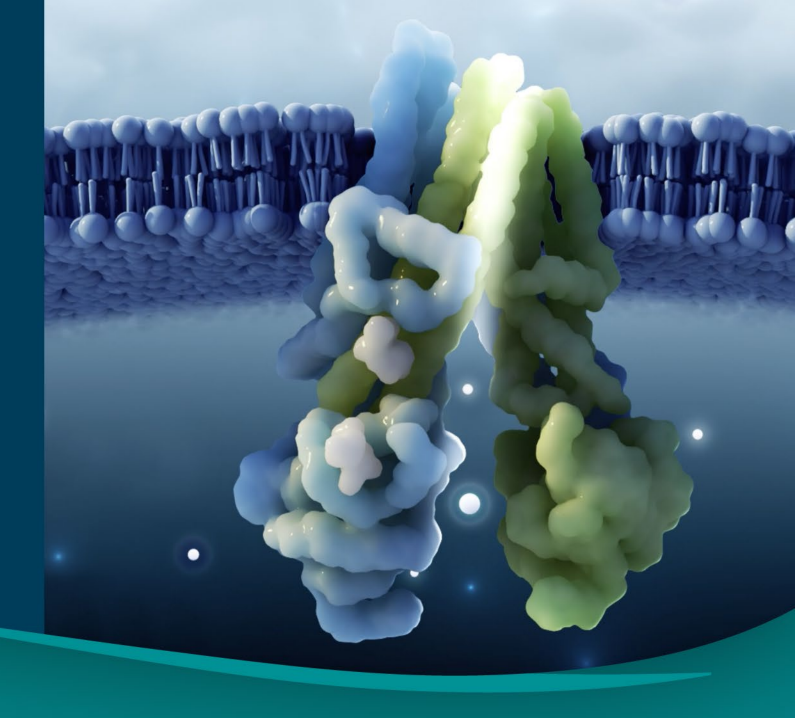
### Sionna Therapeutics

August 2024





# Sionna's differentiated approach focused on NBD1 has a clear path to POC with the potential to deliver best-in-class efficacy

### HIGH UNMET NEED IN LARGE MARKET



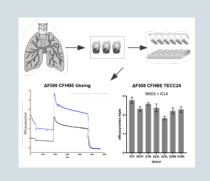
Despite current treatments, unmet need is high in the >\$10B market

### NBD1, THE HOLY GRAIL FOR CFTR



NBD1 is the key to deliver full CFTR function and has been considered 'undruggable'

### PREDICTIVE ASSAYS/BIOMARKERS



chloride biomarker consistently predict clinical efficacy driving near-term value inflection

### FRANCHISE DRIVES STRATEGIC OPTIONALITY



A deep clinical stage
pipeline of NBD1
compounds and
complementary modulators
can significantly raise the
efficacy bar



#### Led by proven management capable of disrupting the CF market



Mike Cloonan
Chief Executive Officer



BAIN (





Charlotte McKee, MD
Chief Medical Officer
VERTEX
PHARMACEUTICALS







**Elena Ridloff**Chief Financial Officer

ACADIA

ALEXION



Jen Fitzpatrick
General Counsel









**Vanya Sagar** Chief People Officer









Greg Hurlbut, PhD
Co-Founder
SVP, Discovery Research

sanofi genzyme



Mark Munson, PhD
Co-Founder
VP, Medicinal Chemistry

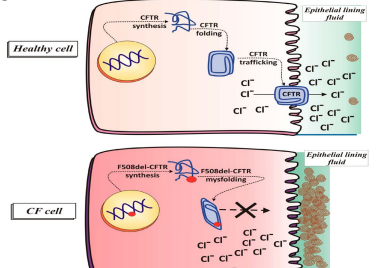
Sanofi AMGEN



# CFTR is a fully validated target, and unlocking NBD1 could deliver optimal clinical benefit in CF

#### The Biology of CF

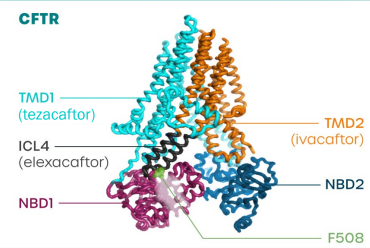
- Driven by mutation of the CF transmembrane conductance regulator (CFTR)
- CFTR is an epithelial chloride channel essential to the production of thin, freely flowing mucus in the airways, digestive system, and other organs



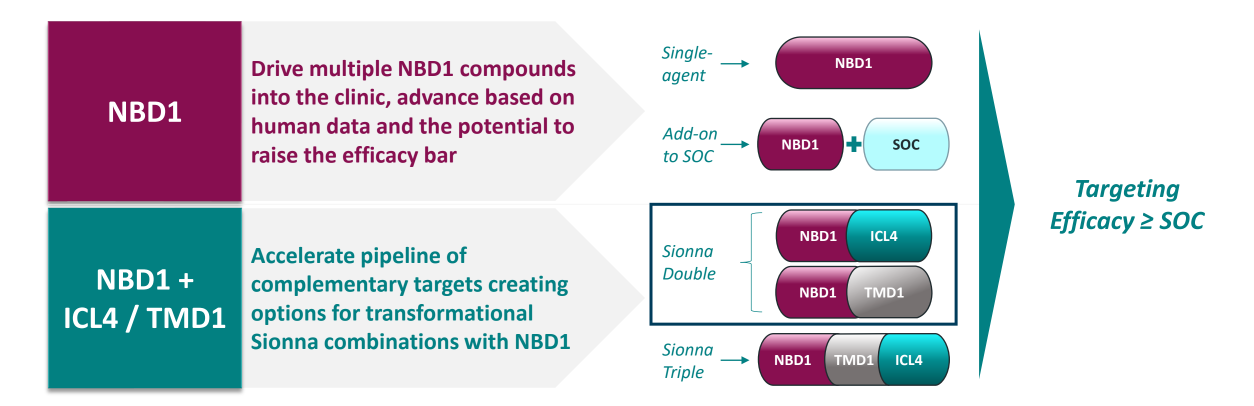
#### The Importance of NBD1

- F508 is present within CFTR's NBD1 domain
- F508del causes NBD1 to unfold at body temperature and weakens NBD1's interface with other regions; these defects cripple CFTR folding, trafficking and function
- None of the existing correctors or potentiators address both ΔF508-CFTR's assembly and its NBD1 instability defects
- ~90% of patients with CF have a F508del mutation

#### NBD1 is the key to full CFTR correction



# Sionna's strategy is to build a CF franchise across MOAs, anchored by novel NBD1, aimed at delivering higher efficacy than SOC



Vision: Deliver transformational option to fully normalize CFTR function, become the SOC



# We are well positioned to execute our strategy to deliver transformational CF treatment options and drive near-term value



### Well Funded with \$182M Series C

- Upsized Series C in March 2024; funds Sionna through YE26 and Ph 2a POC study
- Participation from all existing investors with three new strong investors added to syndicate



### Proven Execution with Ph 1 Advancement

- SION-638 (NBD1) completed
   Ph 1 study; compound is
   advanceable to Ph 2, subject to
   portfolio decision
- SION-109 (ICL4) Ph 1 ongoing on-track for completion by YE24

### Pioneering Next-Gen NBD1 Assets

- Completed GLP tox studies for both next-gen NBD1 compounds, SION-451 and SION-719
- GLP tox demonstrated high margins with no dose limiting toxicity
- Ph 1 initiated for '719 and '451

### Pipeline Expansion with AbbVie Licensing

- Exclusive WW rights for three clinical-stage compounds that expand and de-risk the combination options with NBD1
- Select the best dual combination option with NBD1
- Additional clinical assets become lifecycle development opportunities



# Licensing the ABBV modulators aims to expand and de-risk Sionna's combo development strategy

Compelling Activity in CFHBE Assay



NBD1 dual combinations with ABBV-2222, ABBV-3067, ABBV-2851, and SION-109 demonstrate the potential for superior efficacy to SOC in CFHBE assay

Accelerated and
De-Risked
Dual Combo Strategy



Galicaftor (ABBV-2222), a TMD1 modulator, has positive Ph 2 data in CF patients<sup>1</sup> Plan to advance ABBV-2222 and SION-109, if Ph 1 is successful, as potential dual combination options with an NBD1 stabilizer

LCM Options
with Additional
Clinical Assets



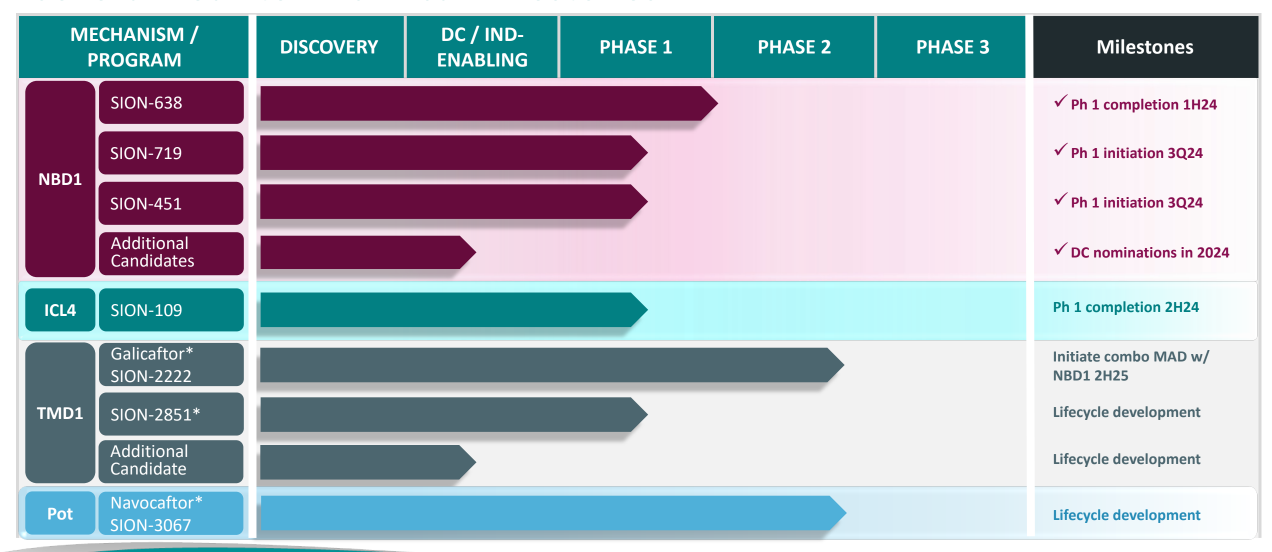
Navocaftor (ABBV-3067), a potentiator, also has positive Ph 2 data; ABBV-2851 is a Ph 1 TMD1 modulator

Both modulators provide lifecycle development opportunities



ClinicalTrials.gov (June 2023)

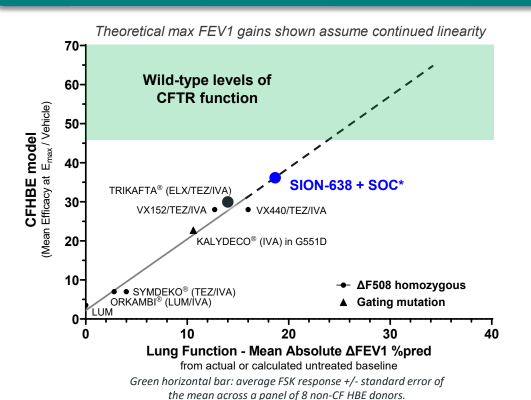
## Sionna has a robust pipeline to drive NBD1 combination strategy with several near-term clinical milestones





# SION-638: First-in-class, clinical stage NBD1 modulator with the potential to deliver higher efficacy

#### SION-638 CFHBE assay data



### Phase 1 human PK supports potential for improved efficacy as an add-on to SOC

- Exposure target for Ph 1 was derived from the CFHBE model\*\* to drive clinically meaningful efficacy
- Dose identified in Ph 1 that achieves target exposure to deliver improved efficacy as add-on to SOC (Trikafta®)
- Progression to Ph 2a will be a portfolio decision informed by Ph 1 data for SION-451 and SION-719

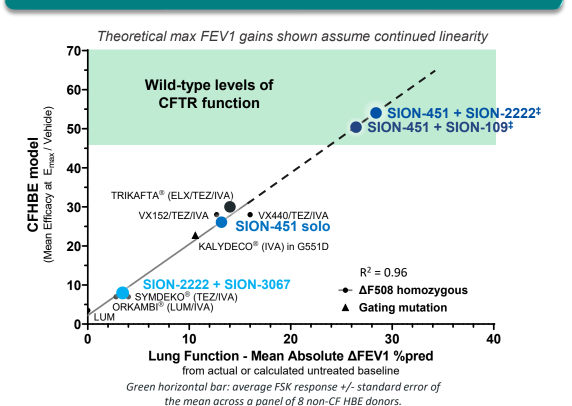


<sup>\*</sup>Based on SION-638 exposure at target dose

<sup>\*\*</sup> Source: Pre-clinical assays conducted by Sionna Trikafta®, Symdeko®, and Orkambi® are registered trademarks of Vertex Pharmaceuticals

### SION-451: Phase 1 ready NBD1 stabilizer demonstrates potential to normalize CFTR function as a dual combo

#### Potential of SION-451 at E<sub>max</sub>



#### Multiple options to raise the efficacy bar

### In the clinically predictive CFHBE assay\*, SION-451 has demonstrated the potential for:

- Single-agent efficacy equivalent to Trikafta<sup>®</sup> at high SION-451 exposures
- Wild-type levels of CFTR function in double combination with a Sionna complementary CFTR modulator
- Wild-type levels of CFTR function as add-on to Trikafta<sup>®\*+</sup>



<sup>\*</sup> Source: Pre-clinical assays conducted by Sionna; + Data not shown

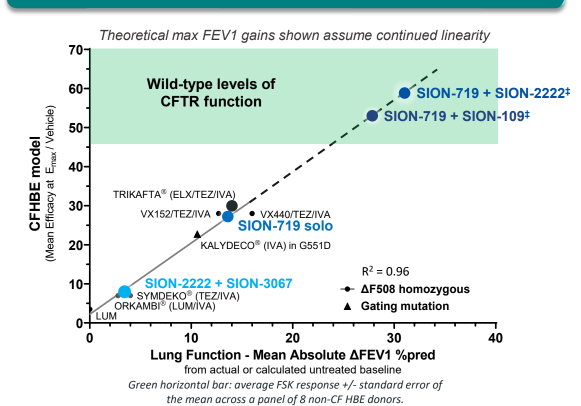
CFHBE - CF Human Bronchial Epithelial primary cells, DC - Development Candidate, FEV- Forced Expiratory

Volume, Gating mutation - G551D CFTR, ELX - elexacaftor, IVA - ivacaftor, LUM - lumacaftor, SOC - Standard

of Care, TEZ - tezacaftor, Trikafta - ELX/TEZ/IVA.

# SION-719: Phase 1 ready NBD1 stabilizer provides another strong option to increase the CF efficacy bar

#### Potential of SION-719 at E<sub>max</sub>



#### Multiple options to raise the efficacy bar

### In the clinically predictive CFHBE assay\*, SION-719 has demonstrated the potential for:

- Single-agent efficacy equivalent to Trikafta® at high SION-719 exposures
- Wild-type levels of CFTR function in double combination with a Sionna complementary CFTR modulator
- Wild-type levels of CFTR function as add-on to Trikafta<sup>®\*+</sup>



<sup>\*</sup> Source: Pre-clinical assays conducted by Sionna; + Data not shown

CFHBF - CF Human Bronchial Epithelial primary cells. DC - Development Candidate. FEV- Forced F

CFHBE - CF Human Bronchial Epithelial primary cells, DC - Development Candidate, FEV- Forced Expiratory Volume, Gating mutation - G551D CFTR, ELX - elexacaftor, IVA - ivacaftor, LUM - lumacaftor, SOC - Standard of Care, TEZ - tezacaftor, Trikafta - ELX/TEZ/IVA.

# Lead Complementary Programs: Galicaftor (SION-2222) & SION-109



# TMD1 directed corrector galicaftor (SION-2222) is an attractive combination agent with NBD1 stabilizers

Mechanism of Action	TMD1-directed CFTR corrector
Rationale and Enthusiasm for Advancement	<ul> <li>Galicaftor (SION-2222) synergizes with NBD1-directed correctors in CFHBE assay</li> <li>Ph 2 demonstrates sweat chloride and ppFEV<sub>1</sub> outcomes in combination with navocaftor (SION-3067, potentiator) comparable to approved duals (Symdeko<sup>®</sup> and Orkambi<sup>®</sup>)</li> <li>API acquired to supply late-stage development</li> </ul>
Status	Phase 2 studies completed by AbbVie*
Key Upcoming Milestones	Combo MAD initiation with NBD1 stabilizer 2H25
Preferred Use Case & TTP	Part of a Sionna proprietary double



# In Ph 1/Ph 2 studies, galicaftor (SION-2222) was well-tolerated in healthy volunteers & CF patients, showed improvement in pulmonary function

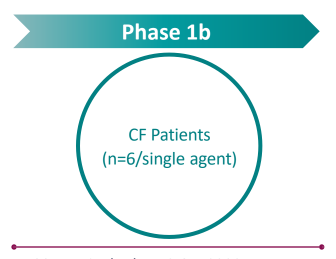
# Healthy Volunteers (n=82/single agent, n=143/combination)

Well-tolerated at all doses as single agent or combination with other modulators

- Up to 600 mg QD, 14d as single agent
- Up to 300 mg QD, 14d as combination

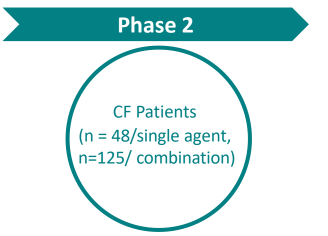
No significant PK DDI with any other modulators tested in combination

t<sub>1/2</sub> ~ **12hr** 



100 mg single dose SION-2222 or PBO

**Well-tolerated**, PK similar to healthy volunteers



50-400 mg QD SION-2222 vs PBO, or 10-300 mg QD SION-2222/150 mg QD SION-3067 vs PBO in F/F CF patients for 4 weeks

200 mg QD **SION-2222** significantly decreased **SwCl** as single agent therapy in F/F patients

200 mg QD SION-2222 significantly improved pulmonary function, and decreased SwCl as dual combination with SION-3067



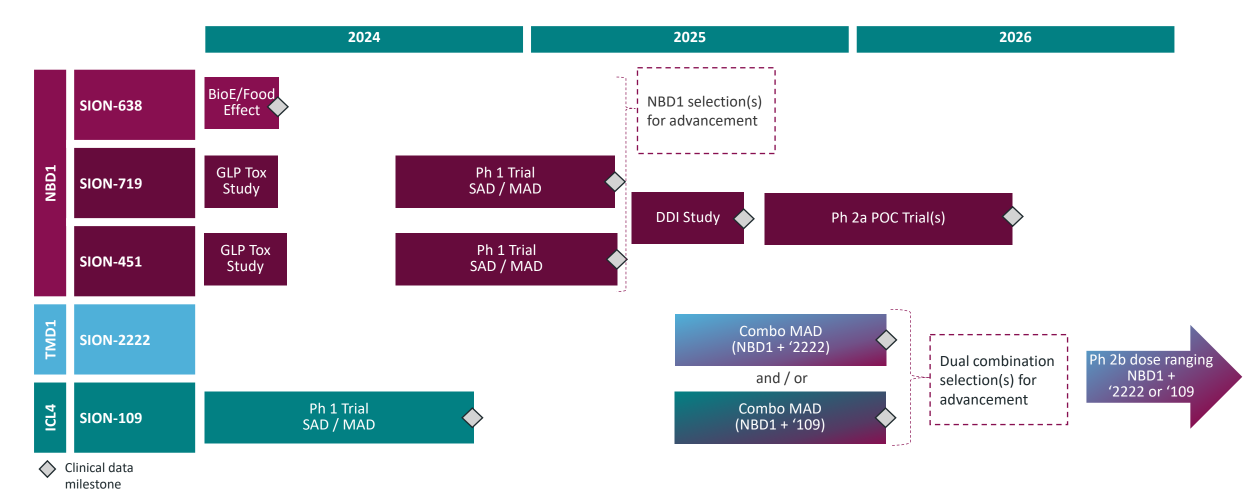
### SION-109, ICL4-directed modulator in Phase 1

Mechanism of Action	ICL4-directed CFTR corrector
Rationale and Enthusiasm for Advancement	<ul> <li>SION-109 synergizes with NBD1-directed stabilizers</li> <li>Promising potency and drug-like profile and tractable predicted target clinical dose</li> <li>No adverse findings in 28-day GLP rat and dog tox, robust margins to target clinical exposures</li> <li>API manufacture completed to support early clinical development</li> </ul>
Status	Phase 1 study initiated in 1Q24
Key Upcoming Milestones	Completion of Phase 1 in 2H24
Preferred Use Case & TTP	Part of a Sionna proprietary double combination





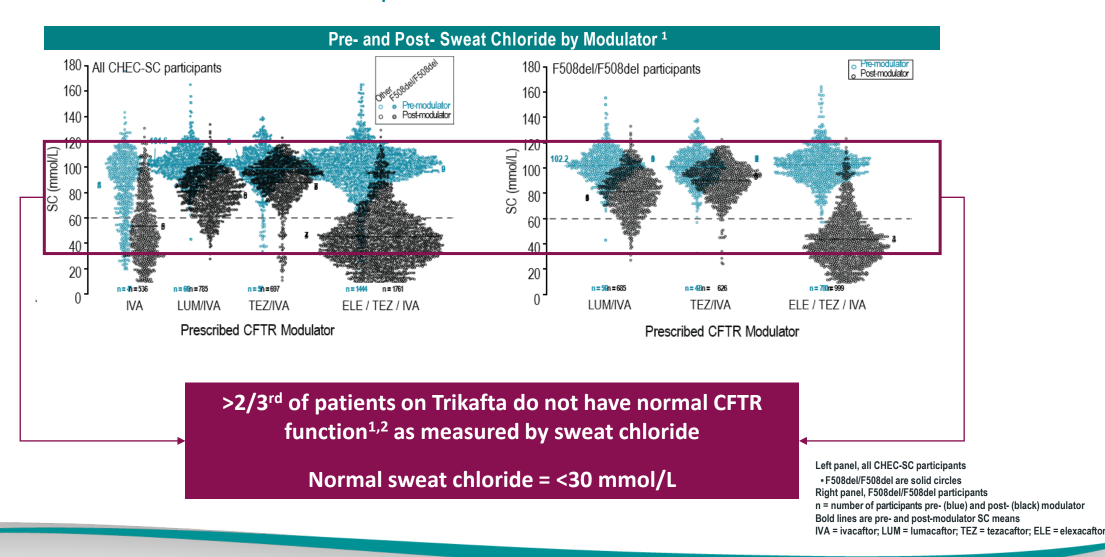
# Sionna's dual combination path will be data driven, selecting the best NBD1 and complementary compounds from our deep pipeline







# The efficacy unmet need remains high, as the goal is to achieve normal CFTR function for CF patients





# Commitment to advancing game-changing therapies, building significant near-term value, and raising the efficacy bar in CF



- SION-638 Phase 1 completed in 1H24
- SION-719 Phase 1 initiated 3Q24
- SION-451 Phase 1 initiated 3Q24



- Portfolio includes multiple clinical stage combo shots on goal:
  - Galicaftor (SION-2222), a TMD1 modulator with Ph 2 data in CF patients
  - SION-109 (ICL4 modulator), Phase 1 data expected 2H24
  - SION-2851 and SION-3067 create options for lifecycle development opportunities
- COMPANY CAPABILITIES
- Goal is to execute the strategy, grow the company, and build capabilities to become a leader in CF
- Cash runway through 2026 and Ph 2a POC

