

S01.2 Update on AAV-mediated CFTR gene delivery (4D-710)

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Disclosures

- Personal financial relationships with commercial interests relevant to medicine, within the past year:
 - As faculty at an institution that is part of the CF TDN, I am/have been site/national PI on studies for 4DMT, Vertex, and Eloxx.
 - I have done clinical trial consulting and/or advisory boards for Insmed, 4DMT, and Vertex.
 - I serve on a DMC for AbbVie.
- Personal financial support from a non-commercial source relevant to medicine, within the past year:
 - I have received grant funding from the CF Foundation.
 - I have no personal relationships with tobacco industry entities.
 - I serve as the adult patient care representative to the CFF Board of Trustees, and on the CF Foundation's Clinical Research Executive Committee, Clinical Research Advisory Board, and as immediate past chair of the CF TDN's Sexual Health, Reproduction and Gender Research-Working Group, on the scientific advisory board for Emily's Entourage, and on the ATS Scientific Grant Review, Awards and Clinical Problems Assembly Programming Committees.

Majority of PwCF are Variant-eligible for Highly Effective Modulator Therapy



Modulator Landscape

Up to 94% of people with CF could ultimately be eligible for a highly effective modulator

It's not known (exactly) how many pwCF are not taking modulators (probably >10%)

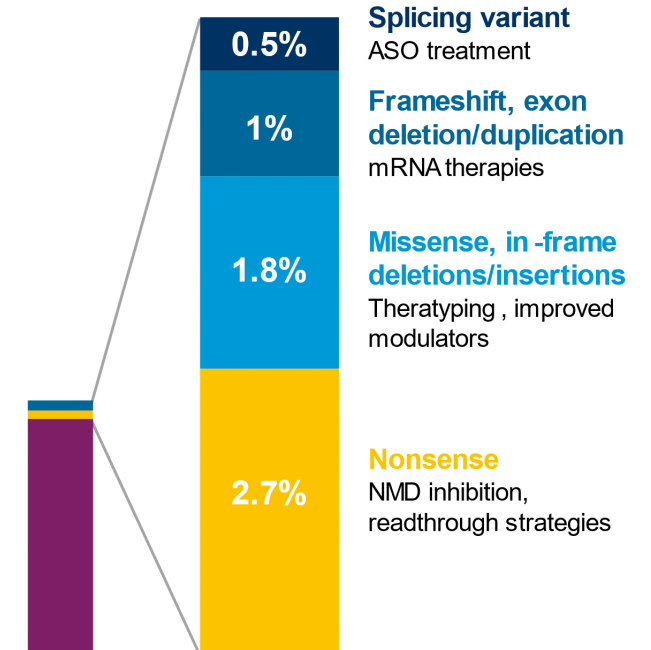
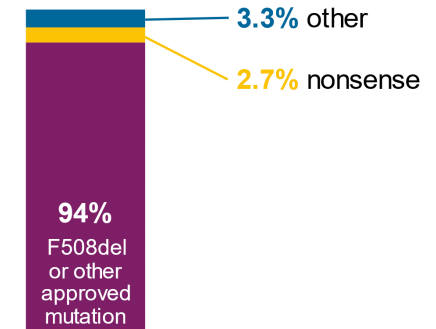
Slide courtesy of JP Clancy

<https://www.cff.org/Research/Developing-New-Treatments/CFTR-Modulator-Types/>

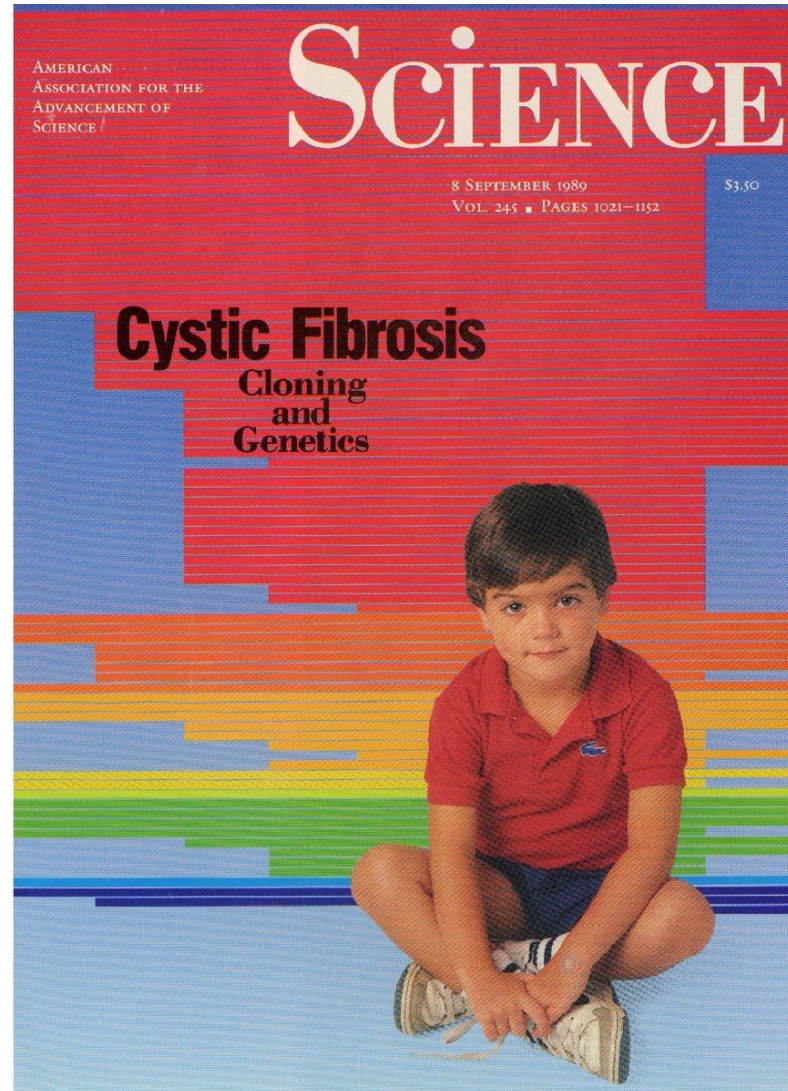
CFTR Modulator Landscape

Up to 94% of people with CF could ultimately benefit from a highly effective modulator (e.g., ETI = Trikafta)

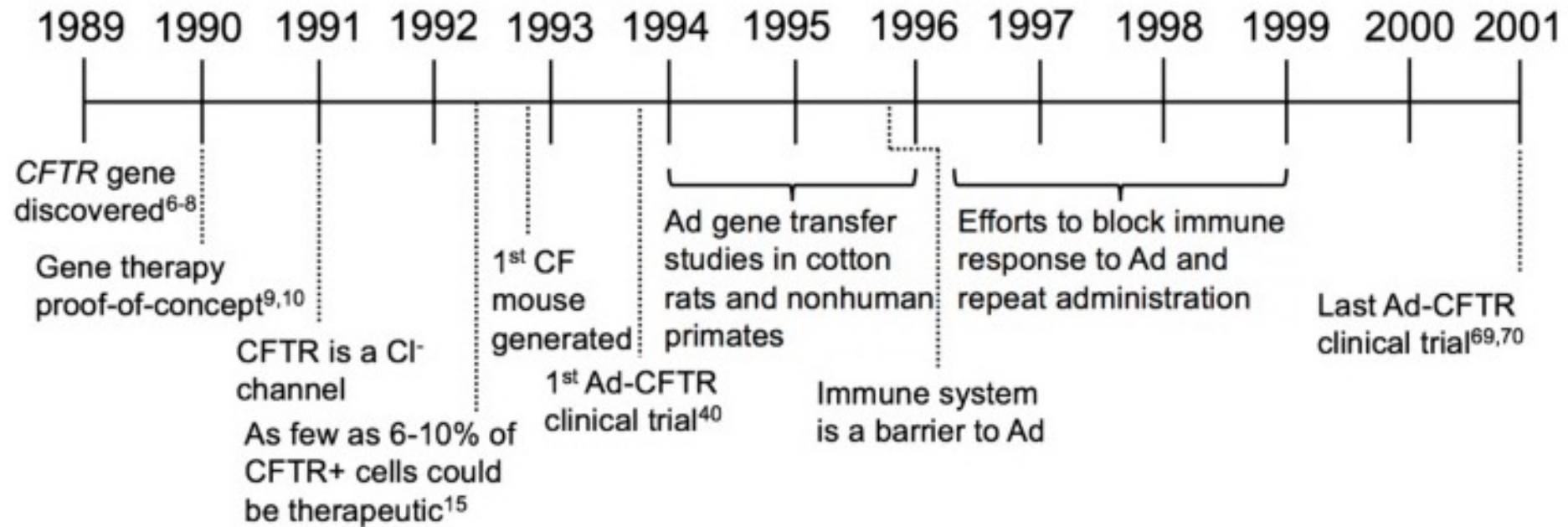
Data provided by Dr. Garry Cutting and Kara Raich, CFTR2, Johns Hopkins University



Cystic Fibrosis Transmembrane Conductance Regulator (CFTR) Gene Discovered



Timeline of CF Gene Therapy 1989-2001



Timelines of CF gene therapy eras: Important milestones impacting the CF field are represented in timelines at the beginning of each era. The timelines are intended to orient the reader to new developments relative to other events and are not comprehensive of all contributions to the field (1989–2001).

Failure of Conventional AAV Gene Therapy in CF Lung

PREVIOUS CLINICAL TRIAL EXPERIENCE WITH TGAAVCF

- Six trials with AAV2-based gene therapy (tgAAVCF) in upper¹⁻⁶ and lower⁵⁻⁸ airways
- Nasal and sinus administration (n=3 trials):
 - Participants dosed: 34
 - Safe and well tolerated
 - DNA: detected
 - **Transgene expression: detected**
 - **CFTR function: detected** (vs contralateral control)
- Aerosol to lung (n=3 trials):
 - Participants dosed: 84 (mild to moderate)
 - Safe and well tolerated
 - DNA: detected
 - **Transgene expression: NOT DETECTED**
 - **ppFEV₁: NO CHANGE vs CONTROLS**

1. Wagner JA et al. *Hum Gene Ther* 1998; 9: 889-909. 2. Wagner JA et al. *Lancet* 1998;351:1702-3. 3. Wagner JA et al. *Laryngoscope* 1999;109:266-74. 4. Wagner JA et al. *Hum Gene Ther* 2002;13:1349-1359. 5. Flotte TR et al. *Hum Gene Ther* 2003;14:1079-88. 6. Flotte TR et al. *Hum Gene Ther* 2005;16:921-8. 7. Aitken ML et al. *Hum Gene Ther* 2001;12:1907-16. 8. Moss RB et al. *Chest* 2004;125:509-21. 9. Moss RB et al. *Hum Gene Ther* 2007;18:726-32.

Advantages and Disadvantages of Conventional AAV as a Vector

Advantages

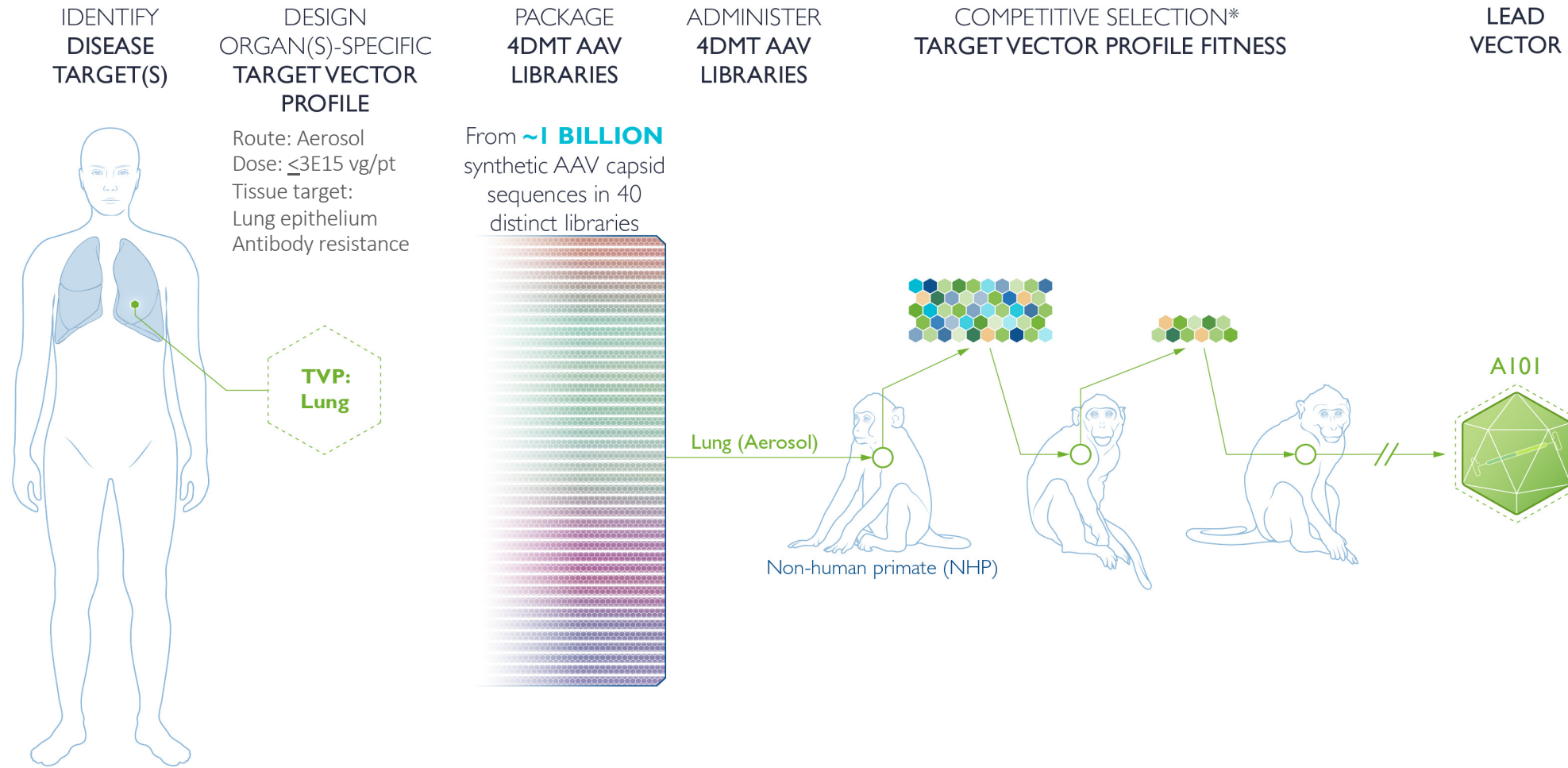
- Lower risk of pathogenicity
- Duration of expression (in non-proliferating cells)
- Broad range of target organs based on # of serotypes
- Lack of strong immune response to many serotypes
- Low risk of insertional mutagenesis

Disadvantages

- AAV receptor on basolateral (rather than apical) surface
- Small genome packaging capacity
- Potential for pre-existing or inducible Ab
- Potential for hepatotoxicity at high dose intravenous delivery
- Unclear if repeat dosing is possible

Therapeutic Vector Evolution: AI01 Aerosol Delivered Synthetic AAV

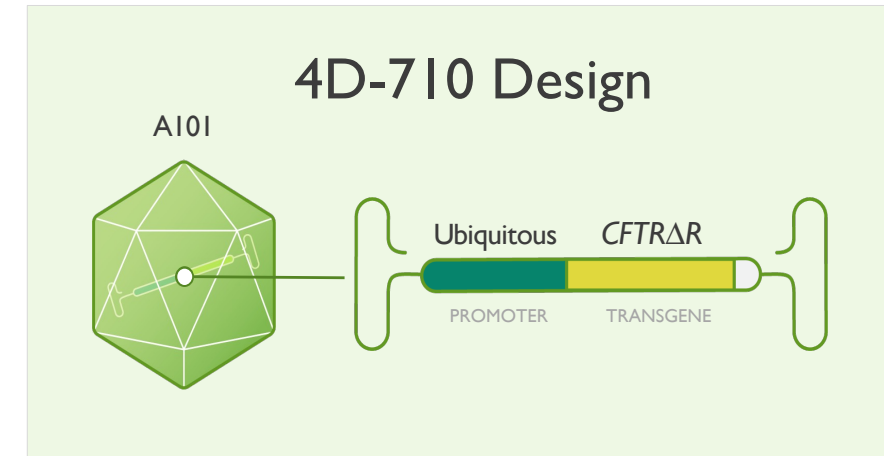
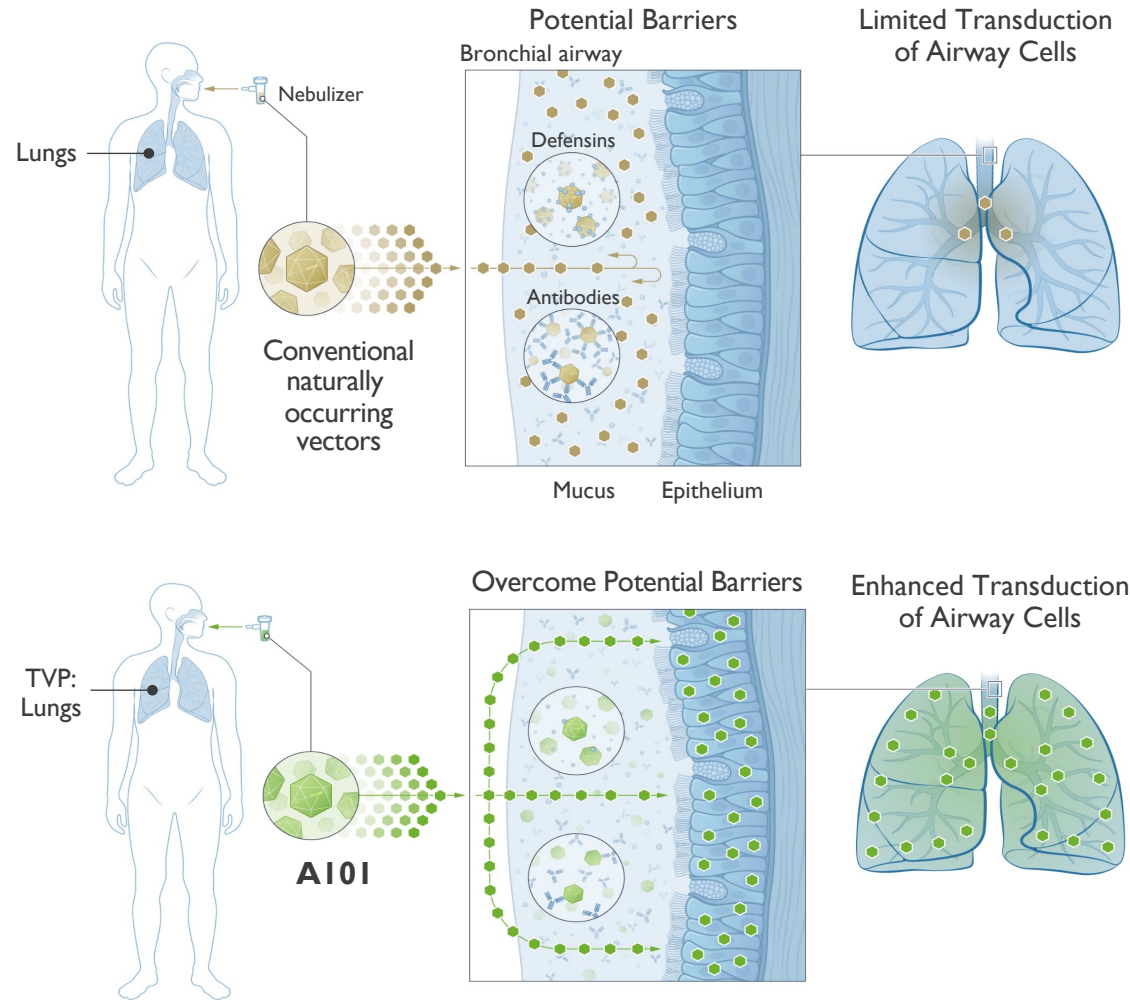
PROPRIETARY SYNTHETIC VECTOR DISCOVERY PLATFORM



TVP, Therapeutic Vector Evolution; vg, vector genome

4D-710: Next-Gen Aerosolized Genetic Medicine for Cystic Fibrosis Lung

AI01 TARGET VECTOR PROFILE & 4D-710 PRODUCT DESIGN AND KEY ATTRIBUTES



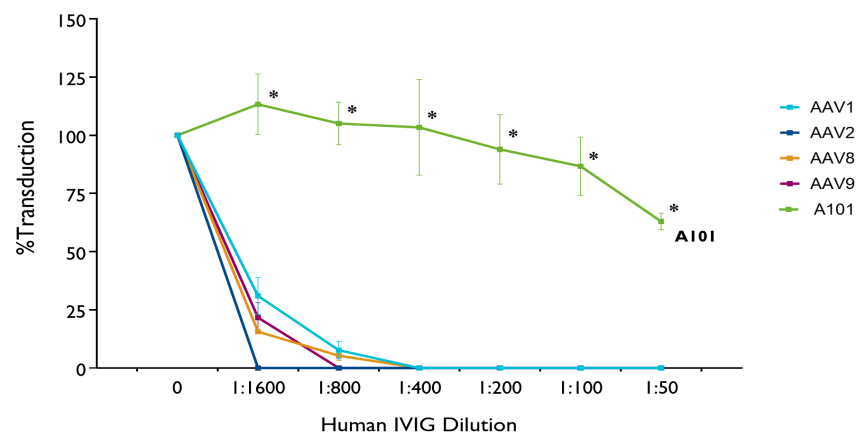
AI01 KEY ATTRIBUTES

- Mucus penetration efficient
- Resistance to pre-existing human AAV antibodies
- Transgene expression efficient
- Specificity for lung (>99.9%)

4D-710 Preclinical Characterization

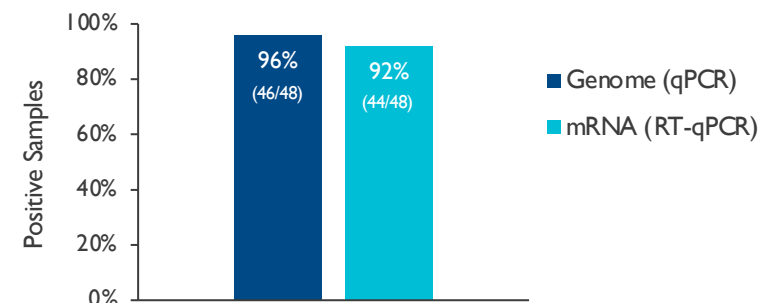
AI01 VECTOR RESISTANCE TO HUMAN IVIG, 4D-710 BIODISTRIBUTION, AND CFTR EXPRESSION IN PRIMATES

Human Antibody Resistance: IVIG

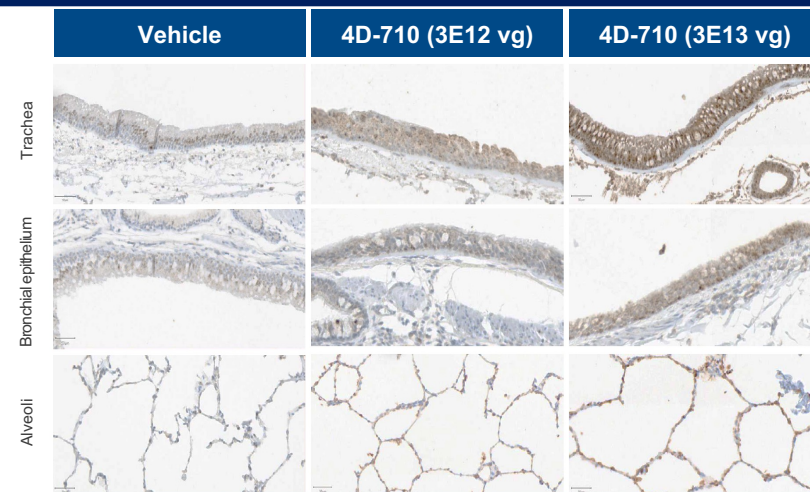
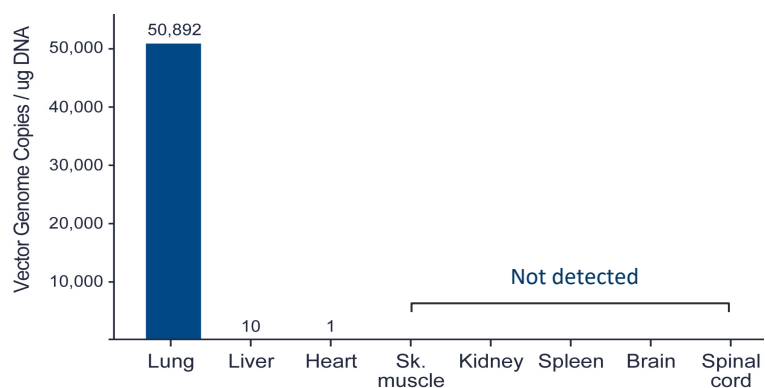


Delivery & Transduction: Aerosol NHP

4D-710 Biodistribution in NHP Lung (n=3 NHP; 48 samples)



Lung-Specific Delivery: Aerosol in NHP



CFTR immunohistochemistry staining of lung tissue samples from nonhuman primates, representative images (10x).

Calton M. American Thoracic Society International Conference, May 14-19, 2021. Abbreviations: NHP, nonhuman primate.

Next-Generation 4D-710 AAV Gene Therapy in Cystic Fibrosis:

4DMT APPROACH TO OVERCOMING HURDLES WITH TGAAVCF

- A101 novel synthetic vector:
 - Directed Evolution in primates for aerosol delivery
 - Efficient mucus penetration & transgene expression
 - High resistance to pre-existing human AAV antibodies
 - Lung retention >99.9%
- 4D-710 genetic medicine product:
 - Targeted and evolved vector (A101): tgAAVCF used AAV2
 - Strong promoter: tgAAVCF had no exogenous promoter

4D-710 Phase I/2 Clinical Trial

STUDY OBJECTIVES AND ELIGIBILITY CRITERIA

Study Objectives

- Evaluate a single nebulized dose of 4D-710 (1E15, 2E15 vg)
 - Safety, tolerability, and immunogenicity
 - Transduction and transgene expression in lung (bronchoscopy samples)
 - Impact on pulmonary function (ppFEV₁)
 - Impact on health-related quality of life
- Identify recommended Phase 2 dose

Key Inclusion Criteria

- Age ≥ 18 years
- Confirmed diagnosis: CF lung disease
- Ineligible for CFTR modulator therapy (per USPI) OR discontinued due to adverse effects
- % predicted FEV₁ $\geq 50\%$ and $< 100\%$
- Resting O₂ sat $\geq 92\%$ on room air

4D-710 Phase 1/2 Clinical Trial: Major Study Endpoints

- Primary endpoint:
 - Incidence and severity of adverse events
- Key secondary endpoints:
 - Transgene transfer and expression in bronchoscopy samples (biopsies, brushings)
 - Change in ppFEV₁ from baseline (through Month 12)
 - Change in Cystic Fibrosis Questionnaire-revised (CFQ-R) scores (through Month 12)

4D-710 Phase 1/2 Clinical Trial: Cohort 1 Participants

DEMOGRAPHICS AND BASELINE CHARACTERISTICS

Characteristic	Cohort 1 (IEI5 vg dose)		
	Participant 1	Participant 2	Participant 3
Age, y	36	24	20
Sex	Male	Male	Female
Race/ethnicity	Non-Hispanic white	Non-Hispanic white	Non-Hispanic white
CFTR modulator eligibility	Hypersensitivity	Ineligible variant	Ineligible variant
Historical sweat chloride, mmol/L	74	103	110
Percent predicted FEV ₁ (ppFEV ₁)	83	69	94

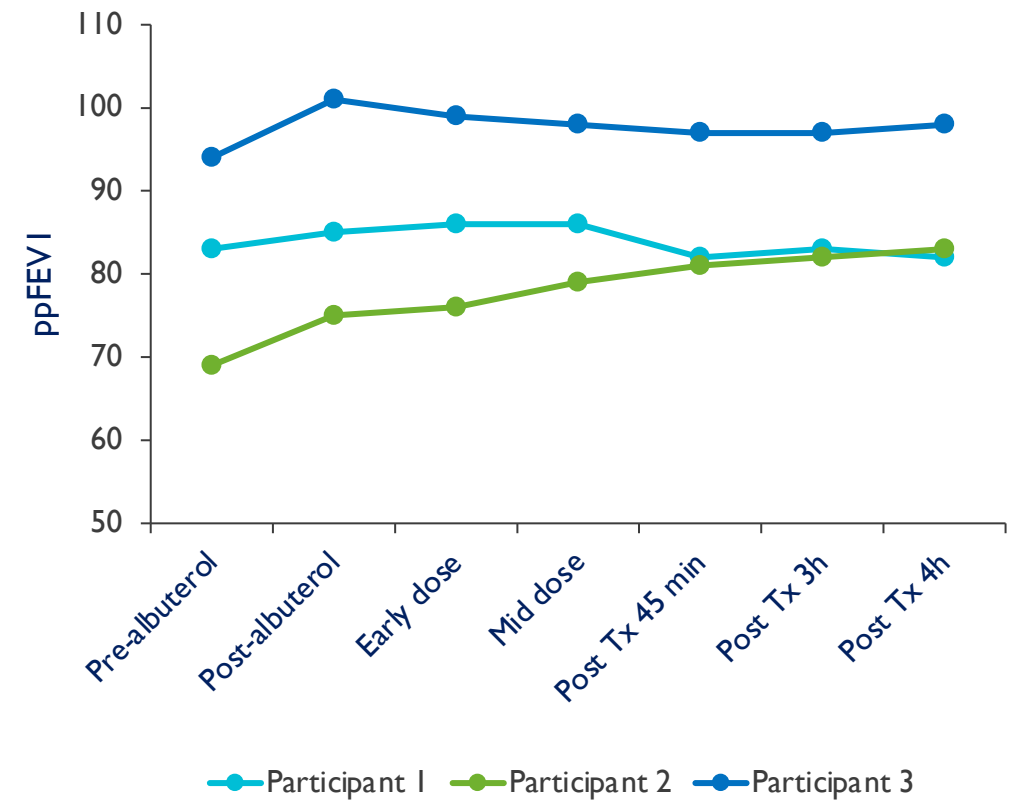
CFTR, cystic fibrosis transmembrane conductance regulator.

4D-710 Phase I/2 Clinical Trial: Cohort I Acute Safety

SERIAL SPIROMETRY AND ADVERSE EVENTS DURING NEBULIZATION OF 4D-710

- Full volume administered (1E15 vg)
- Participant 1: a mild, self-limited AE
 - Grade I dry throat, fatigue during nebulization
- No bronchospasm

Serial Spirometry During 4D-710 Dosing:
Through 4 Hours Post-Nebulization





4D-710 Phase 1/2 Clinical Trial: Cohort 1 Safety Summary

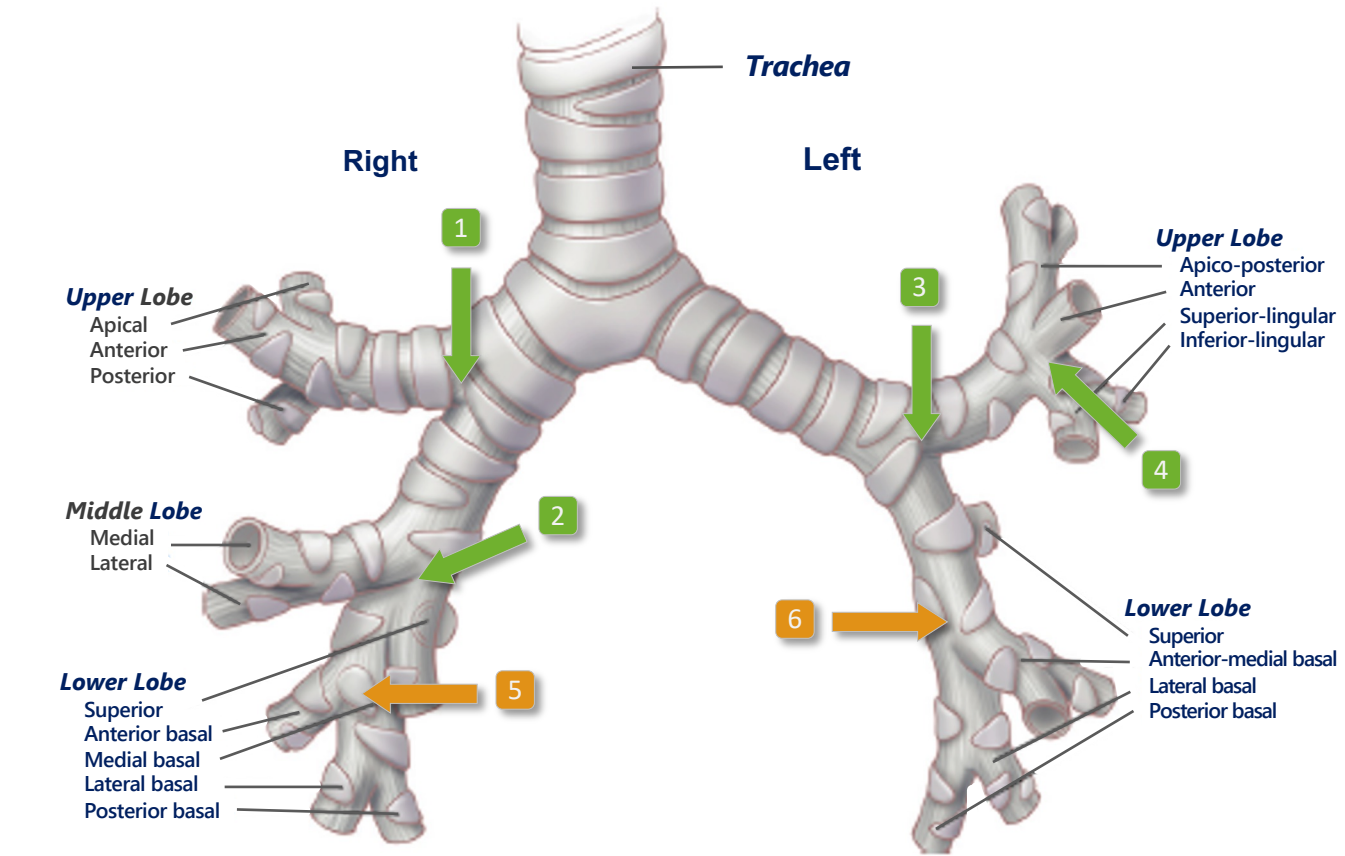
NO 4D-710-RELATED ADVERSE EVENTS AFTER COMPLETION OF DOSING

- No 4D-710-related adverse events
- No 4D-710-related serious adverse events
- No dose-limiting toxicities

4D-710 Phase 1/2 Clinical Trial: Bronchoscopic Sampling Plan

Bronchoscopy: Week 4*

Bronchoscopic Sampling Sites			Biomarker	
			ISH	PCR
Endobronchial biopsy				
	1	Right secondary carina		X
	2	Right middle lobe carina	X	
	3	Left secondary carina	X	
	4	Left upper lobe/lingula carina		X
Endobronchial brushing				
	5	Right lower lobe basal seg x 2	X	
	6	Left lower lobe basal seg x 2	X	



Minnich DJ, Mathisen DJ. Anatomy of the trachea, carina, and bronchi. *Thorac Surg Clin* 2007;17:571-85.

*Participant 3 bronchoscopy conducted at Week 8 due to pulmonary exacerbation (unrelated to study drug).

Widespread Transgene Delivery & Expression: Biopsies

CONSISTENT TRANSDUCTION ACROSS PATIENTS, LUNG REGIONS

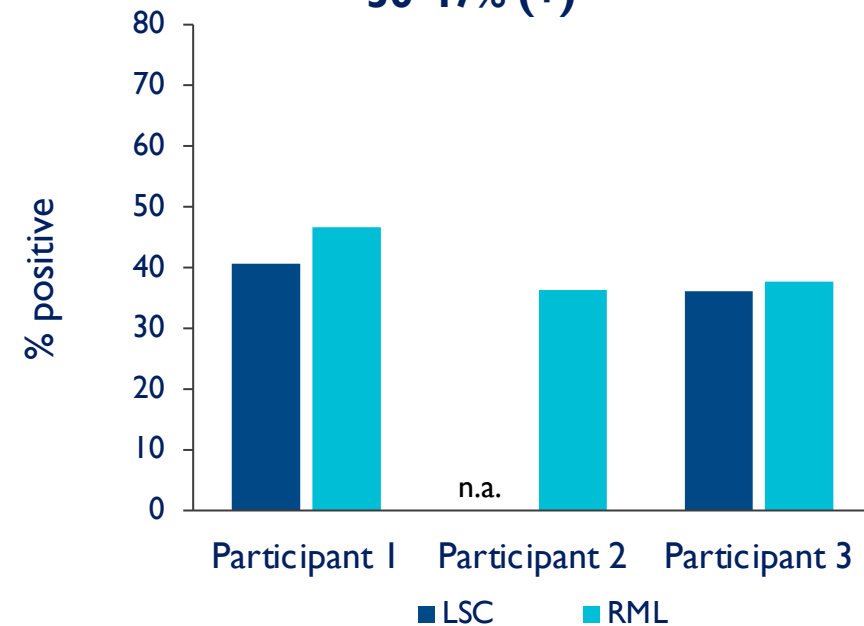
4D-710 DNA (+) Lung Biopsies

CFTR Δ R DNA qPCR¹ Results
5 of 5 biopsies (+) (All 3 pts)

Participant	Left Upper Lobe/ Lingula Carina DNA	Right Secondary Carina DNA
1	Positive	Positive
2	n.a.	Positive
3	Positive	Positive

4D-710 RNA Expression (+) Lung Biopsies

CFTR Δ R RNA ISH
% Positive Epithelial Cells²
36-47% (+)



¹ qPCR assay range: 25 – 25,000,000 copies.

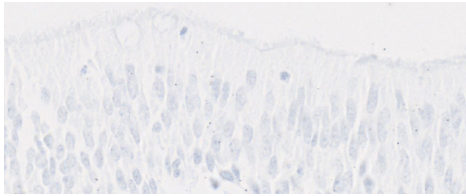
² Participant 2 LSC not sampled. Quantification by Visiopharm AI Machine Learning Analysis. SH, in situ hybridization; LSC, left secondary carina endobronchial biopsy; RML, right middle lobe endobronchial biopsy.

Widespread CFTR Expression in Lung: All 5 Biopsies (+)

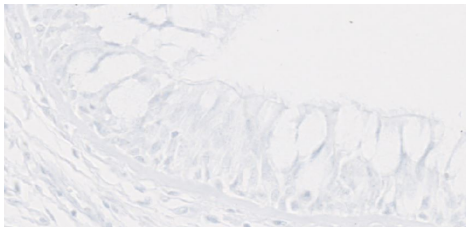
CFTR Δ R RNA EXPRESSION BY ISH

Controls

DAPB ISH
(negative control probe)



CFTR Δ R ISH
Untreated CF lung tissue

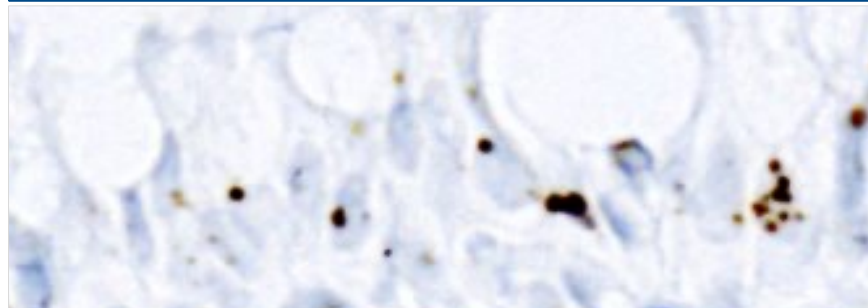


4D-710-Treated
CFTR Δ R RNA probe*

Left Secondary Carina
Endobronchial Biopsy (40X)



Left Secondary Carina
Endobronchial Biopsy (80X)



*Representative images from Participant 1. CFTR Δ R ISH signal observed in all evaluable biopsies from all 3 participants (Participant 2 LSC not sampled). ISH, in situ hybridization; LSC, left secondary carina

Multiple Bronchial Epithelial Cell Types Express CFTR Transgene

INDEPENDENT PATHOLOGISTS' REVIEW: *CFTR* Δ R RNA ISH LOCALIZATION

Transduced cell types*

1. Basal cells
2. Goblet cells
3. Columnar ciliated cells

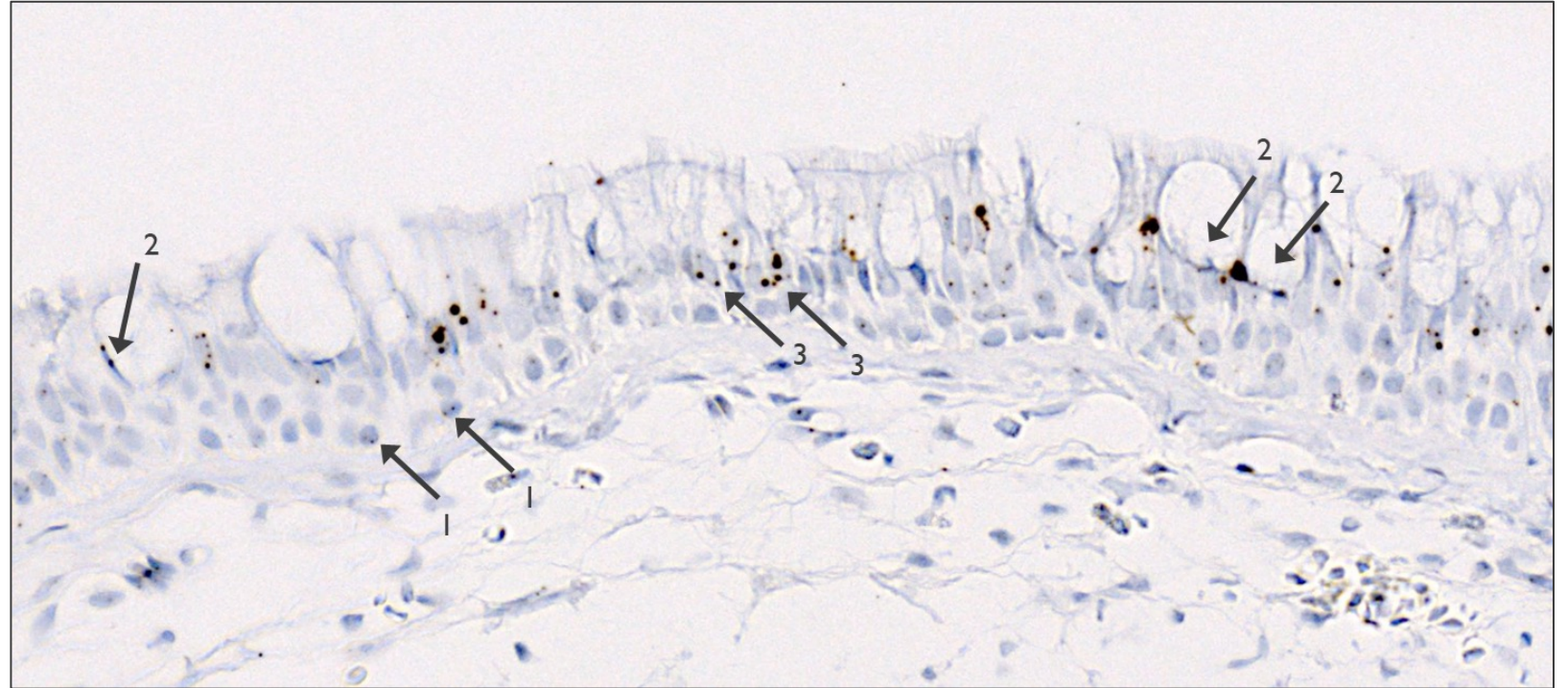


Image from Participant 1.*Assessed by 2 independent pathologists. ISH, in situ hybridization.

Widespread CFTR Expression in Lung: All 6 Brushings (+)

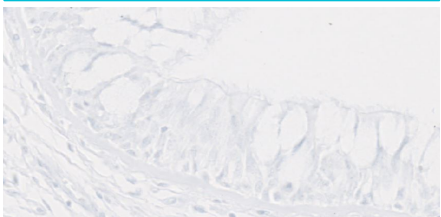
CFTR Δ R RNA EXPRESSION BY ISH

Controls

DAPB ISH BRUSHING
(negative control probe)

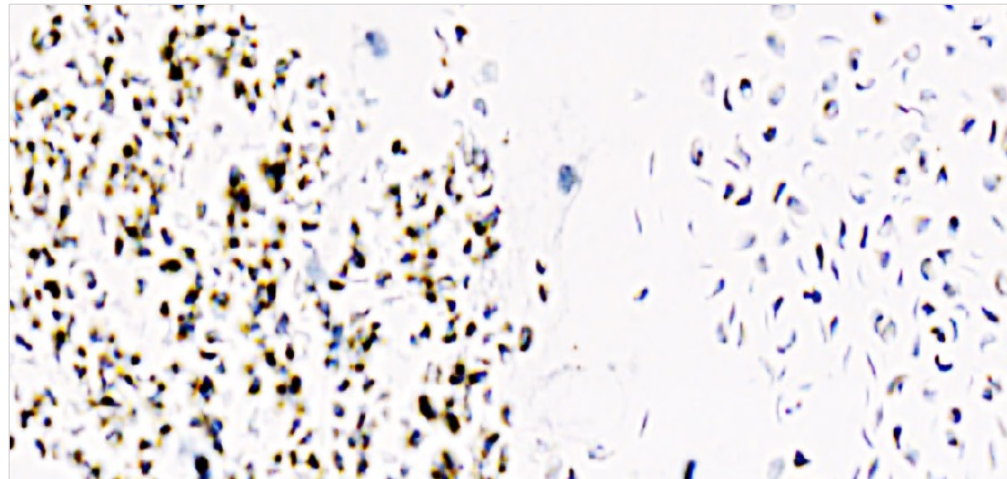


CFTR Δ R ISH
Untreated CF lung tissue



4D-710 Treated
CFTR Δ R RNA probe*

Right Lower Lobe



Left Lower Lobe



*Representative images from Participant 1. CFTR Δ R ISH signal observed in brushings from 2/3 patients (Participant 2 brushings unevaluable). ISH, in situ hybridization.

4D-710 Clinical Data Summary, Implications, and Next Steps

CLINICAL PROOF-OF-CONCEPT FOR SAFETY & WIDESPREAD TRANSGENE EXPRESSION

■ Cohort 1 Summary:

- No 4D-710-related AEs post-dosing
- Widespread CFTR expression (All 11 lung samples)
- ~40% of cells expressed CFTR (Multiple bronchial cell types)

■ Implications:

- A101 lung vector validation
- Clinical proof-of-concept: 4D-710 transgene delivery & expression

■ Next Steps:

- Cohort 2 enrollment underway (2E15 vg dose); Assessment of clinical activity (e.g. ppFEV₁; QoL)
- Consider 4D-710 combination therapy in individuals with CF on CFTR modulators

UNTIL IT'S DONE



- People with CF and their families
- Participating CF clinical and research centers
- JP Clancy and 4DMT for slides

