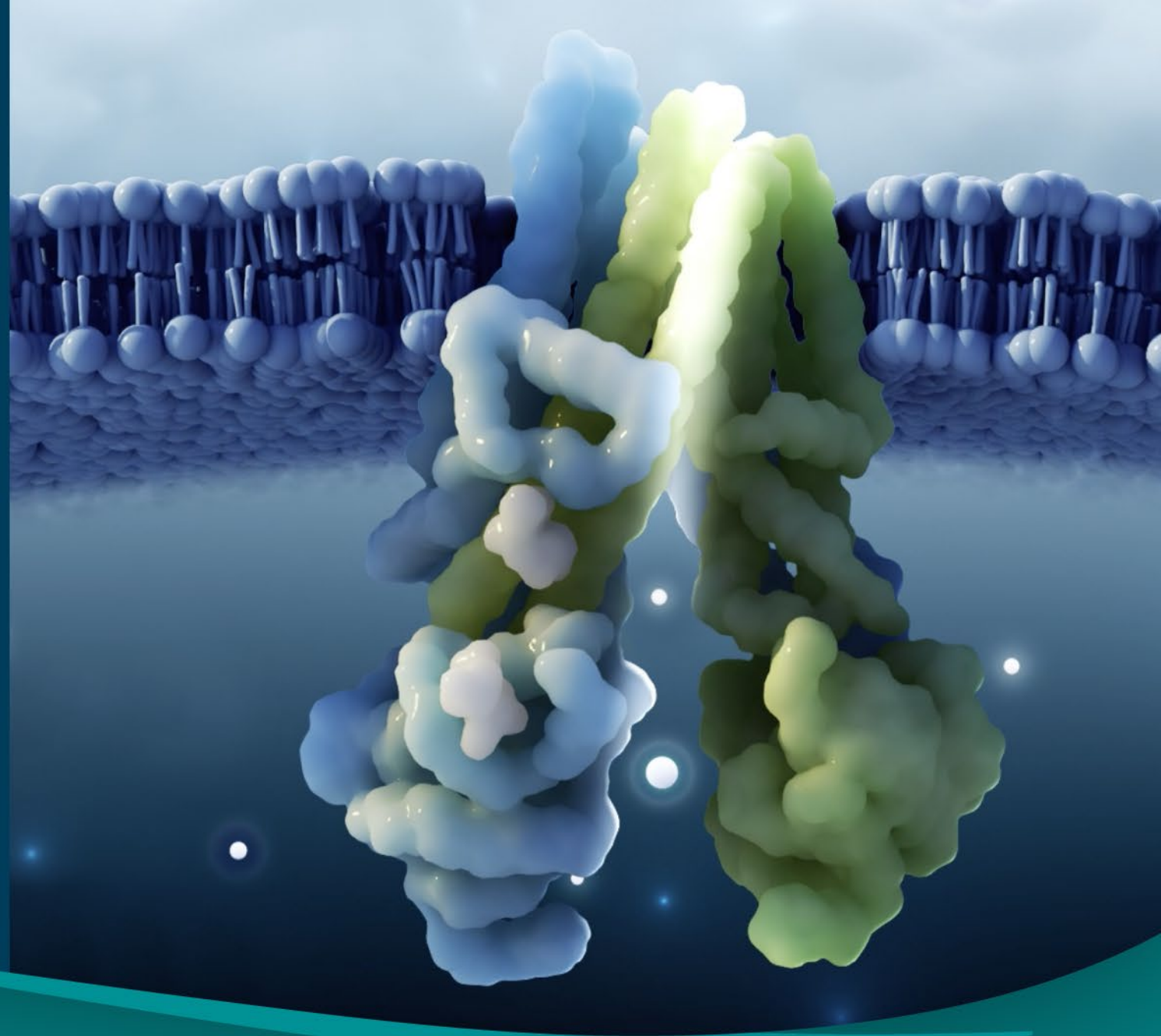


Sionna Therapeutics

August 2024



sionna[™]

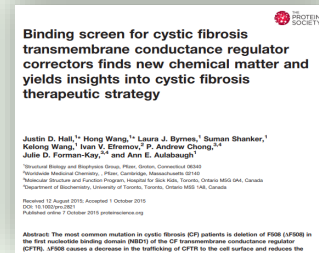
Sionna's differentiated approach focused on NBD1 has a clear path to POC with the potential to deliver best-in-class efficacy

HIGH UNMET NEED IN LARGE MARKET



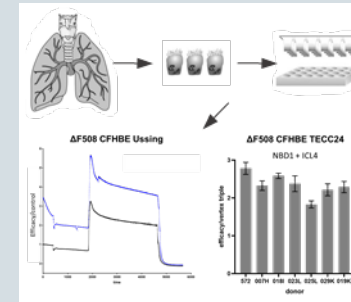
Despite current treatments, unmet need is high in the >\$10B market

NBD1, THE HOLY GRAIL FOR CFTR



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

PREDICTIVE ASSAYS/BIOMARKERS



CFHBE assay and sweat 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



Charlotte McKee, MD
Chief Medical Officer



Elena Ridloff
Chief Financial Officer



Jen Fitzpatrick
General Counsel



Vanya Sagar
Chief People Officer



Greg Hurlbut, PhD
Co-Founder
SVP, Discovery Research



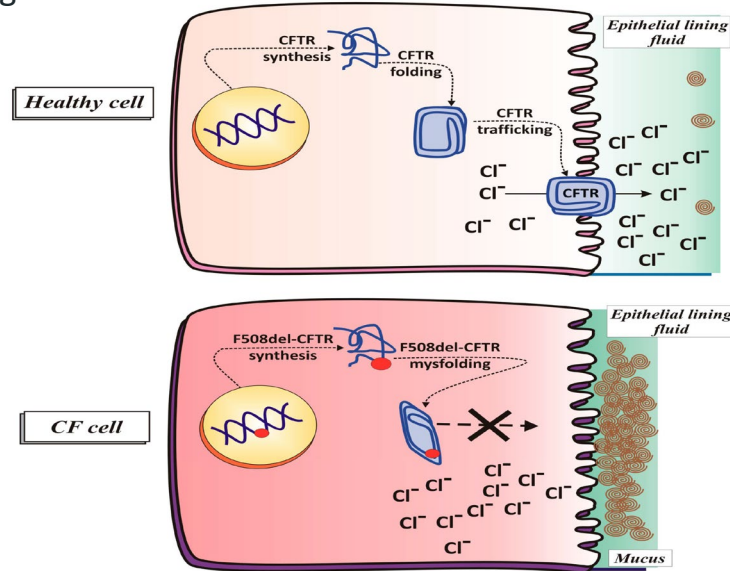
Mark Munson, PhD
Co-Founder
VP, Medicinal Chemistry



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

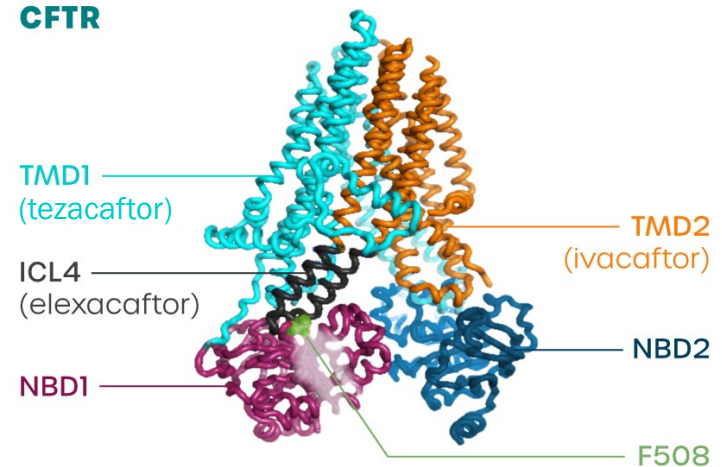


The Importance of NBD1

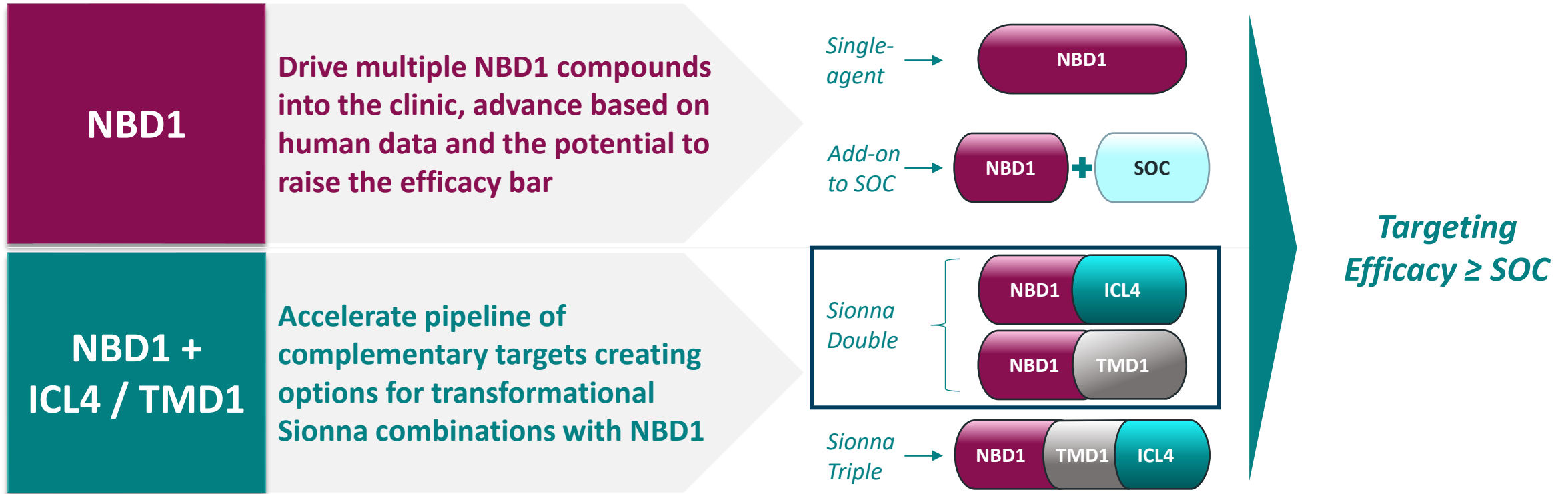
- F508 is present within CFTR's NBD1 domain
- F508del causes NBD1 to unfold at body temperature and weakens NBD1's interface with other regions; these defects cripple CFTR folding, trafficking and function
- **None** of the existing correctors or potentiators address both $\Delta F508$ -CFTR's assembly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

NBD1 is the key to full CFTR correction

CFTR



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

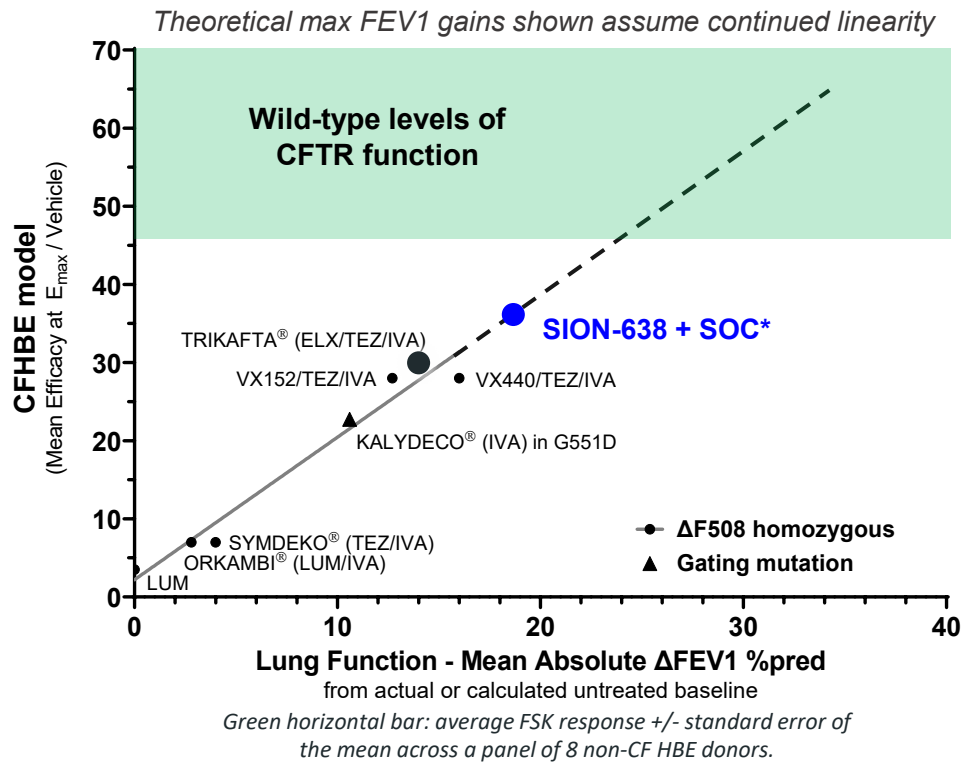
Galicaftor (ABBV-2222), a TMD1 modulator, has positive Ph 2 data in CF patients¹
Plan to advance ABBV-2222 and SION-109, if Ph 1 is successful, as potential dual combination options with an NBD1 stabilizer

LCM Options with Additional Clinical Assets

Navocafter (ABBV-3067), a potentiator, also has positive Ph 2 data; ABBV-2851 is a Ph 1 TMD1 modulator
Both modulators provide lifecycle development opportunities

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

SION-638 CFHBE assay data

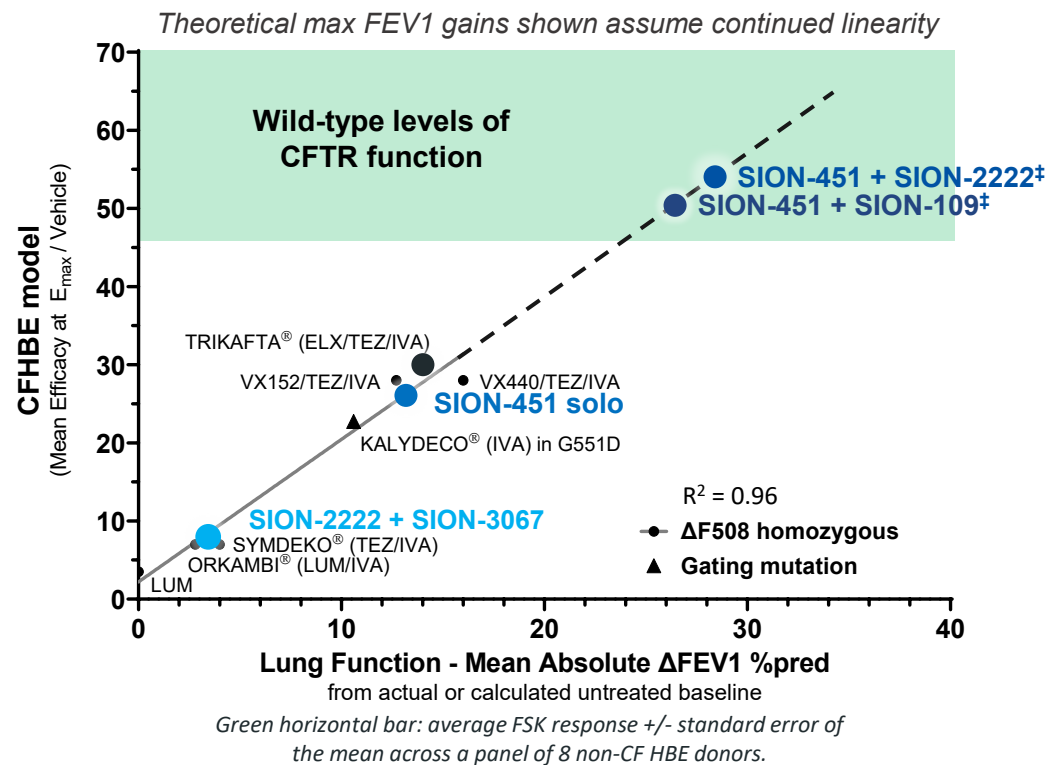


Phase 1 human PK supports potential for improved efficacy as an add-on to SOC

- 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

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

Potential of SION-451 at E_{max}



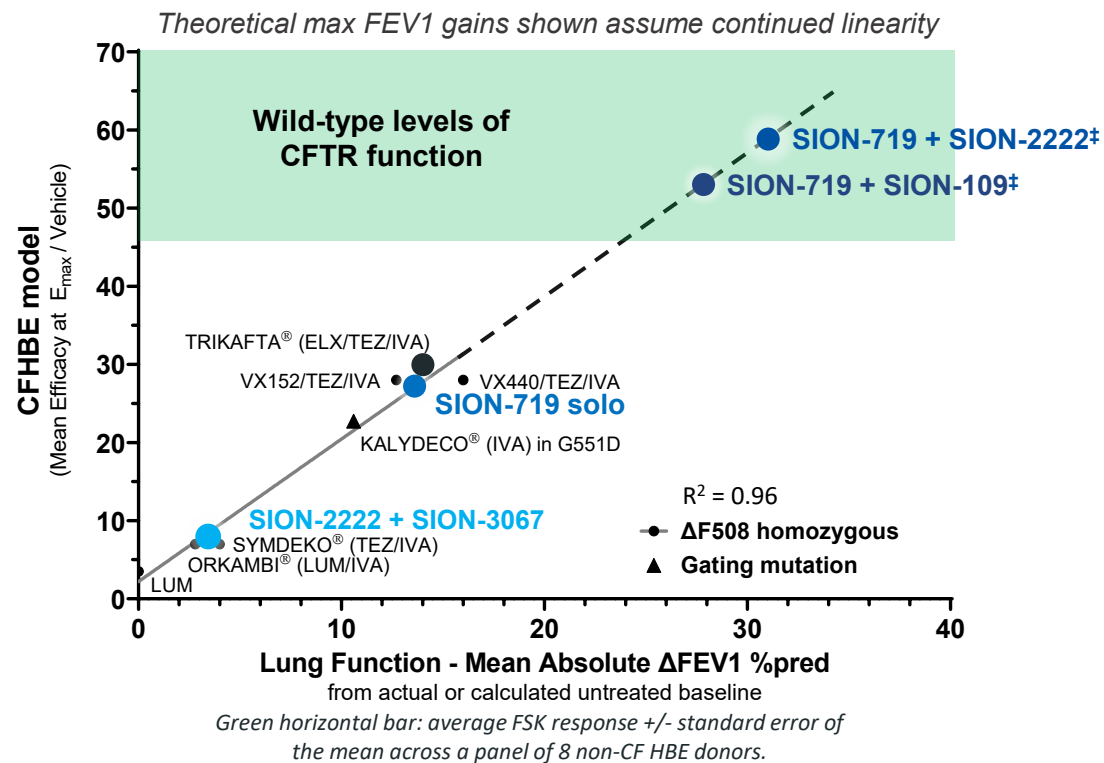
Multiple options to raise the efficacy bar

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

- Single-agent efficacy equivalent to Trikafta® 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®*+

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

Potential of SION-719 at E_{max}



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®*+

*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	<ul style="list-style-type: none">• TMD1-directed CFTR corrector
Rationale and Enthusiasm for Advancement	<ul style="list-style-type: none">• Galicaftor (SION-2222) synergizes with NBD1-directed correctors in CFHBE assay• Ph 2 demonstrates sweat chloride and ppFEV₁ outcomes in combination with navocafator (SION-3067, potentiator) comparable to approved duals (Symdeko[®] and Orkambi[®])• API acquired to supply late-stage development
Status	<ul style="list-style-type: none">• Phase 2 studies completed by AbbVie*
Key Upcoming Milestones	<ul style="list-style-type: none">• Combo MAD initiation with NBD1 stabilizer 2H25
Preferred Use Case & TTP	<ul style="list-style-type: none">• Part of a Sionna proprietary double

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

Phase 1

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

$t_{1/2}$ ~ **12hr**

Phase 1b

CF Patients
(n=6/single agent)

100 mg single dose SION-2222 or PBO

Well-tolerated, PK similar to healthy volunteers

Phase 2

CF Patients
(n = 48/single agent,
n=125/ combination)

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

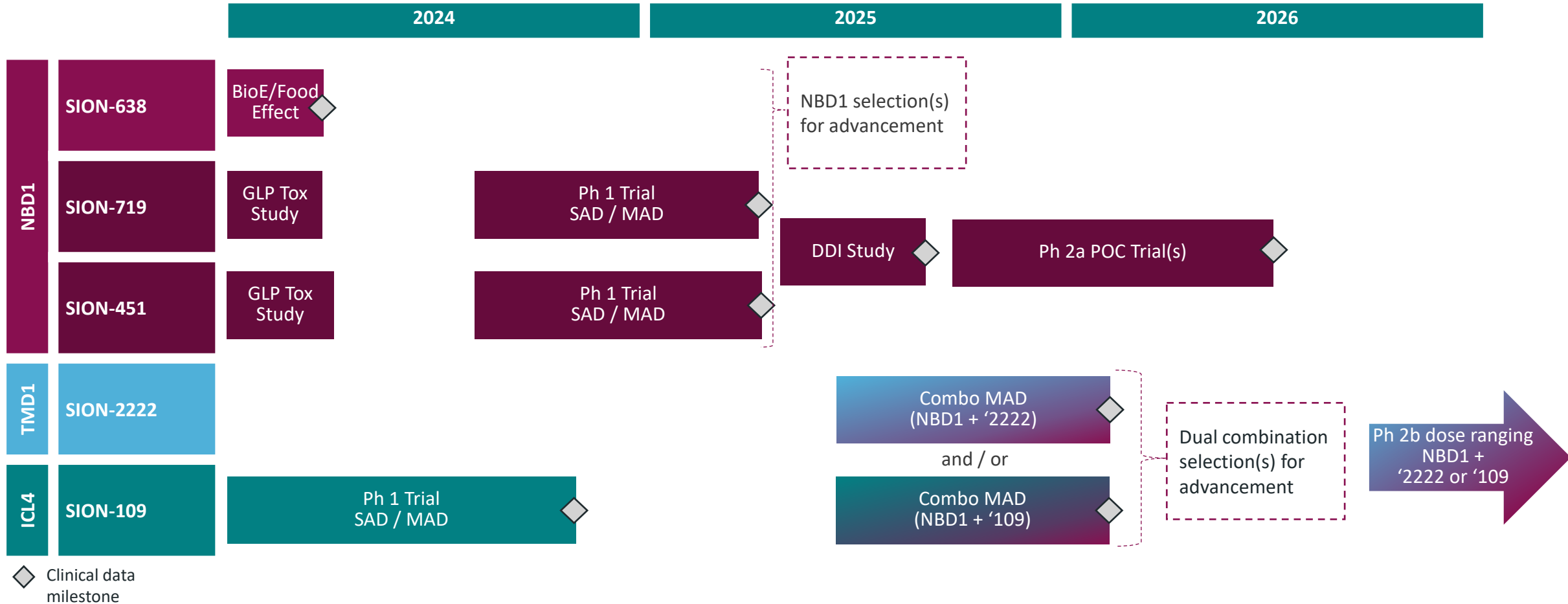
200 mg QD **SION-2222 significantly improved pulmonary function, and decreased SwCl** as dual combination with SION-3067

SION-109, ICL4-directed modulator in Phase 1

Mechanism of Action	<ul style="list-style-type: none">• ICL4-directed CFTR corrector
Rationale and Enthusiasm for Advancement	<ul style="list-style-type: none">• SION-109 synergizes with NBD1-directed stabilizers<ul style="list-style-type: none">• Promising potency and drug-like profile and tractable predicted target clinical dose• No adverse findings in 28-day GLP rat and dog tox, robust margins to target clinical exposures• API manufacture completed to support early clinical development
Status	<ul style="list-style-type: none">• Phase 1 study initiated in 1Q24
Key Upcoming Milestones	<ul style="list-style-type: none">• Completion of Phase 1 in 2H24
Preferred Use Case & TTP	<ul style="list-style-type: none">• Part of a Sionna proprietary double combination

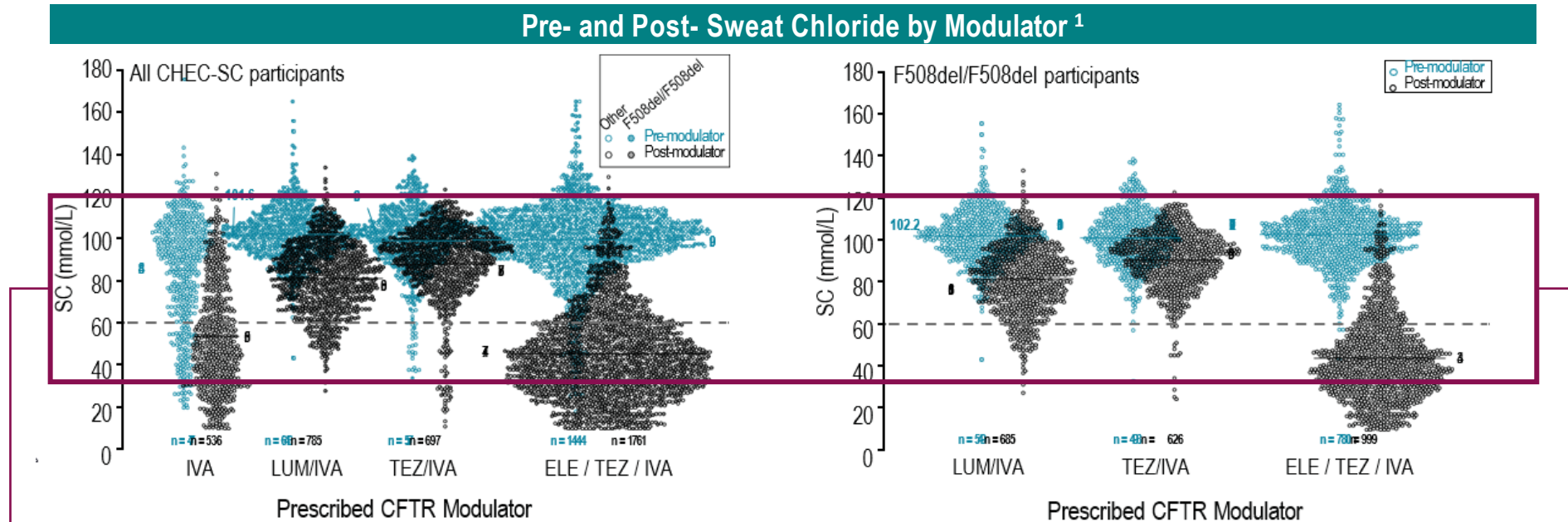
Clinical and Portfolio Strategy

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



Unmet Need

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



>2/3rd of patients on Trikafta do not have normal CFTR function^{1,2} as measured by sweat chloride

Normal sweat chloride = <30 mmol/L

Left panel, all CHEC-SC participants
 - F508del/F508del are solid circles
 Right panel, F508del/F508del participants
 n = number of participants pre- (blue) and post- (black) modulator
 Bold lines are pre- and post-modulator SC means
 IVA = ivacaftor; LUM = lumacaftor; TEZ = tezacaftor; ELE = elexacaftor

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

DELIVER



ROBUST NBD1 PORTFOLIO

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

ADVANCE



COMPLEMENTARY TARGETS

- Portfolio includes multiple clinical stage combo shots on goal:
 - Galicftor (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

BUILD



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