

FORWARD-LOOKING STATEMENTS

This presentation contains "forward-looking" statements that are based on our management's current expectations and assumptions and on information currently available to management.

Forward-looking statements involve known and unknown risks, uncertainties and other factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements.

The risks and uncertainties include, but are not limited to the risk that the preliminary results from our product candidates will not continue or be repeated, the risk that our clinical trials will not be successful. The risk of not obtaining regulatory approval to commence clinical trials on additional UCART product candidates,

the risk that any one or more of our product candidates will not be successfully developed and commercialized.

Further information on the risk factors that may affect company business and financial performance, is included in our annual report on form 20-F and other filings Cellectis makes with the securities and exchange commission from time to time and its financial reports.

Except as required by law, we assume no obligation to update these forward-looking statements publicly, or to update the reasons actual results could differ materially from those anticipated in the forward-looking statements, even if new information becomes available in the future.

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OUR MISSION Leverage our leadership in gene editing and CART therapy to bring new hope to cancer patients through broadly available, off-the-shelf therapies

CELLECTIS - COMMITMENT TO A CURE



INNOVATION

Protein engineering for best-in-class gene editing & CAR technologies, cell engineering and culture technologies



LEADERSHIP

First clinical proof-ofconcept - UCARTs - for off-the-shelf CAR T therapy that address multiple unmet cancer needs



PIPELINE

Pioneering robust first-in-class allogeneic CAR T-cell programs for different hematological malignancies



MANUFACTURING

Comprehensive, scalable, efficient, and cost-effective manufacturing process generates highly potent CAR T therapies

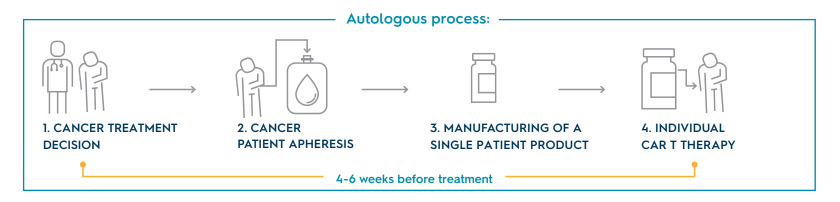


THERAPY

Making cancer therapies cost-effective and available faster to cancer patients

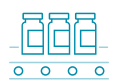


ADVANTAGES OF ALLOGENEIC VS. AUTOLOGOUS CAR T



Allogeneic process:

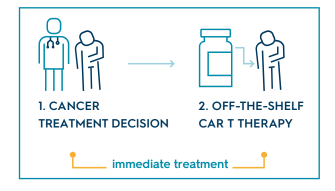




SCALABLE MANUFACTURING OF 100+ DOSES/BATCH

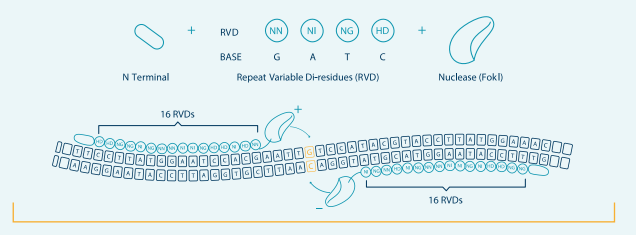


OFF-THE-SHELF
CAR T THERAPIES





TALEN®: BEST-IN-CLASS GENE EDITING



PRECISION

targeting within 6 base pairs of any target in the genome (effective changes)

SPECIFICITY

recognition site is 32 base pairs long (avoids errors)

EFFICIENCY

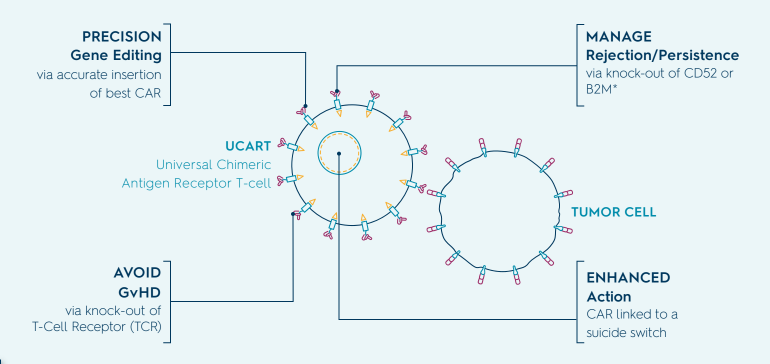
TCR- α can be knocked-out with over 95% accuracy for engineered CAR T-cells (ensures yield)

Editing genes allows disabling a functional gene, correcting a gene, or replacing or inserting a DNA sequence at a chosen location in a genome.

TALEN® has been successfully used in the clinic to solve key challenges with allogeneic CAR T including protection from GvHD, mitigation of rejection, chimerism and enhanced safety via a suicide switch.



UCARTS - FIRST & BEST-IN-CLASS ALLOGENEIC CART-CELLS





PARTNERSHIPS WITH INDUSTRY LEADERS

Development & commercialization partners -



UCART19 (with Allogene)

+ other targets

Up to \$1.1B in development milestones

Royalties on sales



15 LICENSED TARGETS

Up to \$2.8B in development & sales milestones

Royalties on sales

Equity investor —



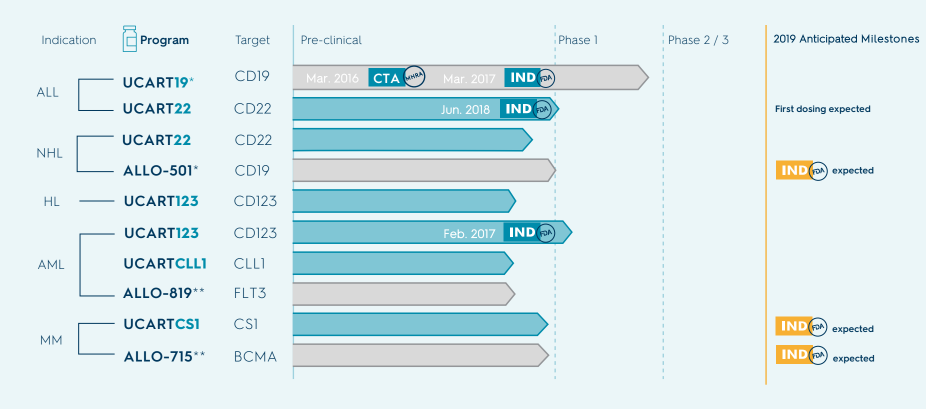
6.6% of outstanding shares

As of September 30, 2018

Up to \$3.9B in potential milestone payments plus royalties



PIPELINE: INNOVATIVE THERAPIES FOR UNMET CANCER NEEDS



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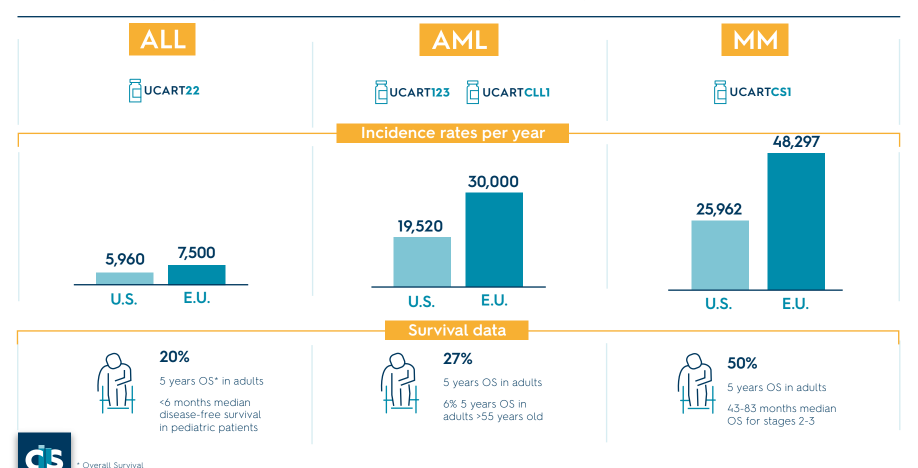
Proprietary development program

Licensed development program

^{*} UCART19 is exclusively licensed to Servier and under a joint clinical development program between Servier and Allogene. ALLO-501 is exclusively licensed to Servier

^{**} Product candidates exclusively licensed to Allogene

PIPELINE TARGETS MULTIPLE UNMET NEEDS IN CANCER



UCART19*: DESIGN OF PHASE 1 STUDIES IN R/R** ALL***

CD19 is a validated target expressed in B-cell malignancies

Adult ALL (CALM study) -

PRIMARY OBJECTIVE

Evaluate safety, tolerability, maximum tolerated dose (MTD) and regimen

SECONDARY OBJECTIVES

Objective remission rate at Day 28. Duration of response, time to remission, progression-free survival



ONGOING



Pediatric ALL (PALL study)

PRIMARY OBJECTIVE

Evaluate safety at a fixed dose in patients aged between 6 months and 18 years old

SECONDARY OBJECTIVES

Determine the ability to achieve molecular remission at Day 28



ONGOING



DL: 1.1 to 2.3x106 cells/kg



- * UCART19 is exclusively licensed to Servier and under a joint clinical development program between Servier and Allogene
- ** Relapsed/Refractory
- *** Acute Lymphoblastic Leukemia
- **** Dose Level

UCART19*: PHASE 1 R/R ALL - DATA** PRESENTED AT ASH 2018

Safety:



- ✓ **14%** Grade 3-4 Cytokine Release Syndrome
- ✓ 0% Grade 3-4 neurotoxicity
- **0%** Grade 3-4 skin Graft vs Host Disease

Efficacy: -

- → 82% CR/CRi rate in FCA***-treated patients
- → 67% overall CR/CRi rate
- → 71% of these patients were MRD-
- → Redosing with UCART19 resulted in cell expansion and MRD- status in 2/3 patients
- → Peak expansion observed mostly at Day 14

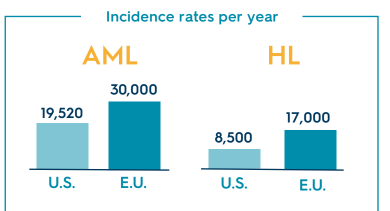


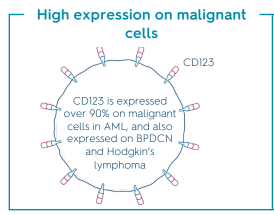
UCART19 is exclusively licensed to Servier and under a joint clinical development program between Servier and Allogene

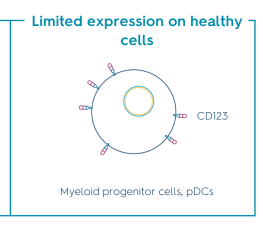
^{**}Pooled data

^{***} Lymphodepletion regimen consisting of fludarabine, cyclophosphamide and an anti-CD52 mAb

CD123 TARGET: RATIONALE FOR THERAPY











UCART123 - PHASE 1 STUDY IN AML

Patient characteristics

Age and fitness: R/R in AML 65 years and older, unfit patients

Mutation status: genetically complex Progression: rapid

progression following relapse

Dose escalation (mTPI*) phase (R/R AML)





28 days between the first 2 patients for each dose**, then 14 days for subsequent patients







DL3: 5.05x106/kg

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Expansion Phase



TOTAL N=64-144

Expected in 2020



R/R AML PATIENTS

N=18-37



FIRST LINE AML PATIENTS

ELN*** Adverse genetic group

V=46-107



- Modified Toxicity Probability Interval Design
- ** 42 days if aplasia
- *** European Leukemia Net

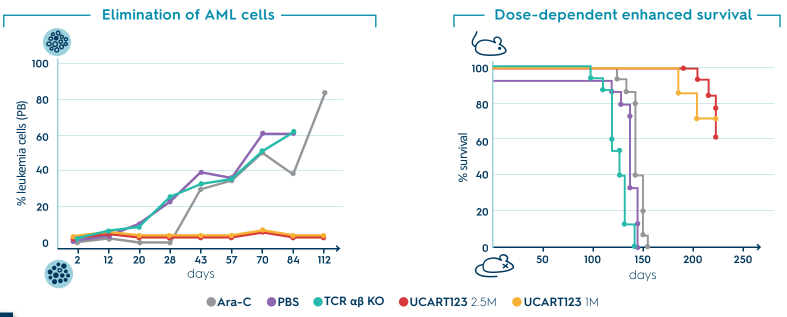
UCART123 - PRECLINICAL RATIONALE IN AML

Development rationale:-

High expression: blasts, independent of mutation status

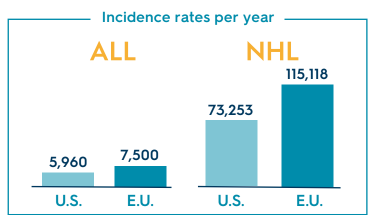
Unmet need: high relapse rate and poor survival in R/R patients

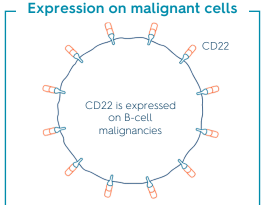
Validated target: CD123 - clinically validated in autologous CAR T-cell trials

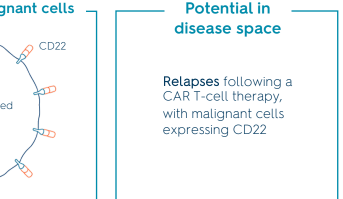




CD22 TARGET: RATIONALE FOR THERAPY















UCART22 - PHASE I TRIAL DESIGN IN ALL

Patient characteristics

Age and fitness:

R/R B-ALL < 65 years

CD19- & CD19+ ALL

high CD22 expressing B-malignant cells

Offers a therapeutic solution to patients who cannot receive, or relapsed, after autologous CD19 CAR T-cell therapy

Dose escalation (mTPI) phase

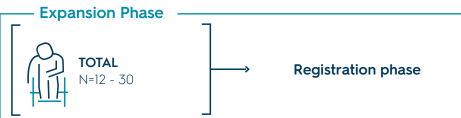


Starting at MD Anderson



28 days between the first 2 patients for each dose, then 14 days for subsequent patients







UCART22 - PRECLINICAL RATIONALE FOR ALL

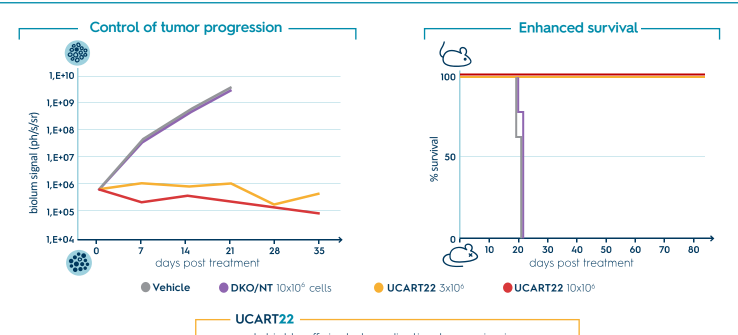
Development rationale:

CD22 expression: in CD19 CD19 negative blasts

Unmet need: high relapse rates (CD19-) after CAR T treatment, poor survival in R/R patients

Validated target in ALL and NHL

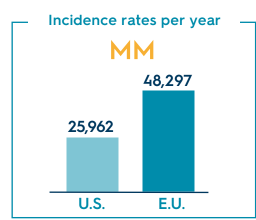
Expandable market: potential expansion into first-line ALL

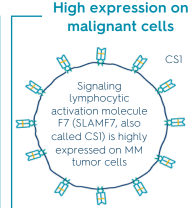




- Is highly efficient at eradicating tumors in vivo
- Result in increased survival in mouse model

CS1-SLAMF7 TARGET: RATIONALE FOR THERAPY





Limited expression on healthy cells CS1 T cells, B cells, macrophages and NK cells

Monoclonal antibody validation

- → **Elotuzumab** is a monoclonal antibody targeting CS1
- → Elotuzumab is safe and effective in MM patients
- → Elotuzumab in combination with lenalidomide and dexamethasone in R/R MM patients shows: 5.5% CR rate and 35% partial remissions

Competitive approaches















MB-104 [CS1]

bb2121/bb21217 [BCMA]

JNJ-68284528 [BCMA]

JCARH125 [BCMA]

AMG 420/701 [BCMA]

GSK2857916 [BCMA] Antibody drug

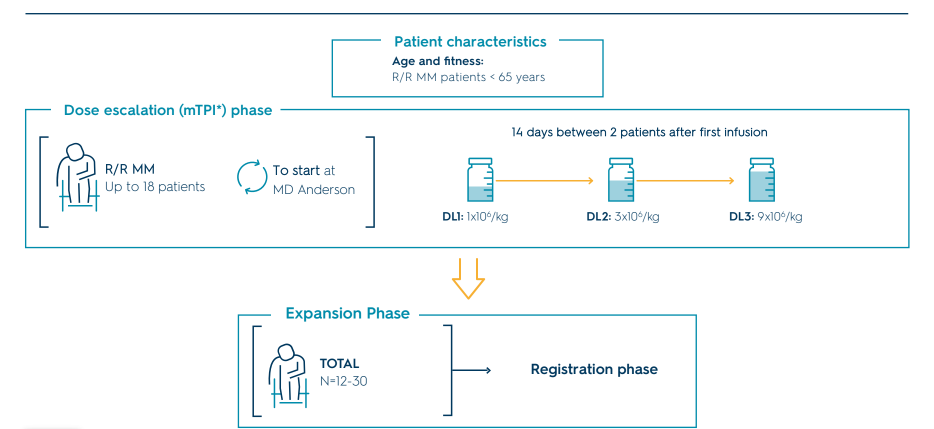
Autologous CAR T

Bispecific antibodies



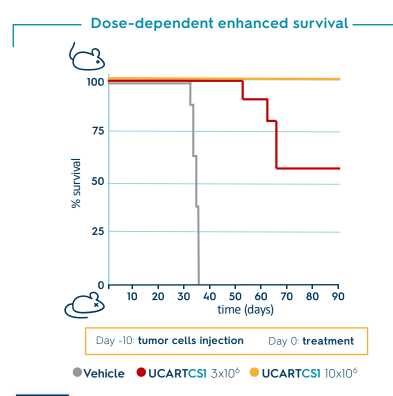


UCARTCS1 - PHASE 1 TRIAL DESIGN IN MULTIPLE MYELOMA

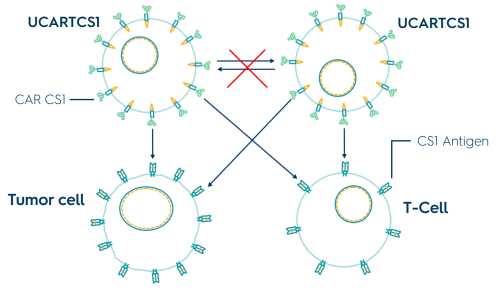




UCARTCSI - PRECLINICAL RATIONALE IN MULTIPLE MYELOMA



Knock-Out of CSI on CAR T-cells to suppress cross T-cell reaction between UCARTCSI

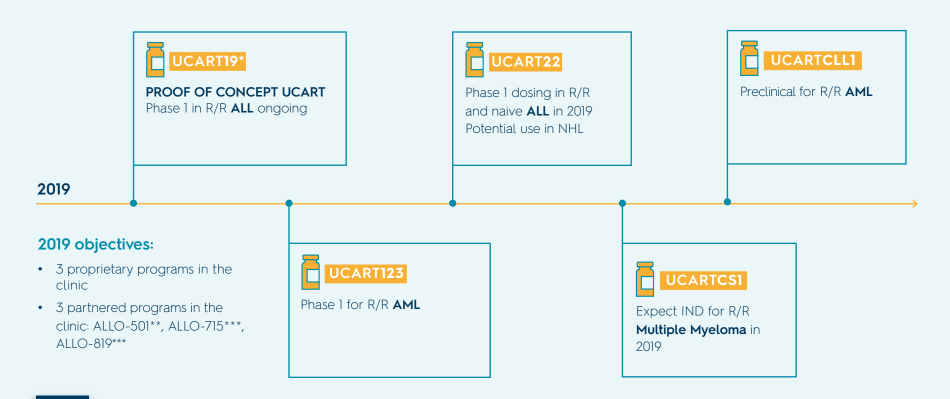


Preclinical evidence:

- Strong anti-tumor effect in mice
- Potential engraftment enhancement



BUILDING THE FUTURE OF ALLOGENEIC CAR T-CELL THERAPY



- * (
 - * UCART19 is exclusively licensed to Servier and under a joint clinical development program between Servier and Allogene
 - ** ALLO-501 is exclusively licensed to Servier
 - *** ALLO-715 and ALLO-819 are exclusively licensed to Allogene

TALEN® GENE EDITING - ADVANTAGES

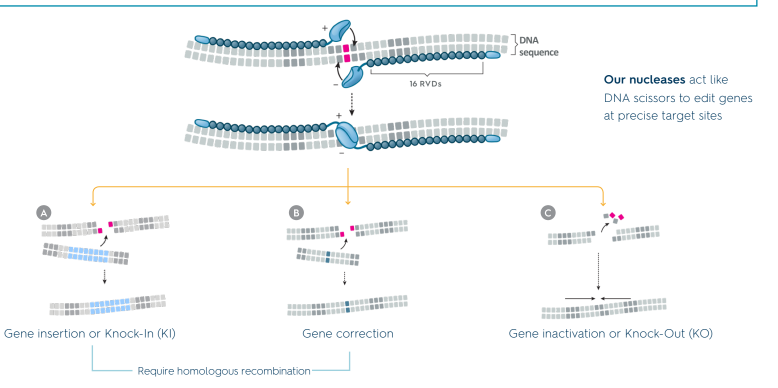
TALEN®:

Driven by protein/DNA

interactions to work on potential off-site cleavage

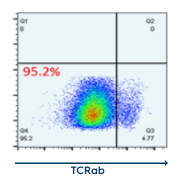
Releases DNA ends accessible to homologous recombination to perform gene insertions and corrections

Over 25 years of building a strong patent portfolio with umbrella patents on gene editing



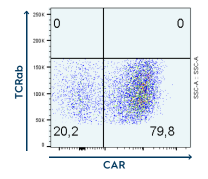
OPTIMIZING YIELD THROUGH HIGHEST GENE EDITING EFFICIENCY

High Knock-Out and Knock-In efficiency and specificity



95.2% single targeted gene Knock-Out

- → TRAC Knock-Out
- → High Specificity
- → Prevents GvHD



~80% single gene integration

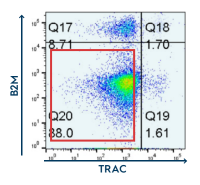
- → CAR Knock-In at TRAC Locus
- → High specificity
- → Enables efficiency

Enables efficiency & protection from GvHD



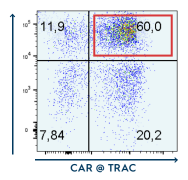
POWER OF TALEN® GENE EDITING: MULTIPLEXING GENE REPLACEMENT

Multiple advantages from combined Knock-Out, Knock-In



88% double targeted gene Knock-Out

- → TCR and B2M
- → B2M Knock-Out exposes cells to potential killing by NK cells – which is prevented as shown



60% double targeted gene insertion

- → CAR insertion at TCR
- → NK inhibitor at B2M
- → Provides protection from NK cell-mediated rejection

Provides protection from GvHD and avoids rejection

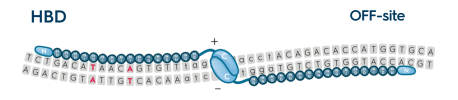


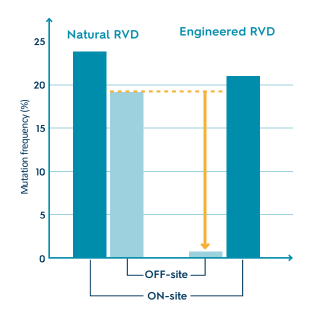
WITH TALEN® WE CONTROL OFF-TARGET CLEAVAGE

Discrimination between ON and OFF-site prevents OFF-site cleavage

Utilization of engineered RVDs to discriminate HBB* and HBD** loci preventing unwanted OFF-site cleavage



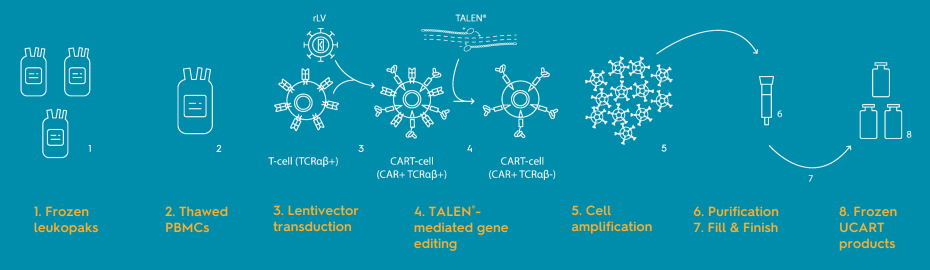






^{**} HBD - Hemoglobin subunit delta

UCART MANUFACTURING



- → 5 years of experience in allogeneic CAR T manufacturing
- → Validated gene editing technology for cell manufacturing
- → 4 UCART product candidates manufactured so far
- → Full QC system in place, 3 products cleared for 5 clinical trials by the U.S. and E.U. agencies



BUILDING 2 STATE-OF-THE-ART PLANTS TO SECURE AUTONOMY

SMART - Starting MAterial Realization for CAR-T products

- ~14,000 sqft in-house manufacturing in Paris, France
- Raw material for clinical supply: plasmids and vectors
- Potential to supply commercial starting material
- Operational "go-live" targeted in 2020

IMPACT - Innovative Manufacturing Plant for Allogeneic Cellular Therapies

- ~60,000 sqft facility located in the U.S. East Coast
- Production of clinical and commercial UCART products
- Operational "go-live" targeted in 2021



ANTICIPATED 12-MONTH MILESTONES

- 12 months -----

Clinical programs: -

UCART19: Phase 1 updates in R/R ALL in 2019 (licensed to Servier)

UCART123: Phase 1 for R/R AML Expansion phase expected in 2020

UCART22: Expect Phase 1 first patient dosing in R/R ALL in 2019

UCARTCS1: Expect IND in 2019 in R/R MM

ALLO715: Expect IND in 2019

Manufacturing: _____

Focusing on refinements to improve efficiency and capacity to support future commercial launch of UCART products

Internalizing large parts of our proprietary manufacturing chain for clinical supplies: SMART plant in Paris, France

Planning to start building a proprietary GMP, commercial scale manufacturing facility in 2019: IMPACT plant in the U.S.

Gene editing: -

Explore applications into new areas: solid tumors and outside oncology space



CELLECTIS HIGHLIGHTS



INDUSTRY LEADER IN GENE EDITING & ALLOGENEIC CAR T (UCART) TECHNOLOGY

- → First clinical proof-of-concept: UCART19 treated the first pediatric ALL patient in June 2015
- → Innovative gene editing (TALEN®) platform: to generate best-in-class allogeneic CAR T-cells
- → Bringing innovative off-the-shelf therapies to a broader market, without treatment delays



BEST-IN-CLASS MANUFACTURING

- → Scalable, efficient, greater consistency and potency
- → Two facilities being built to ensure manufacturing autonomy



PARTNERSHIPS WITH LEADERS: UP TO \$3.9B IN POTENTIAL MILESTONES PLUS ROYALTIES

- → **UCART19** Licensed to Servier (U.S. rights to Allogene)
- → 15 licensed targets to Allogene and other licensed targets to Servier



ROBUST PROPRIETARY PIPELINE

- → **UCART123** Phase 1 AML ongoing; dose escalation in AML in 2019; *wholly-controlled asset*
- → UCART22 Phase 1 first dosing in ALL in 2019; wholly-controlled asset
- → UCARTCS1 IND in Multiple Myeloma in 2019; wholly-controlled asset
- → UCARTCLL1 Preclinical development for AML; whollycontrolled asset



FINANCIAL POSITION:

- → Cash into 2022
- → ~70% ownership of CLXT*



* As of September 30, 2018 P30

THE CELLECTIS GROUP



~70%* ownership



- → NASDAQ: CLLS
- → EURONEXT GROWTH: ALCLS
- → \$476M** cash as of September 30, 2018
- → Expected to fund operations into 2022
- → Based in Paris, France & New York, USA
- → Patient focused

- → NASDAQ: CLXT
- \rightarrow \$101.8M cash as of September 30, 2018
- → Based in Minnesota, USA
- → Consumer focused
- → High value asset

Gene editing is the link



^{*} As of September 30, 2018

^{**} Including \$101.8M of cash from Calyxt for plant activities

THANK YOU

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