (fostamatinib) ^[6]	43% (12 weeks)	16-18% (4/6 visits, weeks 14-24)	30% (vs 45%)	\$76 million (US revenues, approved for ITP)

[1] Lambert MP, Gernsheimer TB. Clinical updates in adult immune thrombocytopenia. *Blood*. 2017 May 25;129(21):2829-2835; [2] Mannering N, et al. Survival in adult patients with chronic primary and secondary immune thrombocytopenia: A population-based study. *Transfusion*. 2023;63(2):415-426. doi:10.1111/trf.17212; [3] Adapted from Wang RT, et al. [A physician-patient survey for primary immune thrombocytopenia: Chinese subgroup analysis of I-WISh International Survey]. *Zhonghua Xue Ye Xue Za Zhi*. 2021;42(5):369-375. doi:10.3760/cma.j.issn.0253-2727.2021.05.004; [4] Provan D, Arnold DM, Bussel JB, et al. Updated international consensus report on the investigation and management of primary immune thrombocytopenia. *Blood Adv*. 2019;3(22):3780-3817. doi:10.1182/bloodadvances.2019000812; [5] company reports; [6] USPI; [7] Study 773A and B from US PI; [8] RAISE study from US PI; [9] Mei H, et al. A multicenter, randomized phase III trial of hetrombopag: a novel thrombopoietin receptor agonist for the treatment of immune thrombocytopenia. *J Hematol Oncol*. 2021;14(1):37. Published 2021 Feb 25. doi:10.1186/s13045-021-01047-9.



Sovleplenib 2L ITP: NDA filing around YE2023

Highly differentiated oral Syk inhibitor with breakthrough therapy designation in China

ESLIM-01 Trial NDA filing with NMPA planned around YE2023 Primary endpoint: durable response rate ✓ Platelet count ≥50×10⁹ /L on at least 4 of 6 scheduled visits during week 14 to 24 **Secondary endpoints:** ✓ ORR ✓ Safety Safety profile consistent with previously reported studies Eligible patients FCOG PS score of 0~1 • Duration of disease is > 6 months • Intolerance or insufficient response, or recurrence after at least one standard drug therapy A history of response to previous ITP therapy Sovleplenib 300mg QD Placebo OD N ≈ 120 $N \approx 60$

Sovleplenib encouraging Phase I/II results

Results from China Phase I/II in R/R primary ITP

- Oral, fast onset of efficacy ORR 80%, Durable ORR 40%
- Robust efficacy in heavily pre-treated patients
- Similar efficacy with or without prior TPO/TPO-RA therapies

Breakthrough Therapy Designation in China

	Sovleplenib – 300 mg, once daily						
	Double-blinded Pts O-24 weeks Cross-over P 9-24 weeks		Total				
ORR: n (%)	75.0% (12/16)	100.0% (4/4)	80.0% (16/20)				
Durable ORR: n (%)	31.3% (5/16)	75.0% (3/4)	40.0% (8/20)				
Use of rescue medication	6% (1/16)	0	5% (1/20)				

Liu X, et al. Sovleplenib (HMPL-523), a novel Syk inhibitor, for patients with primary immune thrombocytopenia in China: a randomised, double-blind, placebo-controlled, phase 1b/2 study [published online ahead of print, 2023 Apr 4]. Lancet Haematol. 2023;S2352-3026(23)00034-0. doi:10.1016/S2352-3026(23)00034-0

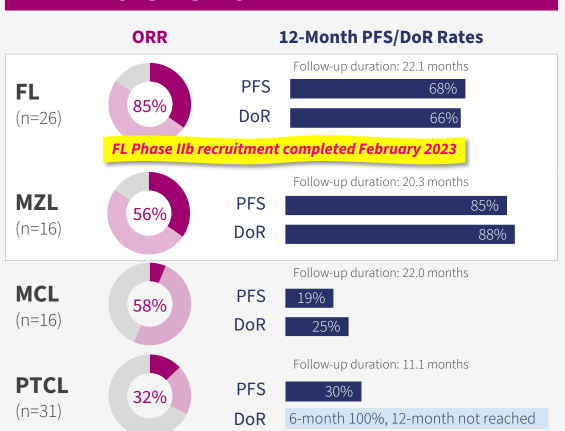
Amdizalisib progressing towards NDA in 2023 in FL



profile

China registration studies supported by differentiated proof-of-concept data

Encouraging single agent activity in indolent NHL [1]



ML 🗲	
23	Highly favorable safety
25	mignly lavorable salety
A N	

All AEs / ≥Gr3 AEs	Amdizalisib ^[1]	Aliqopa®	Copiktra®	Linperlisib ^[3]	Parsaclisib	
30mg QD (copanlisib) (duv		(duvelisib) ^[2]	80mg QD	CITADEL-203 (FL) ^[4]	China cohort ^[5]	
N	153	244	442	84	103	61
Neutropenia*	37% / 12%	32% / 29%	63% / 43%	46% / 16%	48% / 16%	49% / 16%
Leukopenia	5% / 2%	36% / 27%	29% / 8%*	36% / 5% *	<10%	33% / 2%
Anemia	22% / 5%	na	20% / 11%	<10%	34% / 3%*	<20%
Thrombocytopenia	5% / 1%	22% / 8%	17% / 10%	16% / 4%	22% / 0%*	<20%
Diarrhea	16% / 4%	36% / 5%	50% / 23%	16% / 1%	44% / 14%	<20%
Rash	24% / 6%	15% / 2%	31% / 9%	12% / 1%	14% / 3%	<20%
ALT increased	32% / 1%	na / 2%	40% / 8%	23% / 1%	30% / 2%	23% / 0%
AST increased	29% / 1%	na / 2%	37% / 6%	18% / 1%	29% / 0%	<20%
Pyrexia	16% / 1%	na	26% / 2%	<10%	19% / 3%	<20%
Pneumonia	25% / 16%	21% / 14%**	21% / 15%	20% / 19%	<10%	<20%
Hypertension	7% / 1%	35% / 29%	na	na / <2%	<10%	<20%
Hyperglycemia	9% / 0%	54% / 34%	na	13% / 1%	<10%	<20%
Interstitial lung disease	4% / 1%	na	na	5 % / 4 %	<10%	<20%
Lipase increased*	15% / 8%	21% / 8%	36% / 16%	14% / 4%	<10%	<20%
AES leading to:						
Discontinuation	12%	24%	35%	18%	25%	na
Dose reduction	17%	24%	23%	na	20%	na
Dose interruption	46%	64%	64%	43%	48%	na
Current status in China	Ph. II reg study FL & MZL	Approved for 3L+ FL	Approved for 3L+ FL	Approved for 3L+ FL	NDA accepte	ed for 3L+ FL

Phase Ib data as of January 31, 2023



Savolitinib - major late-stage expansion

7 registrational studies – 3 global & 4 in China



GLOBAL – led by AstraZeneca

2/3L TAGRISSO® refractory NSCLC w/ MET aberration

SAVANNAH study – continue evaluation for potential accelerated approval; first data presentation at WCLC

2/3L TAGRISSO® refractory NSCLC w/ MET aberration

Savolitinib + TAGRISSO® Phase III registration study
 - SAFFRON Study initiated in 2022

MET-driven Papillary Renal Cell Carcinoma (PRCC)

- Savolitinib + IMFINZI® vs. SUTENT® monotherapy vs. IMFINZI® monotherapy Phase III registration study
- FPI in October 2021 SAMETA Study

CHINA – led by HUTCHMED

MET Exon14 skipping NSCLC

NDA conditional approval in June 2021

Confirmatory Phase IIIb study – 1L cohort results at WCLC

2L EGFR TKI refractory NSCLC w/ MET amplification

- Savolitinib + TAGRISSO® Phase III registration study
- FPI in November 2021 **SACHI Study**

1L EGFRm+ NSCLC w/ MET overexpression

- Savolitinib + TAGRISSO® Phase III registration study
- FPI in September 2021 **SANOVO Study**

Gastric cancer w/ MET amplification

Single arm study with potential for registration

Registration cohort FPI March 2023

China Breakthrough designation for 3L+ Aug 2023

2



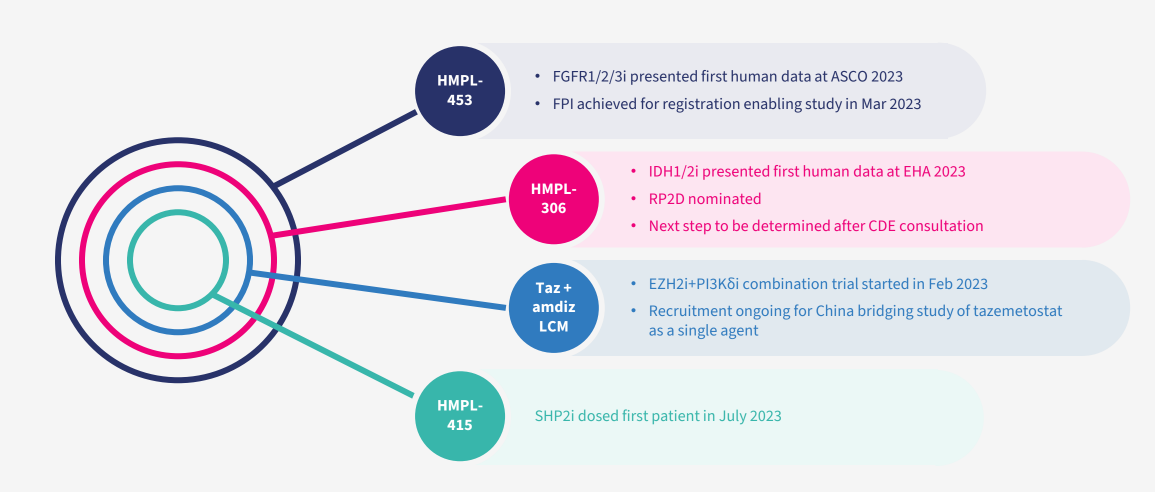




Our innovative engine remains productive



Progression of early-stage programs in 2023



Clinical deliverables in 2023-4



To make significant progress with multiple late-stage programs

Regulatory activities		
Fruquintinib mono US, EU, Japan for 3L+ CRC	\rightarrow	✓ US NDA PDUFA Nov 30, 2023 ✓ EMA MAA validated June 2023 Japan filing to follow in 2023
Fruquintinib + chemo <i>China for 2L GC</i>	\rightarrow	✓ China sNDA filing accepted April 2023
Sovleplenib mono <i>China for 2L ITP</i>	\rightarrow	China NDA filing around YE 2023 ✓ Readout August 2023
Fruquintinib + sintilimab China for 2L EMC*	\rightarrow	✓ China Breakthrough Designation in July 2023

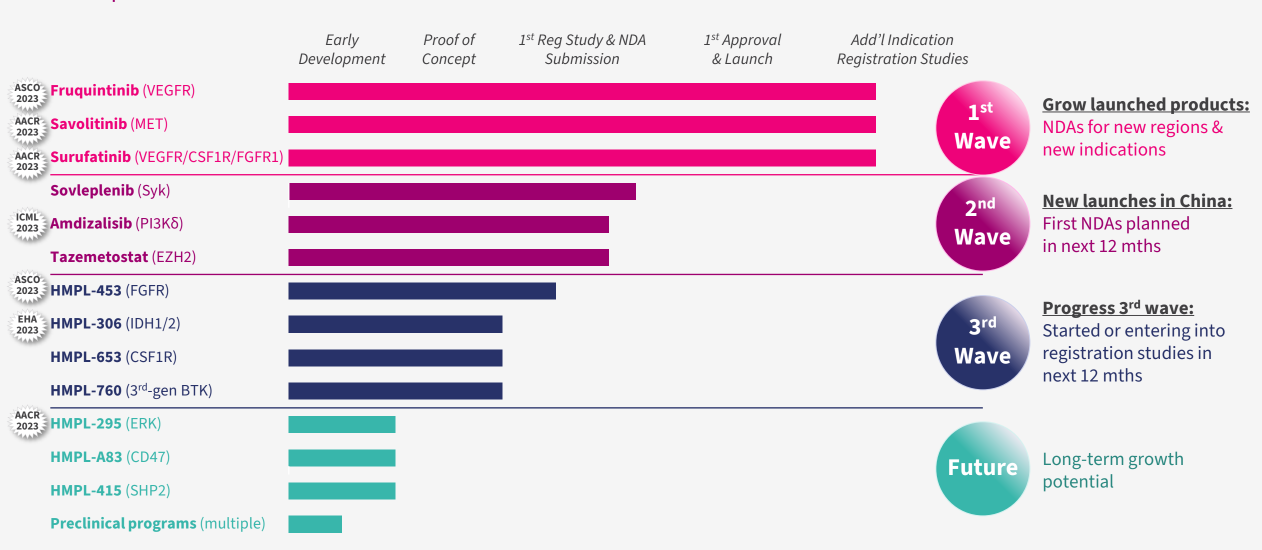
Readouts		
Amdizalisib mono China for 3L FL*	\rightarrow	Readout late 2023 Completed recruitment Feb 2023
Savolitinib mono China confirm. for NSCLC, MET ex14	\rightarrow	Cohort 2 readout YE 2023 Cohort 1 results at WCLC 2023 Completed recruitment H1 2023
Fruquintinib + sintilimab China for 2L EMC*	\rightarrow	H1 2024 Completed recruitment July 2023

Continued progress on additional registration studies				
Tazemetostat mono <i>China for 3L FL*</i>	\rightarrow	Complete recruitment H2 2023		
Savolitinib + osimertinib Intl for 2L NSCLC, MET+*	\rightarrow	Complete recruitment H2 2023		
Fruquintinib + sintilimab <i>China for 2L RCC</i>	\rightarrow	Complete recruitment YE 2023		
Amdizalisib mono China for 2L MZL*	\rightarrow	Complete recruitment H1 2024		
Savolitinib + osimertinib China for 2L NSCLC, MET+*	\rightarrow	Complete recruitment mid 2024		
Savolitinib + osimertinib China for 1L NSCLC, MET+*	\rightarrow	Complete recruitment H2 2024		
Savolitinib mono China for Ref. GC, MET+*	\rightarrow	✓ Early readout at AACR 2023 ✓ China Breakthrough Designation Aug 2023 Complete recruitment in H2 2024		
Surufatinib + toripalimab <i>China for 2L NEC</i>	\rightarrow	Complete recruitment H2 2024		
✓ HMPL-453 mono China for IHCC, FGFR2 fusion*	\rightarrow	✓ Ph Ib/II results at ASCO 2023✓ Reg. cohort FPI March 2023		
Sovleplenib mono <i>China for 2L wAIHA*</i>	\rightarrow	✓ Enrolled Phase II part H1 2023 Decision to proceed YE 2023		

Progressing the pipeline to maximize the value of each asset



Next potential new indications & medicines



Thank you



www.hutch-med.com



References & Abbreviations



ADS = American depositary share.

AIHA = autoimmune hemolytic anemia.

ALL = acute Lymphoblastic Leukemia

AML = acute myeloid leukemia.

API = active pharmaceutical ingredient.

ASCO = American Society of Clinical Oncology.

 $ASCO\:GI = ASCO\:(American\:Society\:of\:Clinical\:Oncology)\:Gastrointestinal$

Cancers Symposium

ASH = American Society of Hematology

bsAb = bi-specific antibody

BID = twice daily.

BRAF = B-Raf.

BSC = best supportive care.

BTK = bruton's tyrosine kinase.

CBCL= cutaneous B-cell lymphoma.

CER = constant exchange rate.

CI = confidence interval.

CLD = chronic liver disease.

CLL/SLL = chronic lymphocytic leukemia and small lymphocytic lymphoma

CRC = colorectal cancer.

CRL = complete response letter.

CSF-1R = colony-stimulating factor 1 receptor.

DCO = data cutoff

DDI = drug-drug interactions.

Deutsche Bank AG = Deutsche Bank AG, Hong Kong Branch.

DLBCL = diffuse large B-cell lymphoma

dMMR = deficient mismatch

DoR = duration of response. DRR = durable response rate.

epNET = extra-pancreatic neuroendocrine tumor.

EGFR = epidermal growth factor receptor.

EGFRm+ = epidermal growth factor receptor mutated.

EMA = European Medicines Agency.

EMC = endometrial cancer.

Epizyme = Epizyme Inc.

ERK = extracellular signal-regulated kinase.

ES = epithelioid sarcoma.

EU = European Union.

EZH2 = enhancer of zeste homolog 2. FISH = fluorescence in situ hybridization.

FISH5+ = MET amplification as detected by FISH with MET copy number ≥ 5

and/or MET: CEP signal ratio ≥ 2 .

FISH10+ = MET amplification as detected by FISH with MET copy number \geq 10.

FDA = Food and Drug Administration.

FGFR = fibroblast growth factor receptor.

FL = follicular lymphoma.

FPI = first patient in.

GAAP = Generally Accepted Accounting Principles.

GC = gastric cancer.

GI = gastrointestinal.

HBYS = Hutchison Whampoa Guangzhou Baiyunshan Chinese Medicine Company Limited.

HKEX = The Main Board of The Stock Exchange of Hong Kong Limited.

HL = Hodgkin's lymphoma.

HR = hazard ratio.

HSBC = The Hongkong and Shanghai Banking Corporation Limited.

Hutchison Sinopharm = Hutchison Whampoa Sinopharm Pharmaceuticals

(Shanghai) Company Limited. IDH = Isocitrate dehydrogenase.

In-market sales = total sales to third parties provided by Eli Lilly (ELUNATE*), AstraZeneca (ORPATHYS*) and HUTCHMED (SULANDA* and TAZVERIK*).

 $HCPs = healthcare\ professionals$

ICI = immune checkpoint inhibitor IHC = immunohistochemistry.

IHC50+ = MET overexpression as detected by IHC with 3+ in \geq 50% tumor cells.

IHC90+=MET overexpression as detected by IHC with 3+ in $\geq 90\%$ tumor cells.

iNHL = indolent Non-Hodgkin's Lymphoma.

I/O = Immuno-oncology.

IND = Investigational New Drug (application).

IR = *independent review*.

IRC = independent review committee.

ITP = Immune thrombocytopenia purpura.

Lilly = Eli Lilly and Company.

 $\it MAA = Marketing Authorization Application.$

 $MAPK\ pathway = RAS-RAF-MEK-ERK\ signaling\ cascade.$

 $Mab = monoclonal\ antibody.$

MCL = mantle cell lymphoma.

MDS/MPN = myelodysplastic/myeloproliferative neoplasms

MET = mesenchymal epithelial transition factor.

MRCT = multi-regional clinical trial.

 $\mathit{MSI-H} = \mathit{high}\ \mathit{levels}\ \mathit{of}\ \mathit{microsatellite}\ \mathit{instability}.$

MSS / pMMR = microsatellite stable / mismatch repair proficient.

MZL = marginal zone lymphoma.

na = not available.

NDA = New Drug Application.

NEC = neuroendocrine carcinoma.

NETs = neuroendocrine tumors.

NHL = Non-Hodgkin's Lymphoma.

NME = new molecular entity.

NR = not reached.

NRDL = National Reimbursement Drug List.

NSCLC = non-small cell lung cancer.

ORR = objective response rate.

OS = overall survival.

QD = once daily.

PD = progressive disease.

PD-L1 = programmed cell death ligand 1.

PFS = progression-free survival.

PI3K δ = phosphoinositide 3-kinase delta.

PJP = pneumocystis jirovecii pneumonia.

PMDA = Pharmaceuticals and Medical Devices Agency.

pNET= pancreatic neuroendocrine tumor.

ccRCC = clear cell renal cell carcinoma.

PRCC = papillary renal cell carcinoma.

PTCL = peripheral T-cell lymphomas. R&D = research and development.

SAA = severe aplastic anemia.

SHP2 = Src homology-2 domain-containing protein tyrosine phosphatase-2

SHPL = Shanghai Hutchison Pharmaceuticals Limited.

SOC = standard of care.

Syk = spleen tyrosine kinase.

TKI = tyrosine kinase inhibitor.

TPO-RA = thrombopoietin receptor agonists.

Tx = treatment.

VEGF = vascular endothelial growth factor.

VEGFR = vascular endothelial growth factor receptor.

wAIHA = warm antibody autoimmune hemolytic anemia.

WM/LPL = Waldenström macroglobulinemia and lymphoplasmacytic lymphoma.

WT = wild-type.

WCLC = IASLC World Conference on Lung Cancer.