











May 2024

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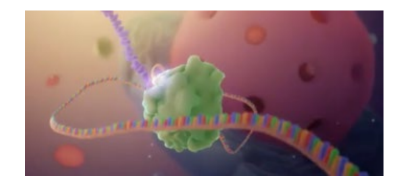
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ARCTURUS"

Arcturus Therapeutics



Global Late-Stage Clinical mRNA Medicines Company



Nasdaq: ARCT

Headquarters: San Diego, CA

Founded: 2013



mRNA Medicine Candidates

LUNAR-OTC Ornithine Transcarbamylase Deficiency

LUNAR-CF Cystic Fibrosis

Additional Earlier Stage Programs

Multiple Strategic Partners













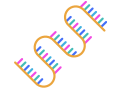
Proprietary mRNA Technologies Driving Therapeutic Programs

Broad Intellectual Property Portfolio

mRNA Technology

mRNA for protein replacement
Self-amplifying mRNA (STARR®)
low-dose vaccine technology





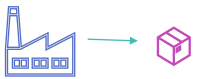
LUNAR® Delivery

Hepatocytes – *intravenous*Myocytes – *intramuscular*Bronchial Cells – *inhaled*



Manufacturing Know-How

mRNA Drug Substance Production
mRNA Purification
LNP Drug Product Production
Fill Finish / Lyophilization







Pipeline of Arcturus-Owned mRNA Therapeutic Candidates

Franchise	Candidate	Funded By	Indication	Global Prevalence	Upcoming Milestone
Hepatic	LUNAR-OTC (ARCT-810)	ARCTURUS° therapeutics	Ornithine Transcarbamylase Deficiency (OTC)	> 10,000	Phase 2 Interim Data H1 2024
Respiratory	LUNAR-CF (ARCT-032)	CYSTIC FIBROSIS FOUNDATION®	Cystic Fibrosis	85,000-100,000	Phase 1b Interim Data H1 2024



Pipeline of Partnered Self-amplifying mRNA Vaccines

Candidate	Partner	Indication	Stage
Kostaive® (ARCT-154) Monovalent: Ancestral	CSL	COVID-19	Approved in Japan; EMA MAA Approval Q3 2024
Kostaive® Bivalent (ARCT-2301) Ancestral / Omicron BA.4/5	CSL	COVID-19	Phase 3
Kostaive® XBB.1.5 (ARCT-2303) Monovalent: XBB.1.5	CSL	COVID-19	Phase 3
LUNAR-FLU (ARCT-2138) Quadrivalent	CSL	Seasonal Influenza	Phase 1
LUNAR-FLU (ARCT-2304) Pandemic	BARDA	Pandemic Influenza	Pre-clinical



CSL: Arcturus Therapeutics Global Vaccine Partner



- \$13.3 Billion USD Annual Revenue
- Operating in 40+ Countries Worldwide
- 32,000+ Employees Worldwide
- 13 Phase III programs including ARCT-154
- Focused on four strategic technology platforms plasma protein; recombinant technology; cell and gene therapy; and vaccines
- Therapeutic areas of focus of immunology, hematology, respiratory, cardiovascular and metabolic, transplant, nephrology and vaccines

CSL Segirus A World Leader in Flu Vaccines • \$2.03 Billion USD Annual Revenue

CSL Segirus is one of the Three Core Businesses of CSL





Deal Value: Up to \$4.5 billion

- Collaboration combines CSL's global vaccine commercial and manufacturing infrastructure with Arcturus' expertise in mRNA design and modification, LUNAR[®] lipid nanoparticle (LNP) technology and manufacturing know-how.
- Deal terms encompass the development, manufacture, and commercialization of mRNA-based vaccines targeting COVID-19, Influenza and three additional respiratory infectious disease vaccines.



Terms of the Partnership





\$200 million

\$1.3 billion

\$3.0 billion

Upfront Payment

Development Milestones

Commercial Milestones

40% profit sharing* for COVID-19 vaccines

Up to **double digit royalties** for influenza and three additional respiratory infectious disease vaccines

* 40% share of net profits (i.e., gross profits less COVID development costs)





Meiji: Background Information



The Meiji Group provides food and pharmaceuticals indispensable to their customers

- \$7.9 Billion USD Net Sales (As of March 31, 2023)
- 113 Locations Worldwide with 17,290 Employees

meiji Meiji Seika Pharma Co., Ltd.

Meiji Seika Pharma provides antibacterial drugs, vaccines, central nervous system drugs, and generic drugs

- \$1.4 Billion USD Net Sales (As of March 31, 2023)
- Received rights in Q4 2022 to conduct ARCT-154 clinical study in Japan
- Granted significant subsidy from Japanese government in Q4 2022
- Entered into agreement with CSL Seqirus in April 2023, responsible for obtaining regulatory approval, distribution, sales and marketing of ARCT-154 in Japan

Meiji Seika Pharma, a Subsidiary of Meiji Holdings Co. Ltd., Funded and Conducted the ARCT-154 Phase 3 Comparator Booster Study and Obtained Regulatory Approval in Japan





ARCALIS: Arcturus' Joint Venture mRNA Manufacturing Partner





ARCALIS is a CDMO Specializing in Manufacturing of mRNA Vaccines and Therapeutics

- Joint Venture Founded in 2021
- Major Equity Owners: Axcelead & Arcturus, subject to dilution
- Meiji Seika Pharma is collaborating with ARCALIS for domestic mRNA vaccine production

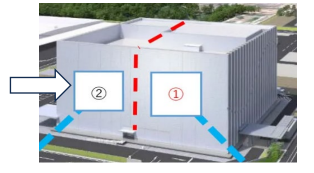


ARCALIS' cGMP mRNA Drug Substance Manufacturing Plant

- Completed July 2023; Located in Minamisoma City, Japan
- Capacity: Up to 5 kg in bulk mRNA drug substance per year
- 78,059 sq ft (7,252 sq m) floor space

ARCALIS' cGMP mRNA Drug Product Manufacturing Expansion

Capacity: 30 L (3 Lines); building to 100 L (2 Lines)



Kostaive® Phase 3 Clinical Studies



Kostaive® (Monovalent, ARCT-154)

Phase 3 Non-inferiority safety and immunogenicity trial

- Kostaive® administered at an 83.3% lower dose than Comirnaty® (N = 828)
- 50% of participants received Kostaive® (5 mcg); 50% of participants received Comirnaty® (30 mcg)
- Conducted in Japan

Achieved Primary Endpoint of non-inferiority of neutralizing antibody response against SARS-CoV-2 Ancestral strain compared to Comirnaty[®] Achieved Secondary Endpoint of superiority of ARCT-154 in neutralizing antibody response against SARS-CoV-2 Omicron BA.4/5 variantIncreased immunogenicity associated with ARCT-154 versus Comirnaty[®] at Day 29, with a geometric mean ratio of neutralizing antibodies against the vaccine strain of 1.43

Generally safe and well tolerated Phase 3 Study published in *The Lancet Infectious Diseases*¹



Kostaive® (Bivalent, ARCT-2301)

Bivalent Kostaive® (ARCT-2301: ancestral D614G and Omicron BA.4-5)

- Results consistent with monovalent Kostaive®
- Phase 3 clinical booster vaccination study was also conducted in Japan

Bivalent Kostaive® was assessed in comparison with bivalent conventional mRNA vaccine (Comirnaty®):

- Day 29 superiority of neutralizing antibody response against SARS-CoV-2 Ancestral strain was established
- Day 29 superiority of neutralizing antibody response against SARS-CoV-2 Omicron BA.4/5 subvariant was established
- Day 29 neutralizing immune response against SARS-CoV-2 Omicron XBB.1.5 subvariant was higher compared to Comirnaty

Kostaive® Received Approval Nov 2023 from Japan's Ministry of Health, Labor and Welfare (MHLW)



Historic Approval of World's First sa-mRNA Product



CSL-Arcturus Collaboration Results in Groundbreaking Approval

First Arcturus Approval

ARCT-154, self-amplifying mRNA COVID vaccine, was approved in Japan by the MHLW in November 2023

The STARR® vaccine was created, optimized, clinically developed and approved in under 4 years



Enduring Vaccine with Strong Clinical Data

Approval based on positive clinical data from several ARCT-154 studies

• 18,000+ subjects have received sa-mRNA COVID vaccines

Partner **Meiji Seika Pharma** advanced the MHLW approval and is the exclusive distributor of the sa-mRNA vaccine in Japan







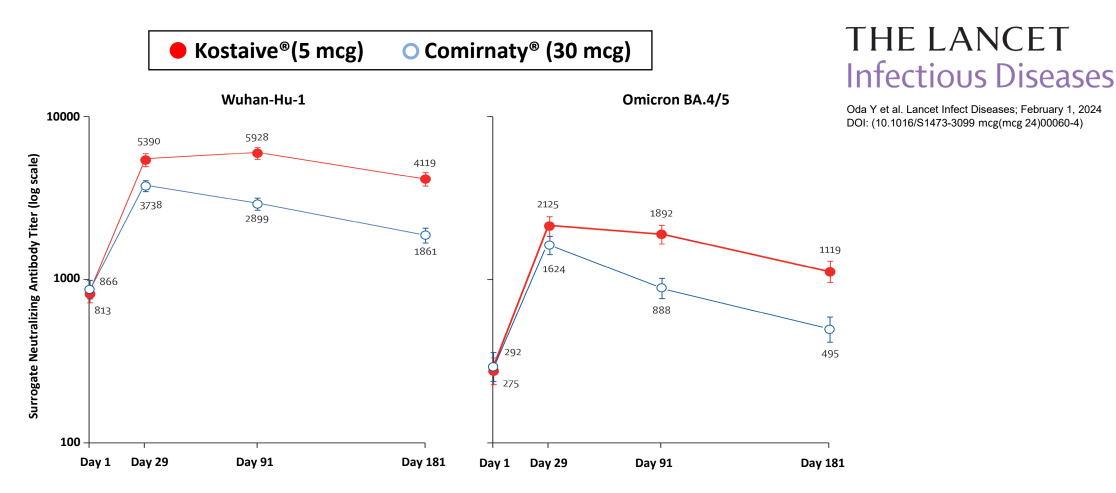
What's the next regulatory approval milestone?

European Medicines Agency (EMA) has validated the marketing authorization application (MAA) for ARCT-154

Kostaive®: More Durable Post-boost Immune Response



Phase 3 Persistence Data Comparing ARCT-154 (5 mcg) to Comirnaty® (30 mcg)



Comirnaty® is the brand name of BNT162b2



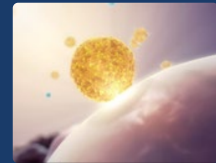


LUNAR® - Lipid Nanoparticle (LNP) Delivery Technology

Proprietary, Biodegradable, Optimized for Each Cell Type



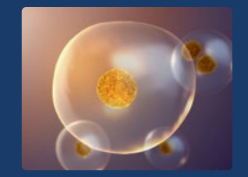
LUNAR[®] interacts with cell membrane



mRNA release



LUNAR® internalized inside endosome



mRNA translated into protein of interest







Ornithine Transcarbamylase (OTC) Deficiency

ARCT-810 Market Opportunity



The most common urea cycle disorder

The urea cycle converts neurotoxic ammonia to water-soluble urea that can be excreted in urine

Deficiency in OTC causes elevated blood ammonia, which can lead to neurological damage, coma, and death

10,000 worldwide prevalence

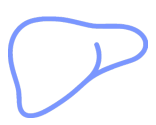


Unmet Medical Need

Present standard of care involves a strict diet (low protein, high fluid intake) plus ammonia scavengers (e.g. glycerol phenylbutyrate)

Present standard of care does not effectively prevent life-threatening spikes of ammonia

Severe OTC Deficiency patients are referred for liver transplant, currently the only cure



LUNAR-OTC Aims to Restore Enzyme Function

Establishing expression of OTC enzyme in liver has potential to restore urea cycle activity to detoxify ammonia, preventing neurological damage and potentially removing need for liver transplantation

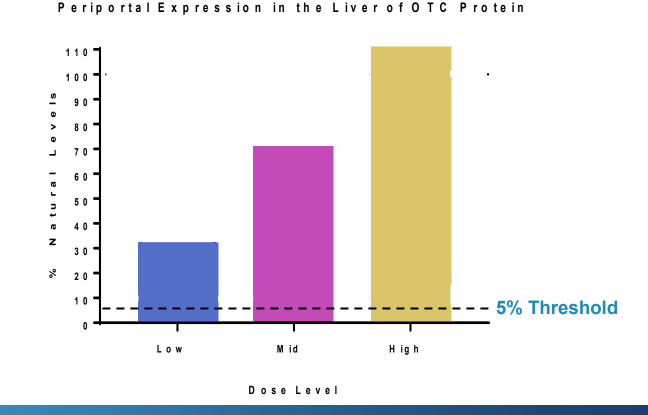


LUNAR-OTC



Exceeds Target of 5% Enzyme Replacement in OTC-Deficient Mouse Model

- OTC deficiency impacts ureagenesis (ammonia detoxification)
- The main site of ureagenesis is the periportal region of the liver*
- The critical threshold of 5% residual enzymatic OTC activity helps avoid severe manifestations of the disease (neonatal coma, mortality)*



LUNAR-OTC Treatment Increases OTC Expression in Mouse Periportal Hepatocytes (Main Site of Ureagenesis)



ARCT-810 Clinical Update



Received FDA Fast Track Designation & Rare Pediatric Disease Designation in June 2023

Phase 1 (NZ) Study in Healthy Volunteers

Completed dosing up to 0.4 mg/kg, total number of subjects N = 24, generally safe and well tolerated

Phase 1b (U.S.) Single Ascending Dose (SAD) Study in OTCD Adults

- Completed enrollment and dosing of all cohorts (N=16)
- Dose cohorts were 0.2, 0.3, 0.4 and 0.5 mg/kg; no serious or severe adverse events

Phase 2 (UK and EU) Single and Multiple Ascending Dose, Placebo-controlled Study in OTCD Adolescents & Adults

- Enrolling up to 24 subjects in two dose cohorts
- Up to 6 bi-weekly doses for each participant with the following endpoints
 - Primary Endpoints: Safety and tolerability
 - Secondary Endpoints: PK and PD (ureagenesis assay, plasma ammonia: 24-hr profile and peak level)
 - Exploratory Endpoints: Plasma amino acids and OTC enzyme activity; urine orotic acid



Cystic Fibrosis

A ARCTURUS

ARCT-032 Market Opportunity



Cystic Fibrosis

85,000-100,000 worldwide prevalence

Caused by mutations in the CFTR gene, resulting in poor chloride transport and dehydrated, sticky mucus in the airways

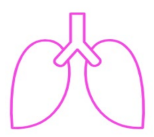
Chronic airway obstruction leads to infection and inflammation, which causes progressive airway damage and ultimately, respiratory failure



Unmet Medical Need

Highly effective CFTR modulators are not approved for treatment of all people with CF and may not be tolerated in others.

Standard of care therapies do not prevent the chronic, progressive loss of lung function that ultimately requires lung transplantation or leads to early death



LUNAR-CF Aims to Restore CFTR Function

An mRNA replacement therapy has the potential to produce wild-type CFTR into the lungs of CF patients, independent of genotype

Functional CFTR protein can restore chloride efflux in the airways, reducing mucus accumulation and airway damage and minimizing the progressive respiratory impairment observed in people with CF



ARCT-032 Clinical Update



Phase 1 Study in Healthy Volunteers (New Zealand)

- Objectives: Assess safety, tolerability and PK of ARCT-032 in healthy adults
- Completed dosing across 4 ascending single-dose cohorts (8 subject per cohort)
- Total number of subjects N = 32
- Safety and tolerability data supported transition to Phase 1b study

Phase 1b Study in Adults with Cystic Fibrosis (NZ) – Enrollment initiated October 2023

- Objectives: Assess safety, tolerability and PK of ARCT-032 in up to 8 adults with CF
- Status: First patient successfully completed two administrations of ARCT-032



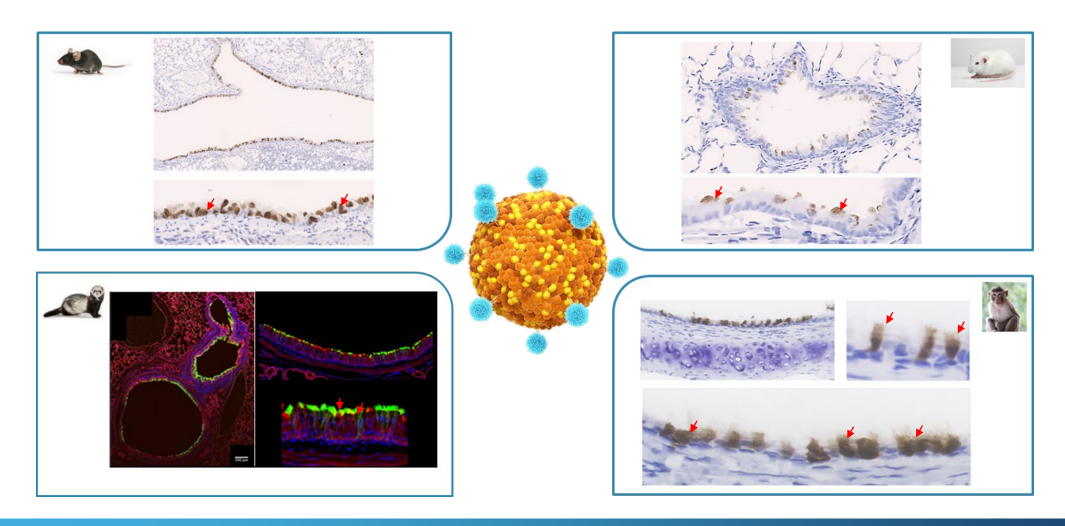
The Cystic Fibrosis Foundation increased its financial commitment to ~\$25 Million to advance ARCT-032

ARCT-032 has received Rare Pediatric Disease Designation and Orphan Drug Designation from the FDA and Orphan Medicinal Product Designation from the European Commission (EC)

LUNAR®-mRNA in Healthy Animals (four different species)



Successful delivery to airway epithelium; transduction demonstrated by Brown and Green staining



LUNAR®-mRNA in Cystic Fibrosis Ferret Model

ARCTURUS

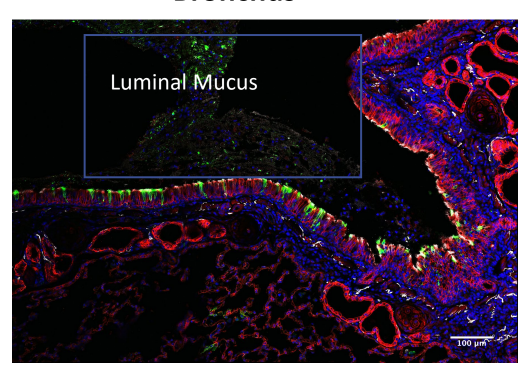
Successfully Transduces Epithelium in the Presence of CF Mucus



Trachea

Low-res High-res

Bronchus



Green denotes functional expression of protein (Cre)

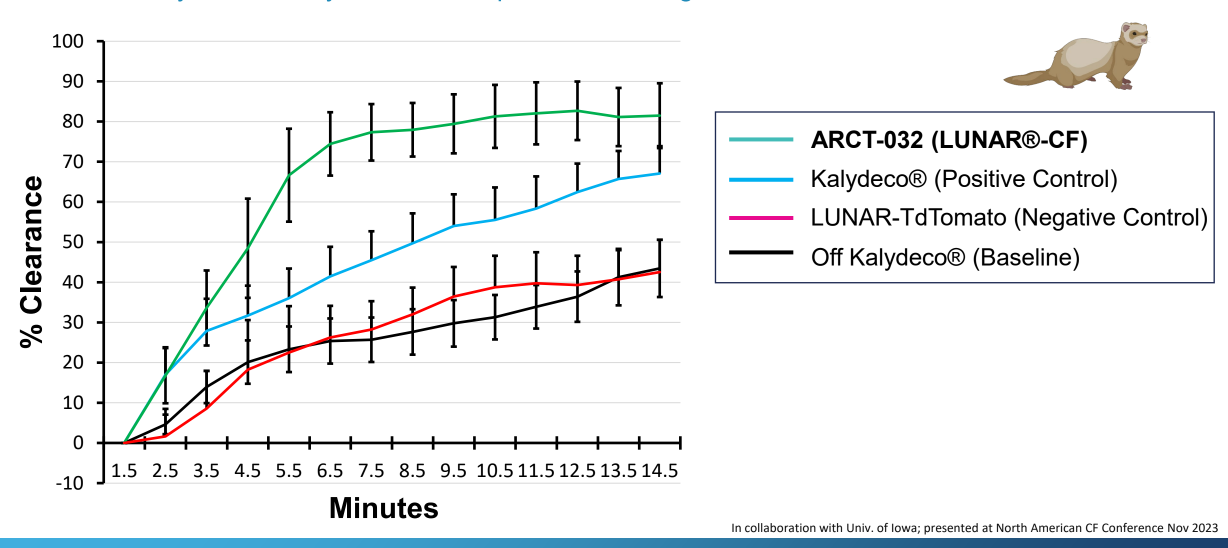
In collaboration with Univ. of Iowa; presented at North American CF Conference Nov 2023





ARCT-032 in a Kalydeco®-responsive CF Ferret Model (G551D)

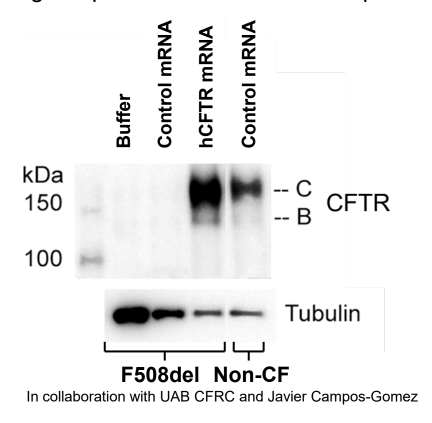
Proof of Activity: Mucociliary clearance improves after single administration of ARCT-032



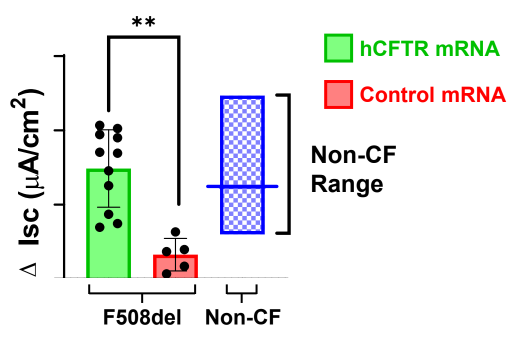


ARCT-032 Restores CFTR Expression & Function

High Expression Levels of CFTR protein



Restored chloride activity (chloride gradient)



**P<0.01; Data from two F508del donors; Using chamber studies performed with chloride secretory gradient

In collaboration with Univ. of Alabama-Birmingham; presented at North American CF Conference Nov 2022



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