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Genscript Biotech Corporation 金斯瑞生物科技股份有限公司*

(Incorporated in the Cayman Islands with limited liability)
(Stock Code: 1548)

VOLUNTARY ANNOUNCEMENT RESEARCH AND DEVELOPMENT UPDATE

This is a voluntary announcement made by GenScript Biotech Corporation (the "Company" and together with its subsidiaries collectively referred to as the "Group").

ESTABLISHMENT OF A NEW MANUFACTURING FACILITY IN EUROPE

The board (the "Board") of directors (the "Directors") of the Company is pleased to announce that, on 22 June 2021 (New York time), Legend Biotech Corporation ("Legend Biotech"), a non-wholly owned subsidiary of the Company, announced the establishment of a manufacturing facility in Belgium ("Belgium Manufacturing Facility"), as part of a joint investment with Janssen Pharmaceutical NV (Janssen), to expand global manufacturing capacity of innovative cellular therapies.

Legend Biotech has a collaboration and license agreement with Janssen Biotech, Inc. to develop and commercialize ciltacabtagene autoleucel (cilta-cel), an investigational CAR-T therapy currently under review by several health authorities around the world including the United States and Europe for the treatment of patients with relapsed and refractory multiple myeloma. For details of the collaboration and license agreement, please see the announcement of the Company dated 22 December 2017.

The Group believes that the new location in Belgium is an ideal choice for Legend Biotech to launch its European manufacturing presence allowing Legend Biotech to tap into the area's vast talent pool and leverage the Belgian life sciences ecosystem. The Group is excited to expand the existing manufacturing network to support the production and delivery of cilta-cel for patients across the globe.

The Belgium Manufacturing Facility adds to Legend Biotech's existing manufacturing facilities and presence in Nanjing, China and New Jersey, United States. The Belgium Manufacturing Facility is anticipated to be operational by 2023.

CORPORATE PRESENTATION OF LEGEND BIOTECH

Reference is made to the announcement of the Company dated 9 June 2021.

On 22 June 2021, Legend Biotech posted an updated version of its corporate presentation (the "**Presentation**") to its website. For details, please refer to the attached Presentation. The attached Presentation is published on the Legend Biotech's website available at https://investors.legendbiotech.com/static-files/5c1f2394-0676-49c3-8530-38436caf485b.

This announcement has been issued in the English language with a separate Chinese language translation. If there is any inconsistency or ambiguity between the English version and the Chinese version, the English version shall prevail.

Shareholders and potential investors of the Company are advised to pay attention to investment risks and exercise caution when they deal or contemplate dealing in the securities of the Company.

By order of the Board

GenScript Biotech Corporation

MENG Jiange

Chairman and Executive Director

Hong Kong, 22 June 2021

As at the date of this announcement, the executive Directors are Mr. Meng Jiange, Ms. Wang Ye and Dr. Zhu Li; the non-executive Directors are Dr. Wang Luquan, Mr. Pan Yuexin and Ms. Wang Jiafen; and the independent non-executive Directors are Mr. Guo Hongxin, Mr. Dai Zumian, Mr. Pan Jiuan and Dr. Wang Xuehai.

* For identification purposes only

Inspired by the human element to advance cell therapy

June 2021





Disclaimer

This presentation has been prepared by Legend Biotech Corporation ("Legend Biotech" or the "Company") solely for information purpose and does not contain all relevant information relating to the Company.

The safety and efficacy of the agents and/or uses under investigation discussed in this presentation have not been established. There is no guarantee that the agents will receive health authority approval or become commercially available in any country for the uses being investigated.

Certain information contained in this presentation and statements made orally during this presentation relate to or are based on studies, publications, surveys and other data obtained from third-party sources and Legend Biotech's own internal estimates and research. While Legend Biotech 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. While Legend Biotech believes its internal research is reliable, such research has not been verified by any independent source.

Forward-Looking Statements

This presentation contains "forward-looking statements" within the meaning of The Private Securities Litigation Reform Act of 1995. The words "anticipate," "believe," "continue," "could," "estimate," "expect," "intend," "may," "plan," "potential," "predict," "project," "should," "target," "will," "would" and similar expressions are intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words.

These forward-looking statements include, but are not limited to, statements relating to the Company's strategies and objectives; the anticipated timing of, and ability to progress, clinical trials; the ability to make, and the timing of, regulatory submissions in the United States, Europe and Asia, including Biologics License Application (BLA) submission to the U.S. Food and Drug Administration (FDA) for ciltacabtagene autoleucel (cilta-cel) for relapsed or refractory multiple myeloma (RRMM), the submission of a marketing authorisation application (MAA) for cilta-cel to the European Medicines Agency (EMA), and the submission of an Investigational New Drug (IND) for LB1901 in relapsed or refractory T-Cell Lymphoma (TCL); the ability to generate, analyze and present data from clinical trials; patient enrollment; anticipated timing regarding regulatory approvals by the FDA, EMA or Center for Drug Evaluation (CDE); and the potential benefits of Legend Biotech's product candidates. Actual results may differ materially from those indicated by such forward-looking statements as a result of various important factors. Legend Biotech's expectations could be affected by, among other things, uncertainties involved in the development of new pharmaceutical products; unexpected clinical trial results, including as a result of additional analysis of existing clinical data or unexpected new clinical data; unexpected regulatory actions or delays, including requests for additional safety and/or efficacy data or analysis of data, or government regulation generally; unexpected delays as a result of actions undertaken, or failures to act, by our third party partners; uncertainties arising from challenges to Legend Biotech's patent or other proprietary intellectual property protection, including the uncertainties involved in the US litigation process; competition in general; government, industry, and general public pricing and other political pressures; the duration and severity of the COVID-19 pandemic and governmental and regulato

Should one or more of these risks or uncertainties materialize, or should underlying assumptions prove incorrect, actual results may vary materially from those described in this presentation as anticipated, believed, estimated or expected.

Any forward-looking statements contained in this presentation speak only as of the date of this presentation. None of the Company nor any of its affiliates, advisers, or representatives has any obligation and does not undertake to update any forward-looking statements to reflect future events or circumstances.

Legend Highlights



>900



Employees

10+



Pipeline Programs Covering:

- Hematologic malignancies
- Solid tumors
- Infectious disease

4



R&D Platforms:

- Autologous CAR-T
- Allogeneic CAR-T
- TCR
- NK

3

Global
Manufacturing Sites:

- United States
- EU
- China



\$462

Million

in Cash and Cash Equivalents as of Q1 2021 \$300

Million

PIPE Investment in May 2021



Cell Therapy Platform Overview

We Are A Fully Integrated Global Cellular Therapy Company



COMPELLING DATA WITH INNOVATIVE PIPELINE

- Lead product candidate ciltacabtagene autoleucel (cilta-cel) may have the potential to deliver deep and durable anti-tumor responses in **RRMM**
- Broad portfolio of earlier-stage autologous product candidates targeting both hematologic and solid cancers, as well as allogeneic CAR-T approaches

FUTURE PIPELINE

GASTRIC

OVARIAN

INFECTIOUS

GLOBAL COLLABORATION WITH JANSSEN*

- Global collaboration with Janssen for the development of cilta-cel established December 2017
 - Received an upfront payment of \$350 million and a total of \$200 million in milestone payments to date
 - Up to an additional \$1,150 million in potential future milestone payments











INTEGRATED CELL THERAPY PLATFORM

- In-house antibody generation and CAR-T specific functional screening technologies
- Early clinical proof-of-concept, leveraging KOL relationships in China, the US and globally
- Building large-scale manufacturing facilities in the United States, Europe and China
- >900 employees worldwide in US. China and Europe



Legend Biotech's Global R&D Strategy









Product Platforms

Autologous

Allogeneic

Disease Areas

Hematologic Malignancies

Solid Tumors

Infectious Diseases

With a Presence in Major Geographies, our Mission is to Improve the Lives of Patients Worldwide

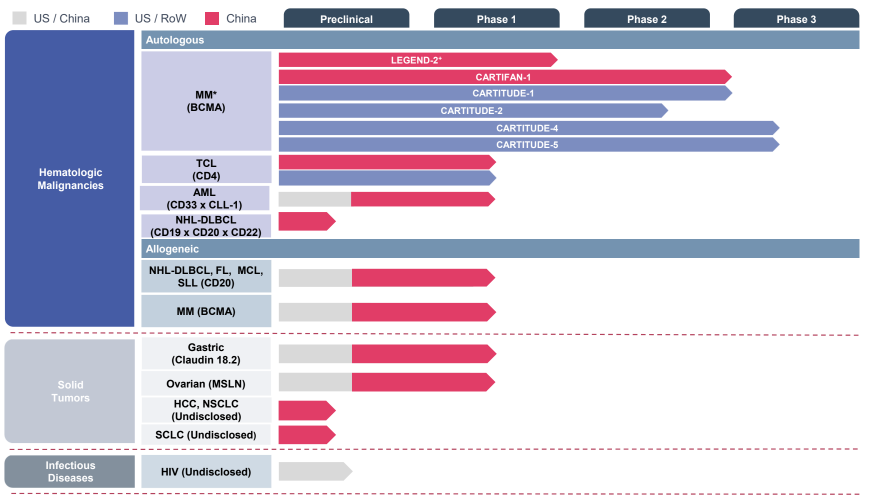


End-to-End R&D Capability

High-throughput antibody screening and Robust in vitro and in vivo engineering capability including screening platforms to single-domain antibodies generated from prioritize pipeline assets Llama and conventional antibodies **Antibody Screening Pre-clinical Platforms Validation Clinical Proof** of Concept **Binding Domain** Selection and Construct Design Proprietary methodology to optimize the selection Efficient clinical translation, of binding domains and design CAR-T constructs leveraging deep relationships with KOLs in US and China with two or more antigen-binding domains



Robust Pipeline of the Next Generation Cell Therapies



AML, acute myeloid leukemia; BCMA, B-cell maturation antigen; DLBCL, diffuse large B-cell lymphoma; FL, follicular lymphoma; HCC, hepatocellular carcinoma; HIV, human immunodeficiency virus; MCL, mantle cell lymphoma; NHL, non-Hodgkin lymphomas; MM, multiple myeloma; MSLN, mesothelin; NSCLC, non small cell lung cancer; RoW, Rest of World; SCLC, small cell lung cancer; SLL, small lymphocytic lymphoma; TCL, T-cell lymphoma

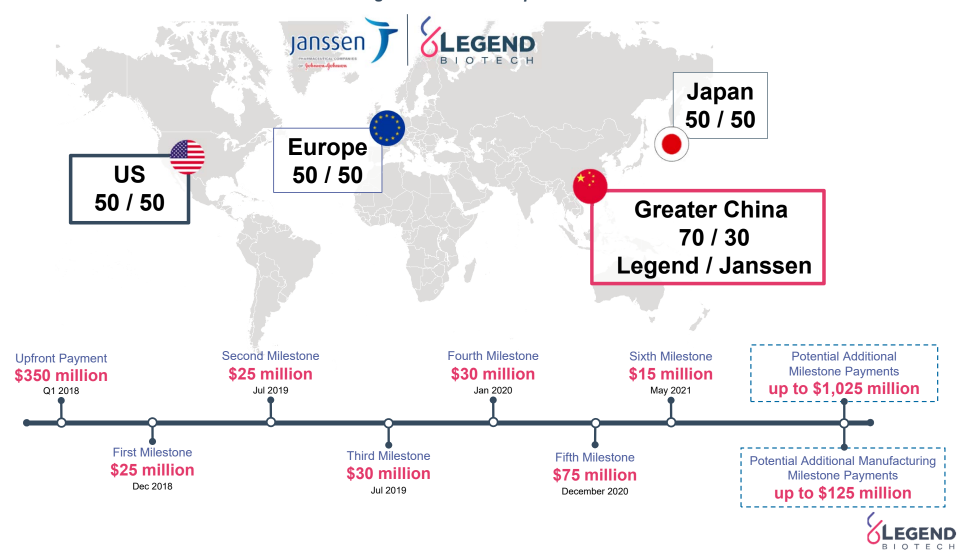


^{*}In collaboration with Janssen, Pharmaceutical Companies of Johnson & Johnson

^{*}LEGEND-2 trial is completed with ongoing follow-up

Legend and Janssen Global Collaboration

Worldwide collaboration and license agreement to develop and commercialize cilta-cel



Highly Experienced Management Team



YING HUANG Chief Executive Officer/ Chief Financial Officer

















Lida Pacaud Clinical Development **U** NOVARTIS Roche Wyeth



DONG GENG Early-stage Drug Development





STEVE GAVEL Commercial Development Celgene ≡IQVIA MILLENNIUM AMGEN



ALAN KICK Global Quality





ELIZABETH GOSEN Global Manufacturing







YUHONG QIU Global Regulatory **b** NOVARTIS Johnson Johnson



MEETA CHATTERJEE Global Business Development







Lori Macomber AMETEK





FRANK FAN Chief Scientific Officer & Co-Founder GenScript SickKids TORONTO



SIMON WU Research & Development GenScript



TRACY LUO Clinical Development AMGEN AstraZeneca 🕏



CHONG YANG Commercial Development Roche BAYER **U** NOVARTIS





Multiple Myeloma: Blood Cancer with a High Unmet Need



3RD MOST COMMON BLOOD CANCER

accounting for 18% of all hematologic cancer¹⁻³

176,404 NEW CASES WORLDWIDE IN 2020.

accounting for 1% of worldwide new cancer cases^{3,4}



32,119, with mortality of 13,426⁵



50,918, with mortality of 32,495⁶

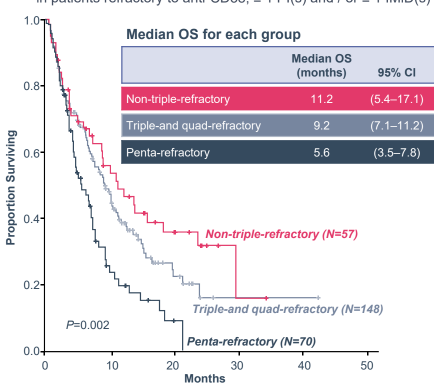


21,116, with mortality of 16,182⁷

POOR SURVIVAL OUTCOMES IN MULTIPLE REFRACTORY MM

Median OS < 12 months

in patients refractory to anti-CD38, ≥ 1 PI(s) and / or ≥ 1 IMiD(s)⁸



CI, confidence interval; PI, Proteasome Inhibitor; IMiD, immunomodulatory drug; MM, multiple myeloma; OS, overall survival

1. Cancer Stat Facts: Myeloma. httml. Accessed June 2021. 2. Facts and Statistics. https://ycv.iarc.fr/today/data/factsheets/concers/35-Multiple-myeloma-fact-sheets.pdf. Accessed June 2021. 4. Globocan 2020 World Fact Sheet: World. https://gco.iarc.fr/today/data/factsheets/populations/900-world-fact-sheets.pdf. Accessed June 2021. 5. Globocan 2020 World Fact Sheet: United States of America. https://gco.iarc.fr/today/data/factsheets/populations/908-europe-fact-sheets.pdf. Accessed June 2021. 7. Globocan 2020 World Fact Sheet: China. https://gco.iarc.fr/today/data/factsheets/populations/908-europe-fact-sheets.pdf. Accessed June 2021. 7. Globocan 2020 World Fact Sheet: China. https://gco.iarc.fr/today/data/factsheets/populations/908-europe-fact-sheets.pdf. Accessed June 2021. 8. Gandhi UH et al. Leukemia. 2019;33:2266-75.



First-in-Human, Phase 1, Dose Finding Study in RRMM LEGEND-2: LCAR-B38M CAR-T cells

Nanjing





Shanghai Shanghai Ruijin Hospital² Chen, ASH 2019 Poster Shanghai Changzheng Hospital² Chen, ASH 2019 Poster

Key Inclusion Criteria^{1,}

- Active MM defined by IMWG criteria with documented disease progression during or within 12 months of most recent anti-MM drugs or auto-HSCT
- Relapsed on prior regimens

Enrollment

- Total: 74 patients (4 sites in China)
- Xi'an: N=57, Wang, et al. ASH 2019
- JS/RJ/CZ sites: N=17, Chen, et al. ASH 2019

Preconditioning

- Cyclophosphamide only (Xi'an, Jiangsu)^{1,2}
- Cyclophosphamide + fludarabine (Changzheng, Ruijin)²

Administered dose (CAR+ viable T cells/kg)

- Xi'an¹ (median)=0.5x106 (0.07-2.1x106)
- RJ/CZ/JS² (mean)=0.70x10⁶ (0.2–1.5x10⁶)

Safety & Tolerability

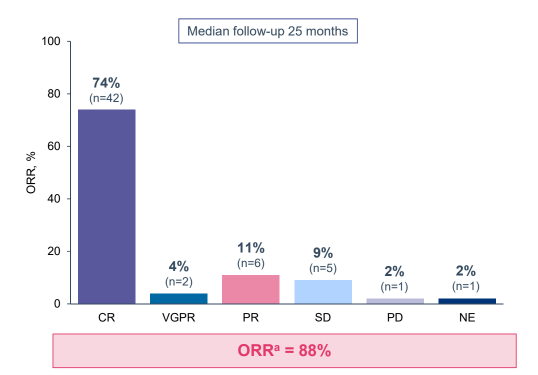
 Cilta-cel CAR-T cells displayed a safety profile consistent with other safety reports of BCMA-targeting CAR-T cell therapy^{1,2}



LEGEND-2: Long-Term Deep Responses and High Response Rate

Xi'an: Best overall response (N=57)¹

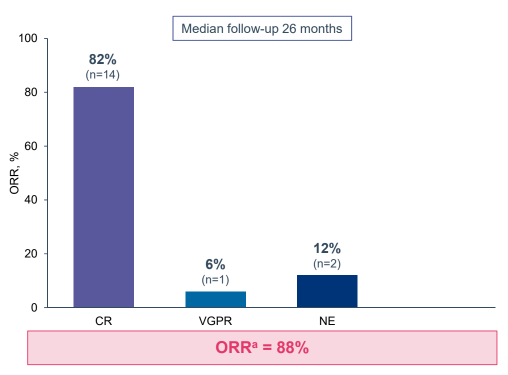
- mDOR= 27.0 months (mDOR for CR= 29.1 months)¹
- Median time to initial response= 1 month¹
- mPFS= 19.9 months (mPFS for CR= 28.2 months)¹
- mOS = 36.1 months (mOS for CR not reached)¹



Ruijin (RJ), Jiangsu (JS), Changzheng (CZ):

Best overall response (N=17)²

- Median time to initial response= 1 month²
- mPFS = 18 months; mOS= not reached²



Data cut-off: 31 July 2019 (N=57) and 31 October 2019 (N=17); Xi'an: NE patient died of PE/ACS prior to first evaluation. RJ,JS, CZ: For NE patients, 1 patient died on Day 13 due to CRS and tumor lysis syndrome; 1 patient received chemotherapy prior to first assessment and was censored. a ORR=PR or better; response assessed per International Myeloma Working Group criteria

CR, complete response; VGPR, very good partial response; PR, partial response; SD, stable disease; PD, progressive disease; NE, not evaluable; mDOR, median duration of response; MRD, minimal residual disease; ORR, overall response rate; mPFS, median progression free survival; mOS, median overall survival.





CARTITUDE-1: Phase 1b/2 Study Design

- Phase 1b: Characterize the safety of ciltacabtagene autoleucel (cilta-cel) and confirm the recommended phase 2 dose
- Phase 2: Evaluate the efficacy of cilta-cel by ORR
- Progressive MM per IMWG criteria
- ECOG PS ≤1
- Measurable disease
- Received ≥3 prior therapies or double refractory
- Prior PI, IMiD, anti-CD38 therapy

Median administered dose:
 0.71x10⁶ (0.51- 0.95x10⁶) CAR+ viable T cells/kg







Bridging Therapy^a (as needed)



Day -5 to -3



Cilta-cel Infusion
Target: 0.75x10⁶ (0.5 – 1.0x10⁶)
CAR+ viable T cells/kg

Day 1



Post-infusion Assessments Safety, Efficacy, PK, PD, Biomarker



Follow-up

Cy, cyclophosphamide; ECOG PS, Eastern Cooperative Oncology Group performance status; Flu, fludarabine; IMiD, immunomodulatory drug; IMWG, International Myeloma Working Group; PI, proteasome inhibitor; PD, pharmacodynamic; PK, pharmacokinetic; MM, multiple myeloma Data cut-off: Feb 11, 2021; a Treatment that was received previously and resulted in at least stable disease.

ULEGEN

CARTITUDE-1: Baseline Characteristics

Characteristic (N=97)		Characteristic	
Age, median (range) years	61.0 (43–78)	Prior lines of therapy, median (range)	
Male, n (%)	57 (58.8)	Prior lines of therapy, n (%)	
Black/African American, n (%)	17 (17.5)	3	
All plasmacytomas, ^a n (%)	19 (19.6)	4 ≥5	
Extramedullary plasmacytomas, n (%)	13 (13.4)	Previous stem-cell transplantation, n (%)	
Bone-based plasmacytomas, n (%)	6 (6.2)	Autologous	
Bone-marrow plasma cells ≥60%, n (%)	21 (21.9)	Allogeneic	
	, ,	Triple-class exposed, ^c n (%)	
Years since diagnosis, median (range)	5.9 (1.6–18.2)	Penta-drug exposed,d n (%)	
High-risk cytogenetic profile, n (%)	23 (23.7)	Triple-class refractory ^c	
del17p	19 (19.6)	Penta-drug refractory ^d	
t(14;16)		Refractory status, n (%)	
t(1 4 ,10)	2 (2.1)	Carfilzomib	
t(4;14)	3 (3.1)	Pomalidomide	
Tumor BCMA expression ≥50%, n (%)	57 (91.9) ^b	Anti-CD38 antibody	
		Refractory to last line of therapy, n (%)	

Data cut-off: Feb 11, 2021; BCMA, B-cell maturation antigen; IMiD, immunomodulatory drug; Pl, proteasome inhibitor.

^aAll plasmacytomas include extramedullary and bone-based plasmacytomas. ^bDenominator n=62, the number of evaluable samples; BCMA expression detected in all evaluable samples. ^cAt least 1 Pl, at least 1 IMiD, and 1 anti-CD38 antibody. ^dAt least 2 Pls, at least 2 IMiDs, and 1 anti-CD38 antibody.

Usmani S, et al. ASCO Annual Meeting (Virtual). June 4-8, 2021. Abstract 8005



6.0 (3-18)

17 (17.5) 16 (16.5) 64 (66.0)

87 (89.7) 8 (8.2) 97 (100)

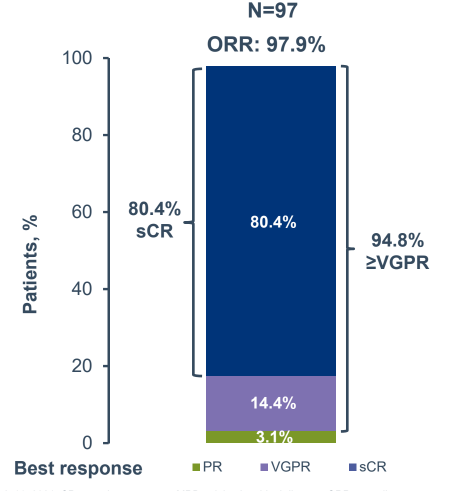
81 (83.5) 85 (87.6) 41 (42.3)

63 (64.9)

81 (83.5)

96 (99.0) 96 (99.0)

CARTITUDE-1: Overall Response Rate



With longer follow-up, responses deepened with increasing rate of sCR

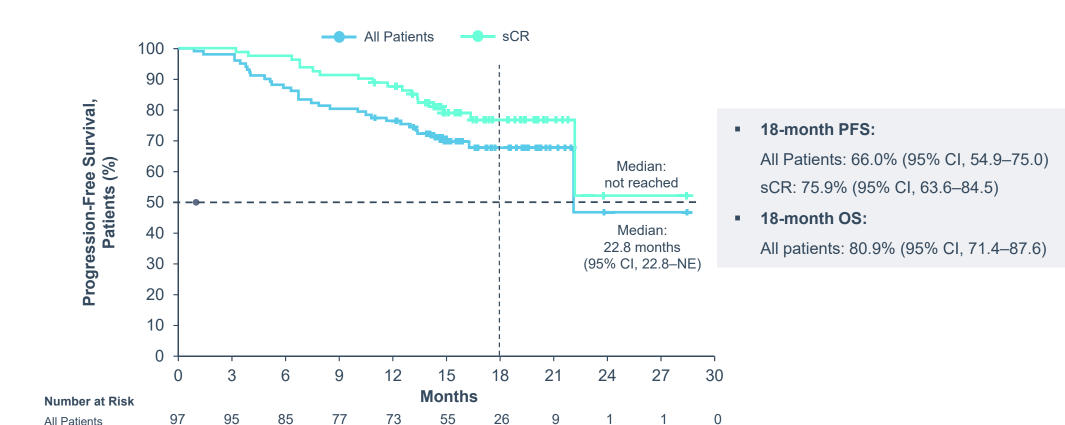
- Median time to first response: 1 month (range, 0.9–10.7)
- Median duration of response: 21.8 months (95% CI, 21.8– NE); not reached in patients with sCR
- Response rates were comparable (range, 95–100%) across different subgroups (eg, number of prior lines of therapy, refractoriness, extramedullary plasmacytomas, and cytogenetic risk)^a
- 91.8% of 61 evaluable patients were MRD negative^b
 - Median time to MRD 10⁻⁵ negativity: 1 month (range, 0.8–7.7)

Data cut-off: Feb 11, 2021; CR, complete response; MRD, minimal residual disease; ORR, overall response rate; sCR, stringent complete response; VGPR, very good partial response. ORR assessed by independent review committee. aSubgroups by number of prior lines of therapy (≤4, >4), refractoriness (triple-class, penta-drug), cytogenetic risk (high risk, standard risk), baseline bone marrow plasma cells (≤30%, >30 to <60%, ≥60%), baseline tumor BCMA expression (≥median, <median), and baseline plasmacytomas (including extramedullary and bone-based). bMRD was assessed in evaluable samples (ie, patients with identifiable clone at baseline and sufficient cells for testing at 10-5 threshold in post treatment samples) by next-generation sequencing (clonoSEQ, Adaptive Biotechnologies) in all treated patients at Day 28, and at 6, 12, 18, and 24 months regardless of the status of disease measured in blood or urine.

Usmani S. et al. ASCO Annual Meeting (Virtual), June 4-8, 2021, Abstract 8005



CARTITUDE-1: Progression Free Survival



Median duration of follow-up: 18 months (range, 1.5-30.5)

0

26

51



76

71

78

Responders With sCR 78

CARTITUDE-1: Safety

	N =	N = 97		
	Any Grade	Grade 3/4		
Hematologic AEs, (≥30%), n (%	%)			
Neutropenia	93 (95.9)	92 (94.8)		
Anemia	79 (81.4)	66 (68.0)		
Thrombocytopenia	77 (79.4)	58 (59.8)		
Leukopenia	60 (61.9)	59 (60.8)		
Lymphopenia	51 (52.6)	48 (49.5)		
Non-hematologic AEs (≥30%),	n (%)			
Hypocalcemia	31 (32.0)	3 (3.1)		
Hypophosphatemia	30 (30.9)	7 (7.2)		
Fatigue	36 (37.1)	5 (5.2)		
Cough	34 (35.1)	0		
CAR-T associated AEs, n (%)				
CRS ^a	92 (94.8)	4 (4.1)		
Neurotoxicity	20 (20.6)	9 (9.3)		

No new safety signals with longer follow-up

- CRS
 - 94.6% of patients experienced low-grade CRS (n=92)
 - Median time to onset of 7 days (range, 1-12)
 - Median duration of 4 days (range, 1-97)^b and resolved in 91 (98.9%) patients within 14 days of onset
- Neurotoxicity
 - 20.6% of patients experienced neurotoxicity in total with overlap between ICANS and Other Neurotoxicities (Grade ≥3: 10.3%)
 - ICANS observed in 16.5% (Grade ≥3: 2.1%)
 - Other Neurotoxicities^c observed in 12.4% (Grade ≥3: 9.3%)
- 6 treatment-related deaths as assessed by the investigator^d

Data cut-off: Feb 11, 2021; AE, adverse event; ALT, alanine aminotransferase; AST, aspartate aminotransferase; ASTCT, American Society for Transplantation and Cellular Therapy; CRS, cytokine release syndrome; HLH, hemophagocytic lymphohisticocytosis. ^aCRS was graded using Lee et al. (*Blood* 2014) in the phase 1b portion of the study and ASTCT in phase 2; in this combined analysis, Lee et al. criteria were mapped to ASTCT criteria for patients in the phase 1b portion. ^bThe patient with 97-day duration died due to CRS/HLH. ^cEvents not reported as ICANS (ie, onset after a period of recovery from CRS and/or ICANS). ^dThere were 21 study deaths: 6 were treatment-related as assessed by the investigator, the remaining were due to AEs unrelated to treatment (n=5) and disease progression (n=10)
Usmani S, et al. ASCO Annual Meeting (Virtual). June 4-8, 2021. Abstract 8005

CARTITUDE-2: Multicohort Study Cohort A: 1 – 3 prior lines, lenalidomide refractory RRMM

 CARTITUDE-2 is a phase 2, multicohort, open-label study assessing the efficacy and safety of cilta-cel in patients with multiple myeloma in various clinical settings

BCMA-binding domains VHH VHH 4-1BB CD3ζ Cilta-cel (CAR-T)

Cohort A:

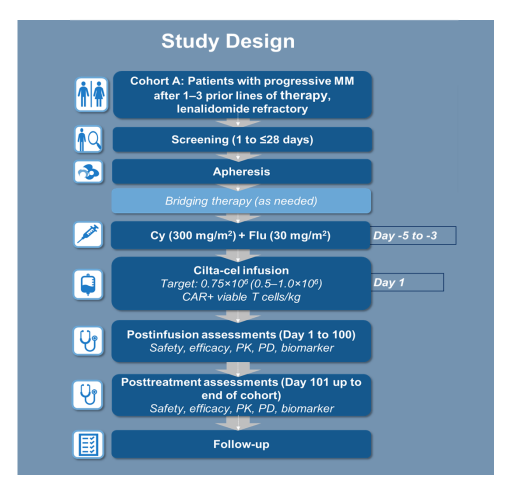
- Cohort A patients had progressive MM after 1–3 prior lines of therapy, and were refractory to lenalidomide
- Despite advances continued unmet need with mPFS of 9.9 months (DPd)¹

Primary objectives

 Minimal residual disease (MRD) 10⁻⁵ negativity

Secondary objectives

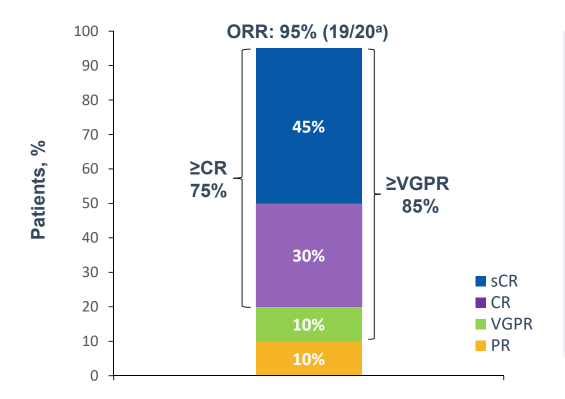
 ORR, duration of response, time and duration of MRD negativity, and incidence and severity of adverse events





CARTITUDE-2: Phase 2 Multi-Cohort Study

- Cohort A included 20 patients who had progressive MM after 1–3 prior lines of therapy and were refractory to lenalidomide
- Median prior lines of therapy: 2 (range, 1-3); 1 patient treated in an outpatient setting



- No progression of disease at median follow-up of 5.8 months (range 2.5-9.8)
- All patients (n=4) with MRD-evaluable^b samples at the 10⁻⁵ threshold were MRD negative at data cut-off
- The safety profile was manageable
 - CRS occurred in 85% (n=17); mostly grades 1/2;
 median time to CRS onset was 7 days (range, 5–9)
 - Neurotoxicities occurred in 20% (n=4) of patients; no grade ≥3; no incidence of movement and neurocognitive TEAEs
 - 1 death occurred 100 days after infusion due to COVID-19 (assessed as tx related by the investigator)

Data cut-off date: Jan 2021; ^aPatient who did not respond had stable disease. ^bMRD was assessed in evaluable samples (ie, patients with identifiable clone at baseline and sufficient cells for testing at 10⁻⁵ threshold in post treatment samples) by next-generation sequencing (clonoSEQ, Adaptive Biotechnologies) in all treated patients.

CR, complete response; MRD, minimal residual disease; ORR, overall response rate; PR, partial response; sCR, stringent complete response; TEAE, treatment-emergent adverse events; VGPR, very good partial response.

Agha M, et al. ASCO Annual Meeting (Virtual). June 4-8, 2021. Abstract 8013.



CARTITUDE Program: Safety

Successful new patient management strategies have been implemented in the CARTITUDE program to prevent and reduce the incidence of neurotoxicity¹⁻³

Movement and Neurocognitive TEAEs^a

 Occurred in 5 of 97 patients in CARTITUDE-1

Risk factors (2 or more)

- High tumour burden^b
- Grade ≥2 CRS
- ICANS
- High CAR T-cell expansion and persistence

Patient Management Strategies

- Enhanced bridging therapy to reduce tumour burden
- Early and aggressive treatment of CRS and ICANS
- Handwriting assessments and extended monitoring

>100 additional patients

have been dosed^c

- Patient management strategies to prevent or reduce these AEs have been successfully implemented in new and ongoing cilta-cel studies
- This is reliant on effective implementation of these patient management strategies

^{1.} Usmani S, et al. ASCO Annual Meeting (Virtual). June 4-8, 2021. Abstract 8005. 2. Agha M, et al. ASCO Annual Meeting (Virtual). June 4-8, 2021. Abstract 8013. 3. Einsele H, et al. ASCO Annual Meeting (Virtual). June 4-8, 2021. Abstract 8028



AE, adverse event; CRS, cytokine release syndrome; ICANS, immune effector cell-associated neurotoxicity syndrome; TEAE, treatment-emergent AE.

^aTwo patients with ongoing symptoms continued to improve at the time of data cutoff; patient management strategies were implemented, including enhanced bridging therapy to reduce baseline tumor burden, early aggressive treatment of CRS and ICANS, handwriting assessments for early detection of neurotoxicity symptoms, and extended monitoring and reporting time for neurotoxicity beyond the first 100 days post-cilta-cel infusion. ^bDefined as having high tumor burden when any of the following parameters were met: bone marrow plasma cell ≥80%, serum M-spike ≥5 g/dL, serum free light chain ≥5000 mg/L. ^cIncluded patients treated in earlier and later line settings across the CARTITUDE program.

Clinical Program: Cilta-cel Studies in Multiple Myeloma

FIH Study in China Long-term Follow-up

LEGEND-21



Registrational **Studies**



















NCT03090659

- Phase 1. multi-center study of LCAR-B38M CAR-T cells in RRMM
- Fully enrolled and ongoing in China

CARTITUDE-1 MMY2001²

CARTIFAN-1

MMY20023

NCT03548207

- Phase 1b/2, multi-center registration study of cilta-cel in RRMM
- Fully enrolled and ongoing in **US** and Japan

CARTITUDE-2 MMY20034

NCT04133636

- Global, multi-cohort study
- Phase 2 open-label study of cilta-cel in various clinical settings to evaluate MRD negativity

NCT04181827

■ Enrolling in US / EU/ Israel

NCT03758417

- Phase 2, multi-center confirmatory study of cilta-cel in RRMM
- Ongoing in China

CARTITUDE-4

MMY3002⁵

- Global, randomized study
- Phase 3 open-label study of cilta-cel vs DPd or PVd in patients with RRMM, 1-3 lines of prior therapy and refractory to lenalidomide
- Enrolling in US/EU/JP/AUS/ Israel/Korea

CARTITUDE-5 MMY3004⁶

NCT04923893

- Global, randomized study
- Phase 3 open-label study of VRd followed by cilta-cel vs. VRd followed by Rd maintenance. in patients with newly diagnosed MM for whom ASCT is not planned as initial therapy
- Planned in US/Canada/EU/AUS/Israel/Brazil

ASCT, autologous stem cell transplant; DPd, daratumumab, pomalidomide, dexamethasone; EU, European Union; JP, Japan; PVd, pomalidomide, bortezomib, dexamethasone; RRMM, relapsed and/or refractory multiple myeloma; SoC, standard of care; US, United States; VRd, bortezomib, lenalidomide, dexamethasone

1 NCT03090659. Clinicaltrials gov website. https://clinicaltrials.gov/ct2/show/NCT03090659. Accessed June 2021; 2 NCT03548207. Clinicaltrials.gov website. https://clinicaltrials.gov/ct2/show/NCT03548207. Accessed June 2021. CARTITIDE-1 is global registration study; 3 NCT03758417. Clinicaltrials.gov website. https://clinicaltrials.gov/ct2/show/NCT03758417. Accessed June 2021. CARTIFAN-1 is registration study for China only; 4 NCT04133636. Clinicaltrials.gov website. https://clinicaltrials.gov/ct2/show/NCT04133636. Accessed June 2021; 5 NCT04181827. Clinicaltrials.gov website: https://clinicaltrials.gov/ct2/show/NCT04181827. Accessed June 2021; 5 NCT04181827. Clinicaltrials.gov/ct2/show/NCT04181827. Accessed June 2021; 5 NCT04181827. Accessed June 2021; 5 NCT0418182. Accessed June 2021; 5 NCT0418182. Ac 2021; 6 NCT04923893. Clinicaltrials.gov website: https://clinicaltrials.gov/ct2/show/NCT04923893. Accessed June 2021.



Global Manufacturing Footprint

US Facilities



EU Facilities



China Facilities

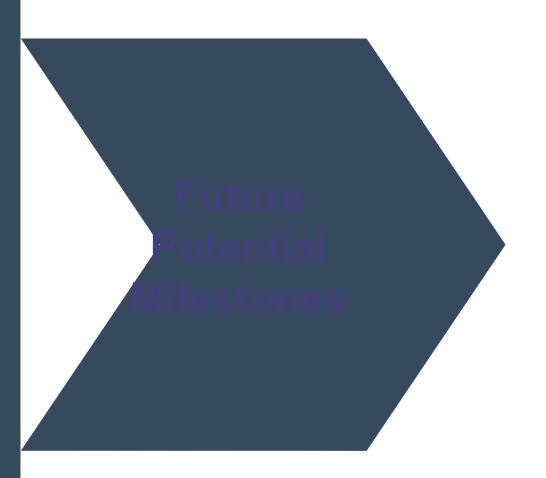


Construction in progress

Building E



Future Potential Milestone Payments



Clinical Milestones: \$105M

\$105 million for the achievement of specified future development milestones

Regulatory Milestones: \$710M

\$710 million for the achievement of specified regulatory milestones

Commercial Milestones: \$210M

\$210 million for the achievement of specified net trade sales milestones.

Manufacturing Milestones: \$125M

Further milestone payments of up to \$125 million for the achievement of specified manufacturing milestones



Program Areas of Development

Legend Biotech is utilizing the extensive cell therapy experience of our leadership and R&D staff, global clinical partners, and expanding research facilities to realize the potential of cell therapy to treat diseases that are thought to be incurable, such as hematologic malignancies, solid tumors and infectious diseases.





LB1901: Investigational CAR-T for T Cell Lymphoma

MoA/ Scientific Rationale

- LB1901 targets CD4 antigen that is expressed in most T cell lymphoma (TCL) subtypes and in subsets of normal immune cells
- LB1901 is a CD8-enriched anti-CD4 CAR-T and contains a unique binder to CD4 leading to potential elimination of CD4+tumor cells

Target

- CD4 is a surface membrane glycoprotein expressed at high levels on TCL and a subtype of normal T cells¹
- Anti-CD4 mAb have been investigated in clinical studies for TCL²

Clinical Development

- US IND cleared with FDA
- Ongoing Phase 1 studies in US and China
- Patient population: relapsed/refractory PTCL and CTCL patients



LB1908: Investigational CAR-T for Gastric Cancer

MoA/ Scientific Rationale

- LB1908 targets Claudin (CLDN) 18.2 through high-affinity VHH antibody
- VHH antibody, identified via in-house, selectively binds to CLDN 18.2

Target

- Claudins are a family of tight junction proteins¹
- CLDN18.2 is commonly expressed on multiple cancers including gastric cancer²

Clinical Development

- Phase I clinical study in China is ongoing for the treatment of adult patients with advanced gastric cancer
- US IND is being developed with planned submission in 2H2021

LB1905: Investigational Allogenic CAR-T

MoA/ Scientific Rationale

- LB1905 targets CD20 that is expressed in B cell lymphoma
- LB1905 applied Legend UniCAR technology which is an unique non-geneediting allogeneic CAR-T platform
- Simple and efficient manufacturing promote product homogeneity and accessibility

Target

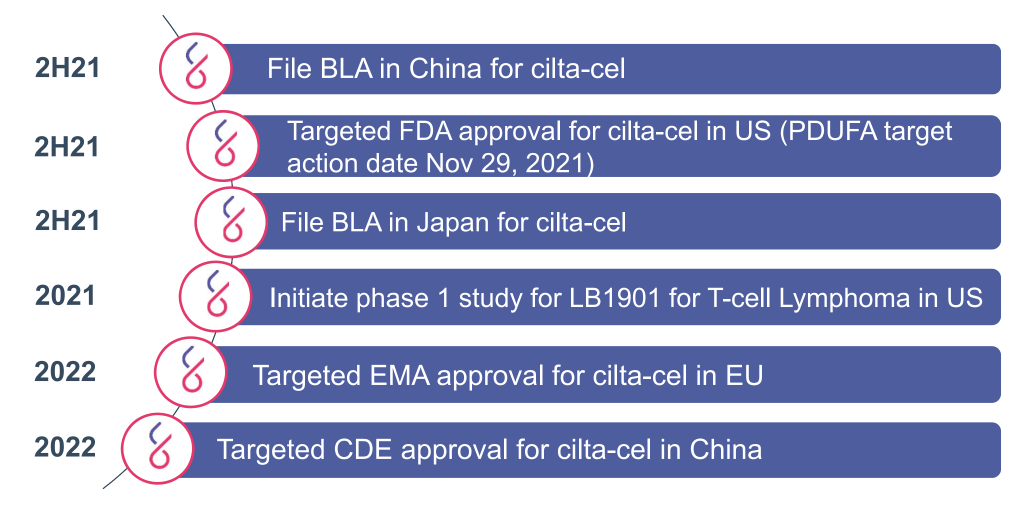
• CD20 is mainly expressed in pre-B cells and mature B cells. It is expressed in more than 95% of B-cell lymphomas and not in hematopoietic stem cells, plasma cells, and other normal tissues

Clinical Development

- Allogeneic CD20 targeted product for the treatment of adult patients with recurred NHL
- Promising allogeneic platform that can potentially be leveraged in Legend clinical development programs

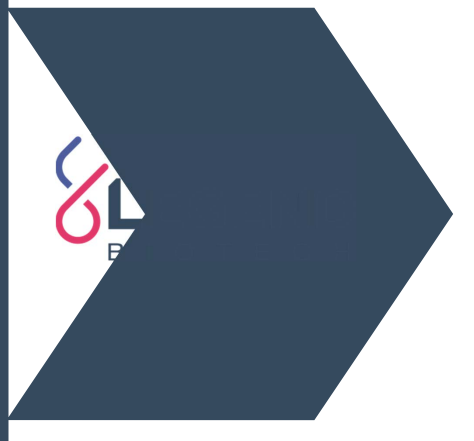


Near-Term Targets for Legend Biotech





Investment Highlights





Global Collaboration

Global collaboration with Janssen for the development of cilta-cel with ongoing clinical trials



Promising Clinical Data

Deep and durable anti-tumor responses observed in heavily pretreated patients with MM; BLA for cilta-cel submitted to US FDA (PDUFA target action date Nov 29, 2021); MAA for cilta-cel submitted to EMA



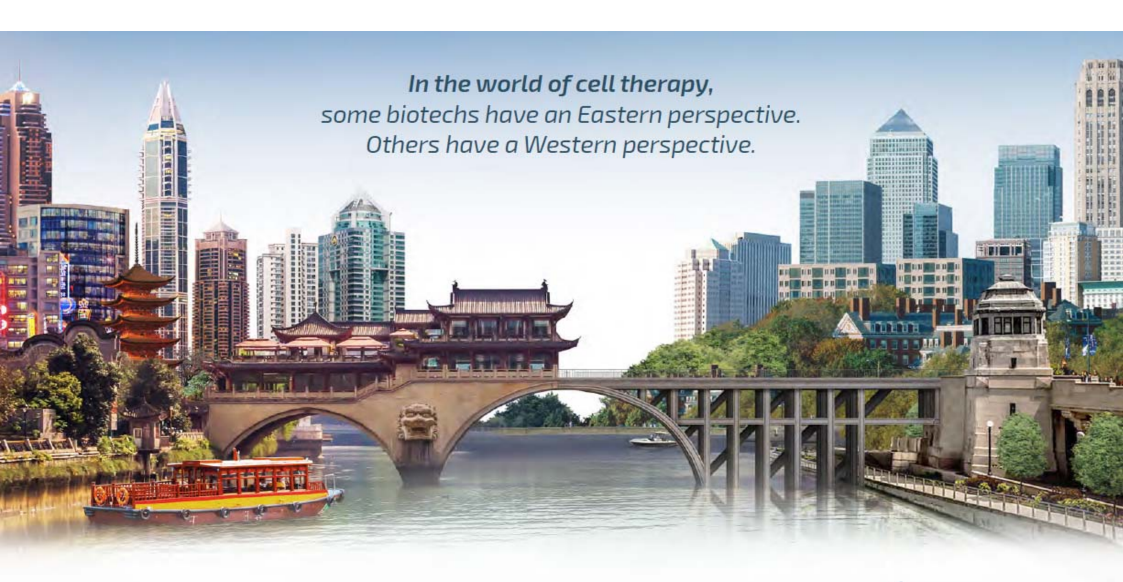
Fully Integrated Platform

End-to-end R&D and manufacturing capabilities with two core technologies (CAR and TCR) and two platforms (Autologous and Allogeneic)



Strong Management

Experienced team with broad involvement in biopharmaceutical drug discovery, development and commercialization



We are bridging the gap between East and West.



Thank You!



