UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 8-K

CURRENT REPORT Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): August 11, 2022

LIANBIO

(Exact name of registrant as specified in its charter)

Cayman Islands (State or other jurisdiction of incorporation) 001-40947

98-1594670 (IRS Employer Identification No.)

103 Carnegie Center Drive, Suite 309 Princeton, NJ (Address of principal executive offices)

(Commission File Number)

08540 (Zip Code)

(Registrant's telephone number, including area code): (609) 486-2308

Not Applicable (Former name or former address, if changed since last report)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions (see General Instruction A.2. below):

0 Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)

Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12) 0

о Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c)) 0

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
American depositary shares, each representing 1 ordinary share, \$0.000017100448 par value per share	LIAN	The Nasdaq Global Market

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

Emerging growth company x

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. \boldsymbol{x}

Item 2.02 Results of Operations and Financial Condition.

On August 11, 2022, LianBio (the "Company") issued a press release announcing its financial results for the fiscal quarter ended June 30, 2022. A copy of the press release is furnished as Exhibit 99.1 to this Current Report on Form 8-К.

Item 7.01 Regulation FD Disclosure.

On August 11, 2022, the Company posted an updated corporate presentation to its website. A copy of the corporate presentation is furnished as Exhibit 99.2 to this Current Report on Form 8-K. The Company undertakes no obligation to update, supplement or amend the materials attached hereto as Exhibit 99.2.

The information contained in this Current Report on Form 8-K, including Exhibit 99.1 and Exhibit 99.2 attached hereto, is being furnished and shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), and shall not be deemed incorporated by reference in any filing under the Securities Act of 1933, as amended, or the Exchange Act, except as expressly set forth by specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits

Exhibit No.

No.	Description
99.1	Press release issued by LianBio, dated August 11, 2022
99.2	LianBio corporate presentation as of August 2022
104	Cover Page Interactive Data File (embedded within the Inline XBRI, document)

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

LIANBIO

By:

/s/ Yizhe Wang Yizhe Wang Chief Executive Officer

Date: August 11, 2022



LianBio Reports Second Quarter 2022 Financial Results and Provides Corporate Update

• Completed enrollment in China Phase 3 trial of mavacamten; topline data expected mid-2023

Submitted mavacamten New Drug Application (NDA) in Singapore

Submitted infigratinib NDA in Hong Kong

Three additional registration-enabling programs to begin in China by year-end 2022

• Cash balance of \$349.4 million at the end of second quarter 2022 with runway into the second half of 2024

Shanghai and Princeton, N.J., August 11, 2022 – LianBio (Nasdaq: LIAN), a biotechnology company dedicated to bringing innovative medicines to patients in China and other major Asian markets, today reported financial results for the second quarter ended June 30, 2022 and provided a corporate update.

"In the second quarter, LianBio achieved meaningful milestones that we believe serve as a testament to our strength in navigating the complex regulatory environments in Greater China and Asia," said Yizhe Wang, Ph.D., Chief Executive Officer of LianBio. "We continue to grow our organization, bringing on key new team members across clinical, medical affairs, quality and commercial functions. Despite clinical site and agency disruptions due to COVID-19 lockdowns in Shanghai and other cities in China, LianBio continued to execute our key clinical development and regulatory priorities, completing enrollment in the Phase 3 EXPLORER-CN trial of mavacamten and submitting marketing applications for both mavacamten and infigratinib in Asia Pacific territories. We believe the enthusiasm for mavacamten's potential as a treatment for obstructive hypertrophic cardiomyopathy (oHCM) among both the clinical community and the world's largest oHCM population is high, and we expect to report topline results from EXPLORER-CN in mid-2023. As we work to complete the EXPLORER-CN study over the coming months to support registration in China, we turn our focus to potential launch and preparations for LianBio's next phase of evolution as a commercial stage company. We remain on track to initiate three additional registration-enabling clinical programs in China this year, solidifying our position as a key partner in cross-border drug development. I am continually proud of our global team's efforts to bring innovative medicines to patients in Asia."

Recent Business Highlights and Clinical Development Updates

Mavacamten progress continues in Asia with enrollment completed in China Phase 3 trial and New Drug Application submitted in Singapore

- In May 2022, LianBio submitted an NDA to the Singapore Health Sciences Authority for mavacamten for the treatment of adults with symptomatic New York Heart Association Class II-III obstructive hypertrophic cardiomyopathy (oHCM). The submission was based on the U.S. Food and Drug Administration (FDA) approval of mavacamten.
- In August 2022, enrollment was completed in the Phase 3 EXPLORER-CN clinical trial of mavacamten in Chinese patients with oHCM.

Infigratinib New Drug Application in 2nd line cholangiocarcinoma submitted in Hong Kong

 In July 2022, LianBio submitted an NDA to the Department of Health, the Hong Kong Special Administrative Region, China, for infigratinib for the treatment of adults with previously treated, unresectable, locally advanced or metastatic cholangiocarcinoma with a fibroblast growth factor receptor 2 (FGFR2) fusion or other rearrangement. The submission was based on the FDA approval of infigratinib.

Development partner ReViral Ltd. acquired by Pfizer Inc.

• In June 2022, Pfizer completed its acquisition of LianBio's development partner ReViral and its respiratory syncytial virus therapeutic candidates, including sisunatovir. LianBio holds development and commercial rights to sisunatovir in mainland China, Hong Kong, Macau and Singapore.

LYR-210 clinical development program refined

LianBio plans to conduct a Phase 3 China standalone trial to support regulatory approval in China, leveraging the results of development partner Lyra's ongoing Phase 3 trial, which is expected to complete enrollment in mid-2023.

Development partner Tarsus advances TP-03 into clinical trial in second indication

• In August 2022, LianBio's development partner Tarsus initiated a Phase 2a clinical trial of TP-03 in patients with meibomian gland disease (MGD).

Development partner Landos Biopharma reports data from NX-13 program

• In August 2022, Landos announced topline results from a Phase 1b clinical trial of NX-13 demonstrating NX-13 was well tolerated. Based on these data, Landos plans to initiate a Phase 2 clinical trial to evaluate the safety, efficacy and optimal dosing of NX-13 in ulcerative colitis patients.

Business is well-positioned to achieve anticipated milestones

• Current cash runway is projected to extend into the second half of 2024.

Key Milestones Anticipated in 2022 and 2023

Mavacamten

LianBio expects to report topline data from the Phase 3 EXPLORER-CN trial of mavacamten in Chinese patients with symptomatic oHCM in mid-2023.

TP-03

• LianBio expects to initiate a Phase 3 study in Chinese patients with Demodex blepharitis in the second half of 2022. LianBio expects this study will support registration of TP-03 in China.

NBTXR3 • L

• LianBio expects to begin dosing patients in Nanobiotix's global Phase 3 NANORAY-312 clinical trial of NBTXR3 for the treatment of locally advanced head and neck squamous cell carcinoma in elderly patients ineligible for cisplatin in the second half of 2022. LianBio expects this study will support registration of NBTXR3 in China and other LianBio licensed territories in Asia.

Infigratinib

 Enrollment is ongoing in LianBio's Phase 2a clinical trial of infigratinib in locally advanced or metastatic gastric cancer or gastroesophageal junction adenocarcinoma with FGFR2 gene amplification and other advanced solid tumors with FGFR genomic alterations. • LianBio expects to begin dosing Chinese patients in Helsinn's ongoing global pivotal Phase 3 PROOF-301 clinical trial of infigratinib in first-line cholangiocarcinoma (CCA) patients with FGFR2 gene fusions/translocations in the second half of 2022.

BBP-398

- LianBio expects to initiate a Phase 1 monotherapy clinical trial of BBP-398 in advanced solid tumors in the fourth quarter of 2022.
- LianBio also expects to initiate a Phase 1 clinical trial of BBP-398 in combination with an EGFR-inhibitor in non-small cell lung cancer in the first half of 2023.

Second Quarter 2022 Financial Results

Research & Development Expenses

Research and development expenses were \$28.6 million for the second quarter of 2022 compared to \$93.0 million for the second quarter of 2021, and \$40.9 million for the six month period ended June 30, 2022 compared to \$146.4 million for the six month period ended June 30, 2021. The decrease was primarily attributable to increased milestone payments in 2021, and was offset by higher development activities to support clinical trials and personnel-related expenses in 2022.

General & Administrative Expenses

General and administrative expenses were \$14.6 million for the second quarter of 2022 compared to \$6.5 million for the second quarter of 2021, and \$30.6 million for the six month period ended June 30, 2022 compared to \$13.6 million for the six month period ended June 30, 2021. The increase was primarily attributable to increases in payroll and personnel-related expenses (including share-based compensation expense) for increased employee headcount and higher expense for legal, consulting and accounting services.

Net Loss

Net loss was \$42.4 million for the second quarter of 2022 compared to net loss of \$100.4 million for the second quarter of 2021, and \$70.1 million for the six month period ended June 30, 2022 compared to \$162.0 million for the six month period ended June 30, 2021.

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Cash Balance

Cash, cash equivalents, marketable securities and restricted cash at June 30, 2022 totaled \$349.4 million compared to \$403.2 million as of December 31, 2021. LianBio projects its current cash, cash equivalents, marketable securities, and restricted cash will be sufficient to fund its current operating plan into the second half of 2024.

About LianBio

LianBio is a cross-border biotechnology company on a mission to bring transformative medicines to patients in China and other Asian markets. Through partnerships with highly innovative biopharmaceutical companies around the world, LianBio is advancing a diversified portfolio of clinically validated product candidates with the potential to drive new standards of care across cardiovascular, oncology, ophthalmology, inflammatory disease and respiratory indications. LianBio is establishing an international infrastructure to position the company as a partner of choice with a platform to provide access to China and other Asian markets. For additional information, please visit the company's website at www.lianbio.com.

Cautionary Note Regarding Forward-Looking Statements

Statements in this press release about future expectations, plans and prospects, as well as any other statements regarding matters that are not historical facts, may constitute forward-looking statements. The words "anticipate," "plan," " believe, " "continue," "expect," "potential," "project," and similar expressions are intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words. Forward-looking statements in this press release include, but are not limited to, statements concerning the Company's plans and expectations with respect to the initiation and completion of its clinical trials, the advancement of its pipeline of therapeutic candidates, the continued growth or its organization, its ability to bring transformative medicines to patients across Asia, its ability to navigate complex regulatory environments in Greater China and Asia, the Company's plans and expectations with respect to preparation for potential commercialization and product launch, and the timeline through which it expects to be able to fund its operating expenses and capital expenditure requirements, as well as statements regarding our partners' announced plans and expectations with respect to their planned product development activities, preclinical studies and clinical trials. Actual results may differ materially from those indicated by such forward-looking statements as a result of various important factors, including: the Company's ability to successfully initiate and conduct its planned clinical trials and complete such clinical trials and admine regulatory enproval and maximize patient reach for its product candidates; the Company's ability to identify new product candidates and successfully acquire such product candidates from third parties; competition from other biotechnology and pharmaceutical companies; general market conditions; the impact of changing laws and regulations and those risks and uncertainties described in LianBio's filings with the U.S. Securities an

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LianBio Consolidated Balance Sheets (In thousands, except share and per share amounts) (Unaudited)

	June 30, 2022	December 31, 2021
Assets		
Current assets:		
Cash and cash equivalents	\$ 134,33	4 \$ 228,182
Marketable securities	194,90	5 155,067
Prepaid expenses and other current assets	7,14	10,354
Other receivable	7,20	6,044
Total current assets	343,70	399,647
Restricted cash, non-current	20,07	20,000
Property and equipment, net	2,99	1,882
Operating lease right-of-use assets	5,00	4,763
Other non-current assets	3	9 51
Total assets	\$ 371,81	3 \$ 426,343
Liabilities and Shareholders' Equity		
Current liabilities:		
Accounts payable	\$ 1,23	4 \$ 3,231
Accrued expenses	17,93	9,976
Current portion of operating lease liabilities	1,80	3 1,125
Other current liabilities	73	8 760
Total current liabilities	21,74	15,092
Operating lease liabilities	3,60	3,709
Other liabilities	20	8 206
Nonrefundable research deposit	20,00	0 20,000
Total liabilities	\$ 45,61	5 \$ 39,007
Commitments and contingencies (Note 8)		
Ordinary shares, \$0.000017100448 par value. Authorized 2,923,900,005 shares as of June 30, 2022; 108,353,831 shares issued and outstanding at June 30, 2022; Authorized 2,923,900,005 shares as of December 31, 2021; 107,275,458 shares issued and outstanding at December 31, 2021		2 2
Additional paid-in capital	724,17	713,269
Accumulated other comprehensive (loss) income	(1,40	2) 526
Accumulated deficit	(430,35	2) (360,235)
Total LianBio shareholders' equity	292,42	4 353,562
Non-controlling interest	33,77	4 33,774
Total shareholders' equity	326,19	387,336
Total liabilities and shareholders' equity	\$ 371,81	3 \$ 426,343

LianBio Statements of Operations and Comprehensive Loss (In thousands, except share and per share amounts) (Unaudited)

	Three Months	Ended June	e 30,	Six Months I	Ended June	: 30,
	2022		2021	2022		2021
Operating expenses:						
Research and development	\$ 28,591	\$	93,030	40,920	\$	146,383
General and administrative	14,551		6,461	30,639		13,607
Total operating expenses	 43,142		99,491	71,559		159,990
Loss from operations	(43,142)		(99,491)	(71,559)		(159,990)
Other income (expense):						
Interest income, net	553		106	833		139
Other income (expense), net	203		(68)	620		(192)
Net loss before income taxes	 (42,386)		(99,453)	(70,106)		(160,043)
Income taxes	5		975	11		1,950
Net loss	 (42,391)		(100,428)	(70,117)		(161,993)
Other comprehensive (loss) income:						
Foreign currency translation (loss) income, net of tax	(421)		122	(814)		130
Unrealized loss on marketable securities, net of tax	(291)		_	(1,114)		_
Comprehensive loss	\$ (43,103)	\$	(100,306)	\$ (72,045)	\$	(161,863)
Net loss per share attributable to ordinary shareholders, basic and diluted	\$ (0.39)	\$	(4.90)	\$ (0.65)	\$	(7.91)
Weighted-average shares outstanding used in computing net loss per share attributable to ordinary shareholders, basic and diluted	107,922,501		20,477,338	107,600,767		20,477,338





The information herein contains statements about future expectations, plans and prospects for LianBio. All statements, other than statements of historical fact, included herein are forward-looking statements. Forward-looking statements are not statements of historical fact nor are they guarantees or assurances of future performance. Forward-looking statements are based on LianBio's expectations and assumptions and are subject to inherent uncertainties, risks and changes in circumstances that may cause actual results to materially and adversely differ from those set forth in or implied by such forward-looking statements, including these risks and uncertainties that are described under the heading "Risk Factors" in our Annual Report on Form 10-K for the year ended December 31, 2021, as well as discussions of potential risks, uncertainties and other important factors in our subsequent filings with the Securities and Exchange Commission. LianBio undertakes no obligation to publicly update or revise any forward-looking statements, whether as a result of new information, future events or otherwise. These forward-looking statements should not be relied upon as representing LianBio's views as of any date subsequent to the date hereof.

Certain information contained in this presentation relates to or is based on studies, publications, surveys and other data obtained from third party sources and LianBio's own internal estimates and research. While LianBio believes these third-party sources to be reliable as of the date of this presentation, it has not independently verified, and makes no representation as to the adequacy, fairness, accuracy or completeness of any information obtained from third party sources. In addition, the third party information included in this presentation may involve a number of assumptions, and there can be no guarantee as to the accuracy or reliability of such assumptions. Finally, while LianBio believes its own internal research is reliable, such research has not been verified by any independent source.

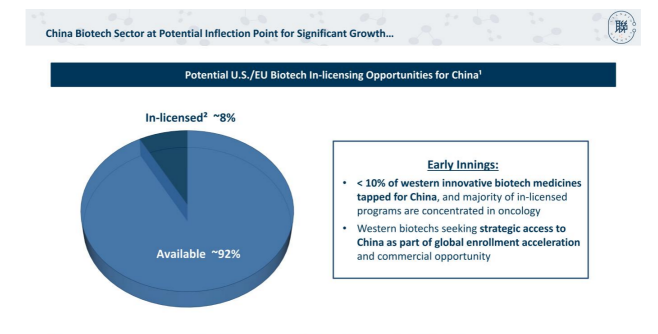


We are a global biopharmaceutical company dedicated to developing and commercializing paradigm-shifting medicines for patients with unmet medical needs in Greater China and other Asian markets

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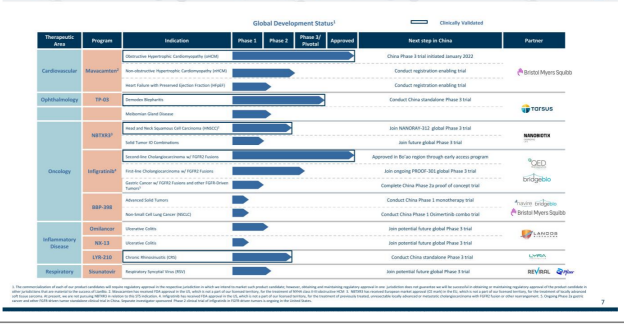




1. CapitalIQ. assumes only one opportunity per company based on 10,794 total US/EU biotech companies as of July 2021. 2.Based on 857 cross-border deals from 2015-2020 per ChinaBio.

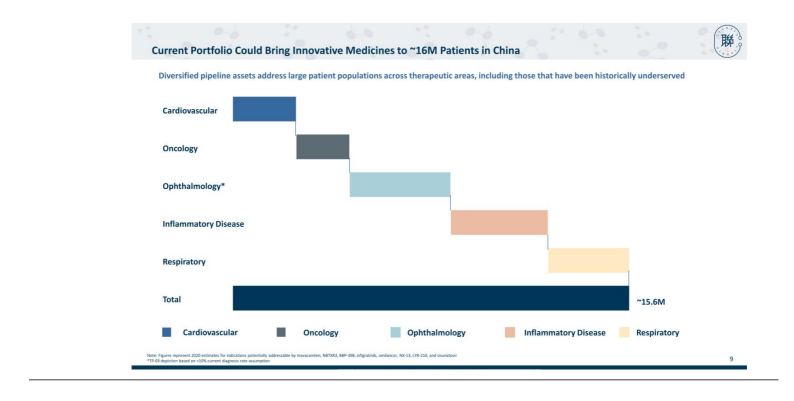


Pipeline of Innovative Medicines – 5 Clinically Validated Therapeutic Candidates



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	Design for the design of the
• BBP-398	Deepen existing TA franchises
LYR-210	Strategic multi-asset partnerships
Sisunatovir	
	Combination opportunities
	LYR-210 • Sisunatovir











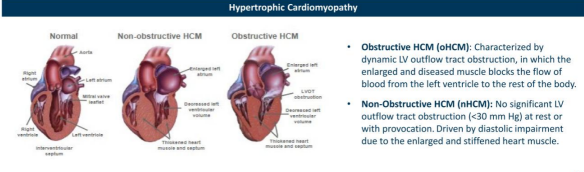
MYOKARDIA (III Bristol Myers Squibb"

Mavacamten for the Treatment of HCM and HFpEF

- Mavacamten is a myosin inhibitor that targets excessive contractility and impaired relaxation, myocardial energetics and compliance
- In development for the treatment of obstructive hypertrophic cardiomyopathy (oHCM), non-obstructive hypertrophic cardiomyopathy (nHCM) and heart failure with preserved ejection fraction (HFpEF)

China Opportunity

- 1.1M 2.8M HCM patients in China (67% oHCM / 33% nHCM)
- 3.7M HFpEF patients, 10-20% of whom may potentially be addressed by mavacamten



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Clinical Activity Demonstrated in oHCM and nHCM



Clinical Data Summary

oHCM:

- Phase 3 EXPLORER-HCM trial demonstrated patients on treatment experienced statistically significant and clinically meaningful improvements
 - Primary endpoint: Improvement of symptoms and functional capacity (improvement in NYHA class and peak VO2)
 - Well-tolerated; safety results were comparable to placebo; only 2% drop out rate

nHCM:

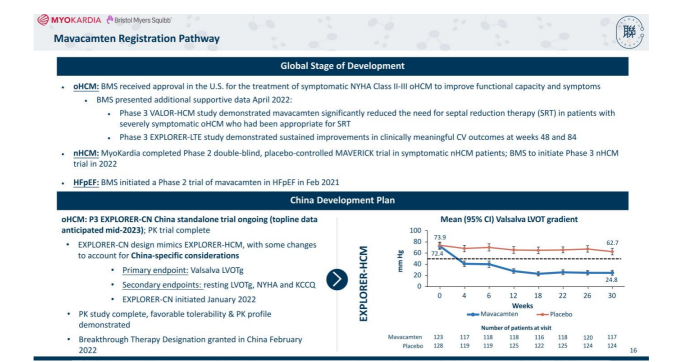
 Phase 2 MAVERICK-HCM trial demonstrated physiologic benefit with dose dependent reduction in serum levels of NT proBNP, with potentially greater benefit in more severe disease

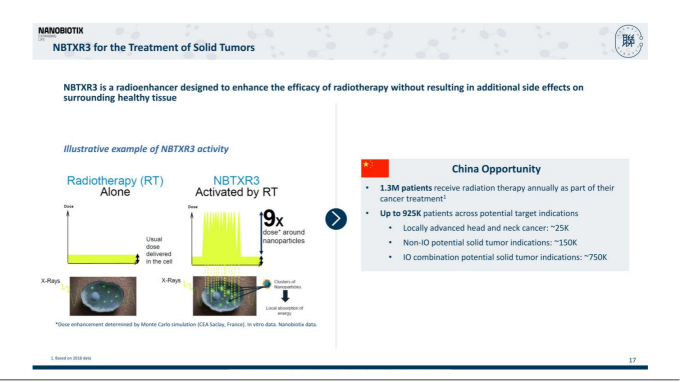
Source: Olivotto et al, Lancet 2020; Ho et al, J Am Coll Cardiol. 2020

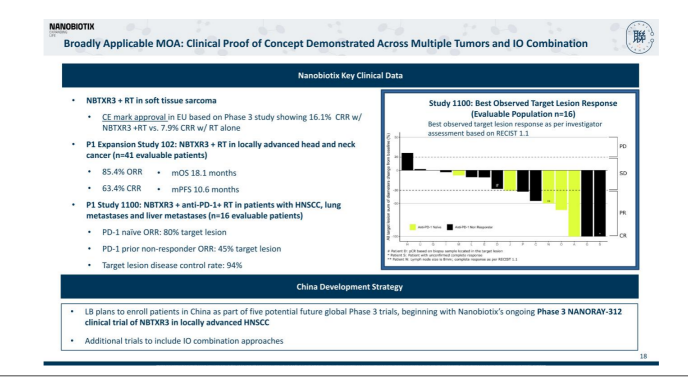
EXPLORER-HCM

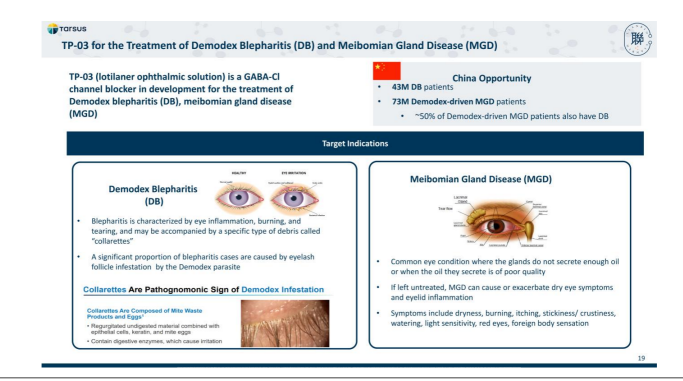
Change from Baseline to Week 30			
Primary Endpoint			
Composite functional, n (%) EITHER ≥1.5 mI/Kg/min increase in pVO2 with ≥1 NYHA class improvement OR ≥3.0 mI/Kg/min increase in pVO2 with no worsening of NYHA class	45 (37%)	22 (17%)	0.0005
Secondary Endpoints			
Post-exercise LVOT peak gradient, mmHg, mean (SD)	-47 (40)	-10 (30)	<0.0001
Peak VO2, mL/kg/min, mean (SD)	1.4 (3.1)	-0.1 (3.0)	0.0006
NYHA improved ≥ 1 class, n (%)	80 (65%)	40 (31%)	<0.0001
KCCQ-CSS, mean (SD)	13.6 (14.4)	4.2 (13.7)	<0.0001
HCMSQ-SoB score, mean (SD)	-2.8 (2.7)	-0.9 (2.4)	<0.0001

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Tarsus

All Pre-Specified Primary and Secondary Endpoints were Met in Tarsus's Saturn-1 and Saturn-2 Pivotal Trials

Tarsus completed two successful pivotal trials with consistency across endpoints

	Saturn-1 ^{N=421} (Pivotal Phase 2b/3)	Saturn-2 ^{N=412} (Pivotal Phase 3)	Combined ^{N=833} Pivotal Data
Primary Endpoint: Complete Collarette Cure	44% vs. 7% (p<0.0001)	56% vs. 13% (p<0.0001)	50% vs. 10%
Clinically Meaningful Collarette Cure (Grade 0 or 1)	81% vs. 23% (p<0.0001)	89% vs. 33% (p<0.0001)	85% vs 28%
Mite Eradication	68% vs. 18% (p<0.0001)	52% vs 14% (p<0.0001)	60% vs 16%
Lid Erythema Cure	19% vs. 7% (p<0.0001)	31% vs. 9% (p<0.0001)	25% vs 8%

Approximately 90% of patients experienced a clinically meaningful benefit with respect to collarettes, collarette grade improvement and mites per lash

Development and Regulatory Status U.S. China • Tarsus has announced plans to submit NDA for TP-03 in DB to U.S. FDA in 2H 2022 • LianBio to conduct pivotal study to support regulatory approval in China, to be initiated 2H 2022 • Phase 2a MGD trial to be initiated 2H 2022 • PK cohort (n=12) • Pase 2a MGD trial to be initiated 2H 2022 • P3 China standalone trial (N=150, 1:1 randomization) • Co-primary endpoints: collarette cure (0-2 collarettes per eyelid) at day 43, mite eradication at day 43

Secondary endpoints: composite cure of collarette and erythema (0-2 collarettes per eyelid and grade 0 erythema) at day 43

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bridgebio [©]<u>GED</u> Infigratinib for the Treatment of FGFR-Driven Cancers

Infigratinib is an orally administered, ATP-competitive, FGFR1-3 tyrosine kinase inhibitor in development for the treatment of patients with FGFR-driven cancers

- QED received FDA approval of infigratinib for the treatment of patients with previously-treated, unresectable locally advanced or metastatic cholangiocarcinoma (CCA) harboring an FGFR2 fusion or rearrangement in May 2021
- Approval based on meaningful clinical activity demonstrated in Phase 2 trial in chemotherapy-refractory CCA patients with FGFR2 fusions
 - BICR cORR of 23.1% (95% CI 15.6 32.2) in 2nd and later line patients
 - BICR cORR of 34.0% in true 2nd line patients
 DOR of 5.0 mos (95% CI 3.7–9.3)
 - DOR of 5.0 mos (95% CI 3.7–9.3)
 Infigratinib administered as third
 - Infigratinib administered as third-and later-line treatment resulted in meaningful PFS and ORR benefit in patients with CCA and FGFR2 fusions ~7 mos
 - Current SoC (chemo) = ~3 mos PFS in 2L CCA

BICR- assessed objective response rate (ORR), % (95% CI)	23.1 (15.6-32.2)
≤1 previous line of therapy (n=50)	34.0
≥2 previous lines of therapy (n=58)	7.4
BICR-assessed best overall response	
Complete Response, n (%)	1 (1.1)
Partial Response, n (%)	24 (22)
Stable Disease, n (%)	66 (61)
Unconfirmed Complete or Partial Response	12 (11)
Progressive Disease, n (%)	11 (10)
Unknown, n (%)	6 (6)
BICR-assessed confirmed or unconfirmed response, % (95% CI)	34.3 (25.4 - 44.0)
BICR-assessed disease control rate, % (95% CI)	84.3 (76.0 - 90.6)
BICR-assessed median duration of response (IQR), months (95% CI)	5.0 (3.7 - 9.3)
BICR-assessed median PFS, months (95% CI)	7.3 (5.6 - 7.6)
Median OS, months (95% CI)	12.2 (10.7 -14.9)

QED's Development and Regulatory Status in the U.S.

- U.S. FDA approval in 2nd line CCA received May 2021
- Ongoing global Phase 3 PROOF-301 trial in 1st line CCA
- Ongoing global Phase 3 trial in urothelial carcinoma
- In Jan 2020 received Fast Track Designation for 1st line CCA
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bridgebio [©] ED Infigratinib Registration Pathway and China Opportunity

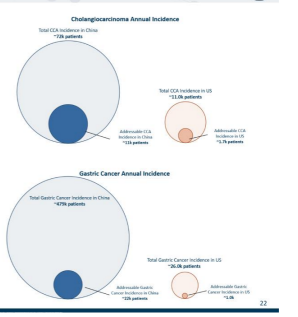


China Opportunity

- Estimated 72,000 patients diagnosed with CCA annually in • China vs. 11,000 diagnosed in U.S.
- Estimated 480,000 patients diagnosed with GC annually in China vs. 26,350 diagnosed in U.S. •

China Development Strategy and Regulatory Pathway

- LB will enroll patients in China as part of QED's ongoing global Phase 3 PROOF trial in first-line CCA
- LB initiated a Phase 2a proof of concept trial in China for . FGFR2-amplified gastric cancer and other solid tumors with FGFR alterations



Dridgebio • D Inavire • Si Inavire • Ir Image: simple s	 SP-398 (SHP2 inhibitor) for the treatment of MAPK pathway-driven solid tumors Differentiated profile with a shorter-half life, attractive PK/PD and clean tox SHP2 inhibitors have broad potential applications across a variety of tumors and are being developed as combination therapy R-210 (implantable drug matrix) for the treatment of chronic rhinosinusitis (CRS) with 3.4M medically refractory tients in China Implantable drug matrix designed to consistently and locally elute mometasone furoate (steroid) to inflamed mucosal sinus tissue for up to six months with a single administration for surgically naïve patients Clinically validated with Ph2 statistically significant symptom improvement vs. control at 16, 20 and 24 weeks
REVIRAL Sisu	itients in China Implantable drug matrix designed to consistently and locally elute mometasone furoate (steroid) to inflamed mucosal sinus tissue for up to six months with a single administration for surgically naïve patients
• P	sunatovir (fusion inhibitor) for the treatment of respiratory syncytial virus (RSV) No SAEs observed across ~200 patients treated to date; no cardiac toxicity observed to date, a key issue leading to failure of prior fusion inhibitors Potential applicability in high-risk patient segments including pediatric, elderly patients
• Or • Or • Ra	nilancor (LANCL2 agonist) for the treatment of IBD Dral, gut-restrictive mechanism (lack of systemic exposure) designed for a safe and convenient route of administration for reatment of moderate to severe IBD Rapidly growing IBD incident population in China -13 (NLRX1 agonist) for the treatment of IBD

Major Validating Milestones Highlight Strength of LianBio Business Development Engine

Partner	LianBio Partnership Date	Asset Milestone Post-Partnership
MYOKARDIA Mavacamten (^{Ill} ı Bristol Myers Squibb	Aug 2020	 Oct 2020: MyoKardia acquired by BMS for \$13.1B Apr 2022: BMS received U.S. FDA approval of mavacamten for patients with symptomatic oHCM
	Mar 2021	✓ Apr 2022: Reviral enters agreement to be <u>acquired by Pfizer for up to</u> <u>\$525M</u>
bridgebio Infigratinib	Oct 2019	 May 2021: QED received FDA approval of infigratinib for patients with previously treated cholangiocarcinoma Mar 2021 & 2022: Helsinn Group and QED enter into and expand infigratinib strategic collaboration
Tarsus TP-03	Mar 2021	 Jun 2021: <u>Positive pivotal results</u> in Tarsus's SATURN-1 trial (P2b/3 DE – all primary and secondary endpoints met May 2022: <u>Positive pivotal results</u> in Tarsus's SATURN-2 trial (P3 DB) – all primary and secondary endpoints met
bridgebio BBP-398 (^{III}) Bristol Myers Squibb	Oct 2019	 May 2022: BridgeBio and BMS enter into BBP-398 strategic collaboration

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A differentiated strategic collaboration that provides sourcing, development and commercial optionality

- Provides LianBio and partners optionality to access
 <u>Pfizer's established commercial infrastructure</u> with a highly compliant, secure commercial engine
- At LianBio's election and Pfizer's ROFN, we can jointly develop and commercialize certain LianBio products
- Companies are also working together to source, select and develop/register leading products for China
 - Pfizer will contribute up to \$70M of non-dilutive capital for in-licensing and co-development activities



Preferential access to an <u>innovative pipeline of more</u> <u>than 20 product development candidates</u>

- BridgeBio is developing transformative medicines to treat patients who suffer from genetic diseases and cancers with clear genetic drivers. BridgeBio is advancing a broad, innovative pipeline across rare disease, oncology, dermatology, and other indications
- LianBio already holds China rights to two of BridgeBio's oncology assets, infigratinib and BBP-398

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Targeting 3 Additional Registrational Trial Initiations and Multiple Catalysts by End of 2022

herapeutic Area	Program	Milestone / Catalyst	Anticipated Timing
		Initiate Phase 3 EXPLORER-CN clinical trial in patients with oHCM	✓ Jan 2022
		 Mavacamten granted BTD for oHCM in China 	✓ Feb 2022
Cardiovascular	Mavacamten	 U.S. FDA approval for the treatment of symptomatic oHCM (BMS) 	V April 2022
		 Completion of PK trial in China, demonstrating favorable safety, tolerability and PK profile 	🗸 May 2022
		 Complete enrollment in Phase 3 EXPLORER-CN clinical trial in patients with oHCM 	✓ Aug 2022
Ontwinding	70.00	 Saturn-2 pivotal trial readout (Tarsus) 	🗸 May 2022
Ophthalmology	TP-03	 Initiate Phase 3 clinical trial in patients with Demodex blepharitis in China 	H2 2022
		Global trial initiation of Phase 3 NANORAY-312 clinical trial in head and neck cancer (Nanobiotix)	🗸 Jan 2022
Oncology	NBTXR3	 Initiate China portion of Phase 3 NANORAY-312 clinical trial in patients with head and neck cancer 	H2 2022
Infigratinib	 Initiate China portion of Phase 3 PROOF-301 clinical trial in patients with first line cholangiocarcinoma 	H2 2022	

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We are a global biopharmaceutical company dedicated to developing and commercializing paradigm-shifting medicines for patients with unmet medical needs in Greater China and other Asian markets

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