# elevar therapeutics

# Elevating Treatment Experiences and Outcomes for Patients

February 2024

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The market data and certain other statistical information used throughout this presentation are based on independent industry publications, governmental publications, reports by market research firms or other independent sources. Some data are also based on our good faith estimates. Although we believe these third-party sources are reliable, we have not independently verified the information attributed to these third-party sources and cannot guarantee its accuracy and completeness. Similarly, our estimates have not been verified by any independent source.

By attending or receiving this presentation and viewing the related video, you acknowledge that you will be solely responsible for your own assessment of the market and our market position and that you will conduct your own analysis and be solely responsible for forming your own view of the potential future performance of our business.

# **Executive Summary**

# Elevar Therapeutics: An Oncology-Focused, Fully Integrated Biopharmaceutical Company

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# PHASE 3 CARES-310 STUDY, WITH POTENTIAL APPROVAL IN MAY 2024<sup>1</sup>

CARES-310 study shows a mOS of 22.1 months in 1L uHCC patients<sup>1</sup> Current combination therapies for uHCC showed mOS of 16.4-19.2 months<sup>2,3</sup>

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### HCC AND uHCC REPRESENT A \$10B+ MARKET OPPORTUNITY

Approximately 15,000+ patients<sup>4-6</sup> receiving 1L uHCC treatment yearly in the US, and incidence and mortality rates increasing<sup>7</sup>



### **EXPERIENCED MANAGEMENT TEAM**

Strong regulatory and commercial team, with a combined 70+ drug approvals and launches



### WELL CAPITALIZED TO EXECUTE ON OUR GOALS

Fully owned subsidiary of HLB Co., Ltd. (KOSDAQ:028300) with a strong balance sheet

References: 1. Qin S, et al. Lancet. 2023;402(10408):1133-1146. 2. Cheng A-I, et al. J Hepatol. 2022;76(4):862-873. 3. Abou-Alfa GK, et al. NEJM Evid. 2022;1(8):doi:10.1056/EVIDoa2100070. 4. Siegel RL, et al. CA Cancer J Clin. 2023;73:17-48. 5. Llovet JM, et al. Nat Rev Dis Primers. 2021;7(1):6. 6. Wang EA, et al. Int J Clin Pract. 2017;71(11). 7. Golabi P, et al. Medicine (Baltimore). 2017;96(9):e5904.

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# **Visionary Management Team**



# **Corporate Highlights**

Elevar is a rapidly growing, fully integrated biopharmaceutical company built on the promise of elevating treatment experiences and outcomes for patients who have limited or inadequate therapeutic options

<b>Resourced</b>	<b>Experienced</b>	<b>Focused</b>
for the evolution to commercialization in the US,	in clinical and commercial development	on solid tumors that respond to anti-VEGF TKIs
and seeking partners to co-develop rivoceranib	and launch experience	that are effective as mono or combination therapy
<ul> <li>Elevar Therapeutics is a majority-owned subsidiary of HLB Co, LTD., a publicly traded company on the Korean KOSDAQ exchange (028300.KQ) and has a market cap of approximately USD\$3B</li> <li>Strong intellectual property protection though 2037</li> <li>Demonstrated success with existing partners</li> </ul>	<ul> <li>Executive Leadership Team with extensive global experience in both big and small Biotech</li> <li>Experienced in building/scaling organizations with more than 70 FDA approvals and launches</li> <li>Plans developed for proven oncology launch footprint to support optimal targeted reach</li> </ul>	<ul> <li>Rivoceranib has the potential to be a best-in-class small molecule, TKI selective for the VEGF receptor; orally administered</li> <li>Camrelizumab is a top-selling anti-PD-1 monoclonal antibody in China generating significant sales with 8 approved indications; intravenously administered</li> <li>CARES-310 study shows a mOS of 22.1 months in 1L uHCC patients<sup>1</sup></li> <li>Elevar has global rights to camrelizumab and rivoceranib for HCC (excluding Greater China and Korea)</li> </ul>

# **Elevar Key Milestones**

Japan filing and approval in 2026+

EU filing and approval in late 2025+

US commercial launch Q3 2024



### **PDUFA date set**

for May 16, 2024, for rivoceranib and May 31, 2024, for camrelizumab



### **BLA/NDA filings**

in May 2023 and accepted in July 2023

# **Company Overview and Transaction Rationale**

Camrelizumab and rivoceranib being developed for HCC and uHCC – areas of large unmet medical need representing a **\$10B+** potential market opportunity

### **Drug Candidates**

### **Rivoceranib**

- Commercialized by Hengrui Pharma in China (under the name apatinib)
- Approved in China for:
  - Gastric cancer 1L monotherapy (2014)
  - Advanced hepatocellular carcinoma (HCC) 2L monotherapy (2020)
  - Unresectable hepatocellular carcinoma (uHCC) in combination with Hengrui Pharma's camrelizumab 1L (January 2023)

### Camrelizumab

- Commercialized by Hengrui Pharma in China (under the brand name AiRuiKa®)
- One of the top-selling anti-PD-1s in China with 8 approved indications

# **Rivoceranib Studied in More Than 6,000 Patients Worldwide** for Multiple Indications<sup>1,2</sup>

	Indication	Therapy/Line	Discovery	Lead Optimization	IND Enabling Pre-Clinical	Phase 1b Clinical	Phase 2 Clinical	Phase 3 Clinical	NDA Accepted
ns	uHCC* (Hengrui Collaboration)	+ Camrelizumab combo/1L							
rograr	ACC*	Recurrent or Metastatic, Monotherapy							
inical p	GC*	Monotherapy/3L/4L							
CI	CRC	+ LONSURF <sup>®</sup> combo/3L							

tory	GC	+ Paclitaxel combo/2L				
Explora	Multiple Solid Tumors (Sarcoma)	+ OPDIVO <sup>®</sup> combo				

\*Orphan Drug Designation (ODD).

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**References: 1.** Elevar Therapeutics. Press release. Accessed September 13, 2023. https://elevartherapeutics.com/2023/08/03/elevar-therapeutics-to-hostaugust-10-virtual-kol-event-on-phase-3-study-of-rivoceranib-in-combination-with-camrelizumab-in-unresectable-hepatocellular-carcinoma-uhcc/ **2.** Elevar Therapeutics. Press release. Accessed September 14, 2023. https://elevartherapeutics.com/2023/07/17/elevar-therapeutics-announces-fda-acceptancefor-filing-of-new-drug-application-for-rivoceranib-in-combination-with-camrelizumab-as-a-first-line-treatment-for-unresectable-hepatocellular-carcinoma/

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## Camrelizumab Has Been Studied in More Than 5,000 Patients Worldwide Across Multiple Indications<sup>1</sup>

	Indication	Therapy	Phase 1 Clinical	Phase 2 Clinical	Phase 3 Clinical	BLA Filed	Approved
	NSQ-NSCLC EGFR(-)/ALK(-)	Chemo combo					
	NPC	Chemo combo					
U	SQ-NSCLC EGFR(-)/ALK(-)	Chemo combo					
Lin	Esophageal Carcinoma	Chemo combo					
-irst	HCC*	Rivoceranib combo					
	GC/GEJC	Chemo combo sequenced by camrelizumab + rivoceranib					
	NSQ-NSCLC EGFR(-)/ALK(-)	Famitinib or Placebo combo plus chemo					
	Cervical Cancer	Famitinib combo					
	НСС	Monotherapy					
Line	Esophageal Carcinoma	Monotherapy					
puo	cHL	Monotherapy					
Sec	NPC	Monotherapy					
ΛΙ	ТИВС	Chemo combo					
		1					
/ant	TNBC	Combo/Neoadjuvant					
var Ijuv							
dju pac							
A S	НСС	Rivoceranib combo/Adjuvant					

Reference: 1. Elevar Therapeutics. Press release. Accessed September 14, 2023. https://elevartherapeutics.com/2023/07/17/elevartherapeutics-announces-fda-acceptance-for-filing-ofnew-drug-application-for-rivoceranib-in-combinationwith-camrelizumab-as-a-first-line-treatment-forunresectable-hepatocellular-carcinoma/ \*1L treatment for HCC approved in China.

Note: All approved treatments are approved in China only, not U.S. or E.U. Note: Hengrui's pipeline is shown, Elevar only has the HCC combination therapy rights globally ex Greater China and South Korea.

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ALK=anaplastic lymphoma kinase; cHL=classic Hodgkin lymphoma; EGFR=epidermal growth factor receptor; GC=gastric cancer; GEJC=gastroesophageal junction cancer; HCC=hepatocellular carcinoma; NPC=nasopharyngeal cancer; NSCLC=non small-cell lung cancer; NSQ=non-squamous; SQ=squamous; TNBC=triple-negative breast cancer.

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# **Technology and Product Candidates**

# **Opportunity in HCC**

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With ~15,000+ patients<sup>1-3</sup> receiving 1L uHCC treatment yearly in the US and incidence and mortality rates increasing, **HCC represents the fastest-rising cause** of cancer-related deaths in the US<sup>4,5</sup>

Typically diagnosed late in its course where **liver function is already severely declining**, survival at diagnosis is only ~6-20 months with very low five-year survival rates<sup>4</sup>

HCC is currently the **2<sup>nd</sup> leading cause of cancer-related death** in Asia and the 6<sup>th</sup> most common in Western countries<sup>5</sup> with ~60-70% of patients being exposed / opting to systemic therapy at some point,<sup>2,6</sup> the number of which should increase with **safer and more tolerable treatment options** 



# **O** Available Treatment

40+% of patients are expected to receive an angiogenesis inhibitor in combination with an ICI, such as atezolizumab + bevacizumab<sup>7,8</sup>

Checkpoint inhibitor and TKI combinations offer promise because toxicity profiles do not overlap<sup>9</sup>

Despite the latest advancements in HCC treatments, an urgent need remains for more efficacious, tolerable treatments due to the disease's severity and low survival rates

References: 1. Siegel RL, et al. CA Cancer J Clin. 2023;73(1):17-48. 2. Llovet JM, et al. Nat Rev Dis Primers. 2021;7(1):6. 3. Wang EA, et al. Int J Clin Pract. 2017;71(11). 4. Golabi P, et al. Medicine (Baltimore). 2017;96(9):e5904. 5. Rawla P, et al. Contemp Oncol (Pozn). 2018;22(3):141-150. 6. Yoo JJ, et al. Sci Rep. 2023;13(1):14584. 7. Data on file. 0001. Fort Lee, NJ: Elevar Therapeutics; February 1, 2024. 8. Sadhu A, Blandy O. Decisions Resources Group. Disease Landscape & Forecast: Hepatocellular Carcinoma. Published November 20, 2020. 9. Zhu XD, et al. Genes Dis. 2020;7(3):328-335.

# uHCC Current Treatment Paradigm<sup>1</sup>



\*Per NCCN Guidelines for 1L uHCC, nivolumab and atezolizumab+bevacizumab are useful in certain circumstances (Child-Pugh Class B only).

References: 1. Leowattana W, et al. World J Gastroenterol. 2023;29(10):1551-1568. 2. Cheng A-I, et al. J Hepatol. 2022;76(4):862-873. 3. Finn RS, et al. N Engl J Med. 2020;382(20):1894-1905. 4. Abou-Alfa GK, et al. NEIM Evid. 2022;1(8):doi: 10.1056/EVIDoa2100070 5. Qin S, et al. Lancet. 2023;402(10408):1133-1146.

# **Complementary Mechanism of Actions (MOAs)** of Camrelizumab + Rivoceranib



Click to View the Camrelizumab MOA Video

### Camrelizumab

Camrelizumab is a **monoclonal antibody** that targets the programmed cell death protein 1 (PD-1) receptor.

In cancer cells, the programmed cell



death ligand 1 (PD-L1) protein binds to the PD-1 receptor, effectively allowing tumor cells to escape immunosurveillance. This is referred to as the PD-1/PD-L1 pathway. By targeting the PD-1 receptor, **camrelizumab blocks this binding** and allows T cells to start attacking tumor cells, boosting **immune response**.

### Rivoceranib

Rivoceranib is a **small-molecule tyrosine kinase inhibitor** (TKI) that works to **inhibit** the vascular endothelial growth factor receptor (VEGFR-2), a primary pathway for tumor angiogenesis (*see illustration*).

By inhibiting VEGFR-2, rivoceranib helps **restrict blood vessels** used in supplying nutrients to the tumor which leads to the **death of tumor cells** and **slows further cancer growth.** 

### **Tumor angiogenesis**



Click to view the Rivoceranib MOA Video

### **Camrelizumab + Rivoceranib**

Camrelizumab and rivoceranib, designated as a targeted combination therapy, is designed to **disrupt cancer cell** pathogenesis at **specific biological points and distinct pathways** that are essential to tumor development as outlined above, via a two-pronged approach. Camrelizumab **reinvigorates the body's immune response** by allowing T cells to attack malignant cancer cells,<sup>1</sup> while rivoceranib targets the VEGFR-2 pathway to **restrict the supply** of blood vessels and nutrients to the tumor.<sup>2,3</sup>

# **CARES-310** Phase 3 Trial Design

### International, Open-Label Phase 3 Trial<sup>1</sup>

#### **Key Eligibility Criteria**

- Unresectable or metastatic HCC
- BCLC Stage B (unsuitable for radical surgery and/or locoregional treatment) or C
- No prior systemic therapy
- ECOG PS 0 or 1
- Child-Pugh A
- At least one measurable lesion per RECIST v1.1



### *Treatment until loss of clinical benefits\* or intolerable toxicity*

#### **Stratification Factors**

- MVI and/or EHS (yes vs no)
- Geographical region (Asia vs non-Asia)
- Baseline serum AFP (<400 vs ≥400 ng/mL)

#### **Primary Endpoints**

- PFS<sup>+</sup>
- OS

#### **Key Secondary Endpoint**

• Objective response rate (ORR)<sup>†</sup>

AFP=alpha-fetoprotein; BCLC=Barcelona Clinic Lever Cancer; BIRC=blinded independent review committee; CARES-310=Camrelizumab plus rivoceranib vs sorafenib as first-line therapy for unresectable hepatocellular carcinoma; ECOG PS=Eastern Cooperative Oncology Group performance status; EHS=extrahepatic spread; HCC=hepatocellular carcinoma; IV=intravenous; MVI=macrovascular invasion; OS=overall survival; PFS=progression-free survival; RECIST=Response Evaluation Criteria in Solid Tumors.

\*Treatment beyond progression allowed if there was evidence of clinical benefits per investigator. <sup>†</sup>By BIRC per RECIST v1.1.

Reference: 1. Qin S, et al. Lancet. 2023;402(10408):1133-1146.

### Camrelizumab + Rivoceranib Demonstrated Significant mOS in a Phase 3 Study vs Sorafenib as First-Line Treatment for uHCC



Please note that head-to-head studies were not conducted between these products. These data are for information purposes only and no comparative claims of non-inferiority or superiority in terms of efficacy or safety are implied or intended.

**References: 1.** NEXAVAR. Prescribing Information. Whippany, NJ: Bayer HealthCare Pharmaceuticals Inc; July 2020. **2.** LENVIMA [package insert]. Nutley, NJ: Eisai Inc. **3.** IMFINZI (durvalumab) [Prescribing Information]. Wilmington, DE: AstraZeneca Pharmaceuticals LP; 2022. **4.** IMJUDO (tremelimumab-actl) [Prescribing Information]. Wilmington, DE: AstraZeneca Pharmaceuticals LP; 2022. **4.** IMJUDO (tremelimumab-actl) [Prescribing Information]. Wilmington, DE: AstraZeneca Pharmaceuticals LP; 2022. **4.** IMJUDO (tremelimumab-actl) [Prescribing Information]. Wilmington, DE: AstraZeneca Pharmaceuticals LP; 2022. **5.** Cheng AL, et al. *J Hepatol*. 2022;76(4):862-873. **6.** Qin S, et al. *Lancet*. 2023;402(10408):1133-1146. doi:10.1016/S0140-6736(23)00961-3

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### The Combination of Camrelizumab + Rivoceranib Has the Potential to Be a Best-in-Class Treatment Option in uHCC Based on Measurable, Clinically Meaningful Data Points

mOS	22.1 months <sup>1</sup>
Relative Risk Reduction (PFS)	<b>48%1</b> HR, 0.52 (95% CI; 0.41-0.65)
Stable Disease	52.9% <sup>1</sup>
Progressive Disease	<b>16.2%</b> <sup>1</sup>
Viral and Non-Viral Etiology	55% and 29% reduction in the risk for mortality for patients with HCV and non-viral etiology, respectively <sup>1</sup>
Albumin-Bilirubin (ALBI) Impact Post-Hoc analysis	<ul> <li>No significant change over time to ALBI Score<sup>1</sup></li> <li>Similar mOS for patients with Grade 1 or Grade 2 ALBI Score<sup>2</sup></li> </ul>
Discontinuation Rate	Lowest: 3.7% <sup>1</sup>
Grade 3-4 Hemorrhage	3.3% rate <sup>1</sup>
Half-life (mean, at steady state)	Rivoceranib: 7.0 hours to 16.3 hours <sup>3</sup> Allows for rapid withdrawal of VEGFR-2 blockade Camrelizumab: 17 days <sup>4</sup>

# Patients Treated With Camrelizumab + Rivoceranib Performed Better Compared With Sorafenib Monotherapy<sup>1</sup>

		No. of events/r	no. of patients					Uns	tratified HR
		Camrelizumab+ rivoceranib	Sorafenib					(95%	% CI)*
Geographic region	Asia	95/225	125/224					0.66	6 (0.51-0.86)
	Non-Asia	16/47	26/47					0.55	6 (0.29-1.02)
ECOG PS	0	47/120	60/116		<b>\</b>	_		0.67	(0.46-0.98)
	1	64/152	91/155		<b>—•</b> —			0.61	(0.45-0.85)
AFP at baseline	<400 ng/mL	58/176	82/171			-		0.64	(0.46-0.90)
	≥400 ng/mL	53/96	69/100			-		0.63	8 (0.44-0.90)
BCLC stage	В	9/38	13/40					0.73	8 (0.31-1.72)
	С	102/234	138/231		-•			0.62	2 (0.48-0.80)
MVI and/or EHS	Yes	86/200	121/200		<b>—•</b> —			0.59	0 (0.44-0.77)
	No	25/72	30/71		•			0.81	(0.48-1.37)
MVI	Yes	18/40	35/52			_		0.56	6 (0.32-0.99)
	No	93/232	116/219					0.68	8 (0.51-0.89)
EHS	Yes	73/175	110/180		<b>—●—</b>			0.56	6 (0.41-0.75)
	No	38/97	41/91		•			0.84	(0.54-1.30)
Etiology	HBV	90/208	113/197		<b></b>			0.66	6 (0.50-0.87)
	HCV	6/22	16/29	_	•			0.45	5 (0.18-1.16)
	Non-viral	15/42	22/45		<b>—</b>			0.71	(0.37-1.36)
Previous local therapy	Yes	67/161	79/150		<b>_</b> _			0.72	(0.52-0.99)
	No	44/111	72/121		<b>—•</b> —			0.57	(0.39-0.83)
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 17.2% of total study patient population were Western

 Hazard ratios (HRs) of OS favored camrelizumab + rivoceranib over sorafenib in the majority of subgroups

# Camrelizumab + Rivoceranib Is Highly Effective for Tumor Reduction

### Best change from baseline in sum of diameters of target lesion<sup>1,2</sup>



**35.2%** of patients had at least a 30% reduction in lesion diameter

72.8% of patients had a reduction in lesion diameter

# Well-Tolerated Safety Profile With Low TRAE-Related Discontinuations<sup>1,2</sup>

### Summary

### **TRAEs with incidence of ≥20%**<sup>§</sup>

	Camrelizumab + rivoceranib (N=272)	Sorafenib (N=269)	Preferred term	Camreliz rivocerani	zumab + b (N=272)	Sorafenib	(N=269)
Modian expective of treatment (IOD) me				Any grade	Grade ≥3	Any grade	Grade ≥3
			Hypertension	189 (69.5)	102 (37.5)	116 (43.1)	40 (14.9)
Camrelizumab	6.9 (3.6-13.4)	-	AST increased	147 (54.0)	45 (16.5)	99 (36.8)	14 (5.2)
Rivoceranib/sorafenib	6.5 (3.4-11.9)	3.8 (1.9-7.4)	Proteinuria	134 (49.3)	16 (5.9)	72 (26.8)	5 (1.9)
	265 (07.4)	240 (02.6)	ALT increased	127 (46.7)	35 (12.9)	80 (29.7)	8 (3.0)
ANY IKAL	203 (97.4)	249 (92.0)	Platelet count decreased	126 (46.3)	32 (11.8)	89 (33.1)	4 (1.5)
Grade 3/4	219 (80.5)	140 (52.0)	Blood bilirubin increased	116 (42.6)	24 (8.8)	75 (27.9)	4 (1.5)
Grade 5	1 (0.4) <sup>+</sup>	1 (0.4)‡	PPE syndrome	102 (37.5)	33 (12.1)	163 (60.6)	41 (15.2)
Serious TRAF	66 (24.3)	16 (5.9)	Diarrhea	83 (30.5)	6 (2.2)	105 (39.0)	14 (5.2)
	00 (2110)	10 (010)	RCEP	79 (29.0)	7 (2.6)	0	0
TRAEs leading to dose modification or interruption of any treatment component	128 (47)	87 (32)	Neutrophil count decreased	73 (26.8)	16 (5.9)	27 (10.0)	3 (1.1)
TRAEs leading to discontinuation of any treatment component	66 (24.3)	12 (4.5)	White blood cell count decreased	73 (26.8)	7 (2.6)	38 (14.1)	3 (1.1)
TRAEs leading to discontinuation of all	10 (2 7)	12 (4 E)	GGT increased	66 (24.3)	27 (9.9)	49 (18.2)	20 (7.4)
treatment components	10 (3.7)	- 3.8 (1.9-7.4) 249 (92.6) 140 (52.0) 1 (0.4) <sup>‡</sup> 16 (5.9) 87 (32) 12 (4.5) -	Hypothyroidism	58 (21.3)	0	16 (5.9)	0

Data are N (%) or otherwise indicated. \*Causality to treatment was determined by the investigator.  $\dagger$ Multiple organ dysfunction syndrome.  $\ddagger$ Respiratory failure and circulatory collapse. Data cutoff: Feb. 8, 2022. IQR=interquartile range; mo=months; TRAE=treatment-related adverse event.

Data are N (%). §TRAEs of any grade occurring in  $\geq$ 20% or of grade  $\geq$ 3 occurring in  $\geq$ 5% of patients in either group are listed. Data cutoff: Feb. 8, 2022. ALT=alanine aminotransferase; AST=aspartate aminotransferase; GGT=gamma-glutamyl transferase; PPE=palmar-plantar erythrodysesthesia; RCEP=reactive capillary endothelial proliferation; TRAE=treatment-related adverse event.

# **Camrelizumab + Rivoceranib Patient-Reported Outcomes<sup>1</sup>**

### Camrelizumab + rivoceranib demonstrated statistically significant differences in patient-reported outcomes vs sorafenib:

- Less deterioration in global health status/quality of life (P=0.012)
- Decreased pain (P=0.045)
- Decreased fatigue (P=0.007)

LS Mean (95% CI) for Change from Baseline in Pain Over Time By Treatment Arm<sup>+</sup>



LS Mean (95% CI) for Change from Baseline in Global Health Status/Quality of Life Over Time By Treatment Arm\*



LS Mean (95% CI) for Change from Baseline in Fatique Over Time By Treatment Arm<sup>+</sup>



CI=confidence interval; EORTC=European Organization for Research and Treatment of Cancer; LS=least squares; QLQ-C30=Quality-of-life Questionnaire Core 30. \*Measured by EORTC QLQ-C30 using data up to week 57. An increase in scores from baseline indicates improvement. †Measured by EORTC QLQ-C30 using data up to week 57. A decrease in scores from baseline indicates improvement. **Reference: 1.** Data on file. 0004. Fort Lee, NJ: Elevar Therapeutics; February 12, 2024.

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# Subgroup Analysis: CARES-310

### **OS SUBGROUP ANALYSIS<sup>1</sup>**

		No. of events/no	o. of patients		Unstratified HR
		Camrelizumab+ rivoceranib	Sorafenib	-	(95% CI)*
Geographic region	Asia	95/225	125/224	_ <b>—</b>	0.66 (0.51-0.86)
	Non-Asia	16/47	26/47		0.55 (0.29-1.02)
ECOG PS	0	47/120	60/116	<b>_</b>	0.67 (0.46-0.98)
	1	64/152	91/155	_ <b>—</b>	0.61 (0.45-0.85)
AFP at baseline	<400 ng/mL	58/176	82/171	<b></b>	0.64 (0.46-0.90)
	≥400 ng/mL	53/96	69/100	_ <b>—</b>	0.63 (0.44-0.90)
BCLC stage	В	9/38	13/40		 0.73 (0.31-1.72)
	С	102/234	138/231	<b>_</b>	0.62 (0.48-0.80)
MVI and/or EHS	Yes	86/200	121/200	<b>_</b>	0.59 (0.44-0.77)
	No	25/72	30/71	<b>_</b>	 0.81 (0.48-1.37)
MVI	Yes	18/40	35/52	<b>_</b>	0.56 (0.32-0.99)
	No	93/232	116/219	_ <b>e</b>	0.68 (0.51-0.89)
EHS	Yes	73/175	110/180	_ <b>—</b>	0.56 (0.41-0.75)
	No	38/97	41/91		 0.84 (0.54-1.30)
Etiology	HBV	90/208	113/197	_ <b>_</b>	0.66 (0.50-0.87)
	HCV	6/22	16/29		 0.45 (0.18-1.16)
	Non-viral	15/42	22/45		 0.71 (0.37-1.36)
Previous local therapy	Yes	67/161	79/150		0.72 (0.52-0.99)
	No	44/111	72/121	_ <b>_</b>	0.57 (0.39-0.83)
					 ,

0.0 0.5 1.0 1.5 2.0

### CARES-310





OS outcomes favored the cam + rivo arm regardless of baseline ALBI scores

- HR 0.62 (0.47-0.83) for ALBI grade 1
- HR 0.62 (0.4-1.0) for ALBI grade 2

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# Subgroup Analysis: IMbrave150 (Roche)

#### **OS SUBGROUP ANALYSIS<sup>1</sup>**



#### IMbrave150<sup>2</sup>



No OS benefit seen in ALBI grade 2 (or mALBI 2a and mALBI 2b) patients treated with atezo-bev vs sorafenib:

- ALBI 1: HR 0.50 (0.35-0.72) mALBI 2a: HR 0.97 (0.59-1.59) • ALBI 2: HR 0.92 (0.66-1.29) • mALBI 2b: HR 0.85 (0.54-1.34)

# Subgroup Analysis: HIMALAYA (AstraZeneca)

### **OS SUBGROUP ANALYSIS<sup>1</sup>**

A STRIDE versus Sorafenib			Hazard ratio
	STRIDE	Sorafenib	(95% CI)
All patients	262/393 (66.7%)	293/389 (75.3%)	0.78 (0.66-0.92)
Sex: male	211/327 (64.5%)	255/337 (75.7%)	0.73 (0.61-0.88)
Sex: female	51/66 (77.3%)	38/52 (73.1%)	1.02 (0.67-1.56)
Age at randomization: <65 yr	134/195 (68.7%)	146/195 (74.9%)	0.82 (0.65-1.04)
Age at randomization: >>65 yr	128/198 (64.6%)	147/194 (75.8%)	0.73 (0.58-0.93)
PD-L1 expression: positive	101/148 (68.2%)	110/148 (74.3%)	0.85 (0.65-1.11)
PD-L1 expression: negative	128/189 (67.7%)	138/181 (76.2%)	0.83 (0.65-1.05)
Etiology of liver disease: HBV	82/122 (67.2%)	98/119 (82.4%)	0.64 (0.48–0.86)
Etiology of liver disease: HCV	73/110 (66.4%)	64/104 (61.5%)	1.06 (0.76-1.49)
Etiology of liver disease: nonviral	107/161 (66.5%)	131/166 (78.9%)	0.74 (0.57-0.95)
ECOG performance status at baseline: 0	147/244 (60.2%)	168/241 (69.7%)	0.79 (0.63-0.98)
ECOG performance status at baseline: 1	114/148 (77.0%)	124/147 (84.4%)	0.74 (0.57-0.95)
Macrovascular invasion: yes	78/103 (75.7%)	83/100 (83.0%)	0.78 (0.57-1.07)
Macrovascular invasion: no	184/290 (63.4%)	210/289 (72.7%)	0.77 (0.63-0.93)
Extrahepatic spread: yes	146/209 (69.9%)	160/203 (78.8%)	0.67 (0.53-0.84)
Extrahepatic spread: no	114/182 (62.6%)	133/185 (71.9%)	0.87 (0.67-1.11)
Macrovascular invasion=yes and/or extrahepatic spread=yes	186/263 (70.7%)	196/251 (78.1%)	0.73 (0.59-0.89)
Macrovascular invasion=no and extrahepatic spread=no	74/128 (57.8%)	97/137 (70.8%)	0.79 (0.58-1.06)
Region: Asia (except Japan)	103/156 (66.0%)	123/156 (78.8%)	0.71 (0.54-0.92)
Region: Rest of world (includes Japan)	159/237 (67.1%)	170/233 (73.0%)	0.82 (0.66-1.02)
Alpha-fetoprotein at baseline: <400 ng/ml	109/167 (65.3%)	130/182 (71.4%)	0.82 (0.63-1.05)
Alpha-fetoprotein at baseline: ≥400 ng/ml	70/98 (71.4%)	60/71 (84.5%)	0.64 (0.45-0.91)
BCLC score: B	44/77 (57.1%)	44/66 (66.7%)	0.87 (0.57-1.33)
	010/016 (60 00/)	240/222 (77 104)	0.76 (0.63-0.91)

### ALBI SUBGROUP ANALYSIS<sup>1,2</sup>

### ALBI grade 1:

- mOS was 23.4 months with STRIDE vs 19.02 months with sorafenib
- OS HR (95% CIs) was 0.79 (0.62-1.01) for STRIDE vs sorafenib, consistent with full analysis set (0.78 [96% CI, 0.65-0.93])

### ALBI grade 2/3:

- mOS was 11.3 months with STRIDE vs
   9.7 months with sorafenib
- OS HR (95% CIs) was 0.83 (0.65-1.05) for STRIDE vs sorafenib

Hazard Ratio (95% Cl

# **Commercial Opportunity and Strategy**

# **Top Unmet Needs in uHCC Today**



### Low Overall Survival<sup>1-3</sup>

With a 5Y overall survival (OS) rate at 18% in the US, **uHCC has one** of the lowest OS rates of all cancers. Many patients when diagnosed with HCC see this as "depressing," "living a nightmare," and a "death sentence." Improved HCC treatment to extend life is needed.



### Quality of Life<sup>3,4</sup>

Severe liver dysfunction because of HCC conjoined with treatment **side effects** (weight loss, fatigue, nausea, etc.) **make quality of life poor for patients.** HCC also can take a severe toll on caregivers and their quality of life—a need for safer and more efficacious therapy is needed.



### **Treatment for Patients With Impaired Liver Function<sup>5</sup>**

Improved treatment for those with higher levels of liver dysfunction (as determined by Child-Pugh or ALBI) is needed, as these **patients are the most underserved**, leading some patients to opt for hospice instead of treatment.



### **Optimal Treatment Selection<sup>6</sup>**

With more novel HCC treatments available (and more clinical trials ongoing), there is an increasing need to understand how to **best optimize treatment choice by patient type**, including factors like biomarkers, liver function, disease etiology, etc., both in 1L and sequencing across lines of therapy.

References: 1. Decisions Resources Group – HCC Survival Rate (Accessed 9/30/23). 2. Vogel A, et al. Lancet. 2022;400(10360):1345-1362. 3. Data on file. 0003. Fort Lee, NJ: Elevar Therapeutics; February 1, 2024. 4. American Cancer Society. Treating liver cancer. https://www.cancer.org/cancer/types/liver-cancer/treating.html Accessed January 31, 2024. 5. NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) for Hepatocellular Carcinoma V.2.2023. 6. Rai V, et al. Hepatobiliary Surg Nutr. 2022;11(4):629-631.

**HCC Unmet** 

Needs

# **Elevar Is Well Positioned to Treat the US uHCC Patient Population**



# **Experienced Oncology Sales Force With Ability to Expand**



## **Appendix** For more information, please visit https://elevartherapeutics.com/

# **Corporate Timeline**

