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ENaC inhibition in cystic fibrosis: potential role in the new era of CFTR modulator therapies

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Take-home message

ENaC inhibition with BI 1265162 is a promising strategy to optimise outcomes in patients with CF either eligible, or ineligible, for CFTR modulator therapy. Phase II clinical trials of BI 1265162 must now show this translates into clinical benefit.

Abstract

Cystic fibrosis transmembrane conductance regulator (CFTR) modulators are the first approved drugs targeting underlying epithelial ion/fluid transport defects in patients with cystic fibrosis (CF). Current CFTR modulators restore mutant CFTR activity to up to ~50% of normal CFTR Cl⁻ channel function, translating into improvements in percentage predicted FEV₁ and other clinical outcomes. In addition, reductions in airway bacterial colonisation are observed; however, patients fail to eradicate bacteria over time and still experience pulmonary exacerbations, and long-term safety of CFTR modulator therapy remains unknown. Currently approved CFTR modulators are predicted to be effective for up to 90% of patients. A mutation-agnostic approach could address the remaining 10% with CFTR mutations unresponsive to CFTR modulator therapy and may act together with CFTR modulator therapy to further improve epithelial ion/fluid transport and clinical outcomes. Together with CFTR and other Cl⁻ channels, the epithelial Na⁺ channel (ENaC) is key to regulating airway surface liquid homeostasis. ENaC activity is limiting for Na⁺/fluid absorption and remains intact or may even be increased in CF airways, leading to increased Na⁺/fluid absorption, airway surface dehydration, impaired mucociliary clearance, bacterial infection, inflammation and progressive lung damage – the major cause of CF-related morbidity and mortality. Inhibition of ENaC in the airways is therefore an attractive therapeutic target to counteract airway surface dehydration and downstream consequences in CF lung disease. This review examines ENaC inhibition in CF therapy, and describes a new ENaC inhibitor with potential mutation-agnostic therapeutic benefit, both alone, and in synergy with CFTR modulators.

Introduction

Small-molecule CFTR modulator drugs for cystic fibrosis (CF) are the first therapies since the disease was initially described by Guido Fanconi in 1936 [1] to target and partially restore the function of the cystic fibrosis transmembrane conductance regulator (CFTR) Cl⁻ channel. CFTR modulator therapy is expected to have significant clinical benefits for many, but it does not result in a cure and is not appropriate or available for all patients with CF [2, 3]. In this review, evidence is described suggesting that inhibiting the epithelial Na⁺ channel (ENaC) responsible for the Na⁺/fluid absorption that contributes to airway surface dehydration and impaired mucociliary clearance (MCC) observed in CF airways may significantly improve clinical outcomes irrespective of the CFTR genotype, and may synergise with currently approved CFTR modulators to further improve clinical outcomes.
The CFTR modulator landscape

Significant progress has been made in the treatment of patients with CF with the introduction of CFTR modulator therapies, which consist of CFTR correctors that improve folding and trafficking of the common F508del-CFTR mutation and potentiators that improve the open probability of mutant CFTR channels at the apical cell membrane [4-6]. Current CFTR modulator drugs vary in efficacy in improving CFTR function and clinical outcomes. The potentiator ivacaftor was the first approved CFTR modulator for CF patients with at least one G551D-CFTR gating mutation and was shown to rescue mutant CFTR function to ~50% of wild-type levels, which was associated with an improvement in mean absolute percentage predicted forced expiratory volume in 1 second (ppFEV₁) of ~11% in the pivotal trial [3, 5, 7]. Subsequently developed corrector-potentiator combinations for patients homozygous for the common F508del-CFTR allele (lumacaftor/ivacaftor and tezacaftor/ivacaftor) or patients with one F508del allele and a residual function allele (tezacaftor/ivacaftor) showed smaller improvements in CFTR function (to ~10–20% of wild-type levels), which were associated with more modest improvements of ppFEV₁ of ~3–4% in F508del homozygous patients [6, 8, 9] and ~7% in patients with one F508del allele in combination with a residual function allele [10]. In addition, it has been shown that bacterial counts and inflammatory markers are reduced in sputum of patients treated with ivacaftor, but that bacteria are not eradicated over time [11, 12]. A triple-agent CFTR modulator drug (elexacaftor/tezacaftor/ivacaftor) has recently been approved in patients with a single F508del-CFTR allele. Elexacaftor, like tezacaftor, is a CFTR corrector, but acts additively at a second site in the F508del-CFTR protein to improve multiple folding defects [13]. Consistent with this additive effect at the molecular level, this triple-combination CFTR modulator therapy resulted in greater improvement in lung function and other clinical outcomes (an approximately 11–14% improvement in ppFEV₁ compared to control) in patients homozygous for F508del or compound heterozygous with a minimal function allele [14, 15] and is predicted to become the ‘gold standard’ for the treatment of up to 90% of patients with CF (those with at least one F508del allele) [2]. However, for most patients with chronic CF lung disease, this improvement in pulmonary function is not necessarily a return to the normal range, and patients continued to have exacerbations, albeit at much reduced rates [14]. Improving clinical outcomes further in patients receiving elexacaftor/tezacaftor/ivacaftor has the potential to have a significant impact on the CF population (Fig. 1). In addition, the approximately 10% of CF patients for which CFTR modulator therapy is ineffective (those who do not carry at least one F508del allele) could benefit from a mutation-agnostic approach. Finally, while long-term safety of treatment with elexacaftor/tezacaftor/ivacaftor remains unknown, other therapeutic avenues should be considered.

ENaC: role in healthy and CF airways, and potential therapeutic target
In the conducting airways, ENaC is expressed at the apical membrane of airway epithelial cells and provides the limiting pathway for transepithelial Na\(^+\) absorption that drives absorption of Cl\(^-\) and water through the paracellular shunt pathway (Fig. 2) [16-18]. In healthy airways, coordinated Cl\(^-\) secretion by CFTR and other Cl\(^-\) channels and Na\(^+\) absorption by ENaC is essential for proper volume regulation of airway surface liquid (ASL), which comprises the periciliary layer (PCL) and the overlying mucus layer [19, 20]. ENaC is regulated by several intracellular and extracellular mechanisms. Intracellular mechanisms include activation by convertase-type proteases such as furin [21], and inhibition by CFTR [22, 23]. Extrinsic activation mechanisms include proteolytic cleavage by neutrophil elastase and other proteases released from neutrophils and other inflammatory cells in the airways [24, 25], as well as by bacterial proteases released in airway infection [26, 27]. Therefore, ongoing inflammation and infection in CF patients with established lung disease, including those treated with CFTR modulators [11, 12], is likely to cause ongoing proteolytic cleavage and activation of ENaC in the airways.

Evidence suggests that ENaC is hyperactivated in CF as a result of CFTR dysfunction and/or proteolytic activation by host- and bacteria-derived proteases [22-29]. Hyperactivated ENaC in the airways of CF patients results in markedly increased Na\(^+\) absorption [20, 30, 31], leading to increased paracellular Cl\(^-\) and water absorption from the airway lumen, ASL volume depletion, hyperconcentration of mucus, reduced height of the PCL with compressed cilia, reduced mucociliary clearance (MCC), mucus accumulation, airway plugging, bacterial colonisation, inflammation, progressive tissue damage and decline in lung function [25, 32]. The pathogenic role of increased ENaC function has been supported by the lung phenotype of mice with airway-specific overexpression of the β-subunit of ENaC (βENaC-Tg) that phenocopy CF-like airway surface dehydration/mucus hyperconcentration and develop CF-like lung disease [33, 34]. Collectively, these results support ENaC inhibition in the airways as an attractive target for CF therapy. In this context, it is noteworthy that patients with pseudohypoaldosteronism with loss-of-function mutations in the α- and β-subunits of ENaC have increased ASL volume and MCC rates [35], and CF patients with a mutation in the δ-subunit of ENaC causing reduced ENaC activity were found to have slow progression of lung disease [36].

Inhibition of ENaC was first demonstrated with amiloride, and in the kidney, this is an effective diuretic drug [37, 38]. In βENaC-Tg mice, which display a CF-like lung phenotype, intrapulmonary treatment with amiloride was effective, leading to a reduction in mucus plugging, airway inflammation and pulmonary mortality when started as preventive therapy immediately after birth, i.e. when the lungs were structurally normal; however, no beneficial effects were observed in adult βENaC-Tg mice with established CF-like lung disease [39]. In CF patients, probably due to amiloride’s
short half-life and limited potency, studies of the effect of inhaled amiloride on MCC and lung function resulted in only moderate and inconsistent improvements, even with multiple doses [40-42]. The amiloride derivative GS-9411, whilst 100 times more potent than amiloride and with a longer duration of action, failed Phase I trials as an inhaled ENaC blocker due to ENaC inhibition in the kidney, resulting in hyperkalaemia [43], which may have cardiac and neurological safety implications. These studies led to the hypothesis that if the poor pharmacodynamics and safety observed with amiloride and GS-9411 could be overcome with a newer-generation compound, ENaC inhibition would be a viable therapy to improve airway surface hydration and pulmonary outcomes in patients with CF, irrespective of CFTR genotype (i.e. a mutation-agnostic therapy), particularly in those with rare CFTR mutations that have no currently approved CFTR modulator therapy [42, 44-46]. Further, more recently, mucus hyperconcentration has also been implicated in the pathogenesis of a spectrum of other muco-obstructive lung diseases including chronic bronchitis and non-CF bronchiectasis, suggesting that ENaC inhibition may be beneficial far beyond CF [47-49]. Importantly, ENaC inhibition may act synergistically with CFTR modulators (Fig. 2e). CFTR can secrete or absorb Cl\(^-\) across epithelial surfaces depending on the electrochemical driving force that is determined by i) the intra- and extracellular Cl\(^-\) concentrations that are tightly regulated and result in a reversal potential for Cl\(^-\) (i.e. the membrane potential at which there is no net flow of Cl\(^-\) from one side of the membrane to the other, also known as the Nernst potential) in the range of −30 mV in CF airway epithelial cells, and ii) the membrane potential of the cell that is set by the relative conductances of Cl\(^-\), Na\(^+\) and K\(^+\) channels and respective intra- and extracellular concentrations of these ions [17, 18, 50]. It is therefore predicted that if ENaC is inactive or not expressed in the same cell, cAMP-mediated stimulation will lead to a concomitant activation of apical CFTR and basolateral K\(^+\) channels (reversal potential for K\(^+\) approximately −90 mV) that drives the membrane potential more negative than the reversal potential of Cl\(^-\), and thus generate a driving force for CFTR-mediated Cl\(^-\) secretion; however, if ENaC is active in the same cell (reversal potential for Na\(^+\) approximately +60 mV) this will depolarise the membrane potential, reduce the driving force for Cl\(^-\) secretion, and may even result in CFTR-mediated Cl\(^-\) absorption as observed in the sweat duct [17, 18, 28, 50-52]. Regarding the combined use of CFTR modulators and ENaC inhibitors in patients with CF, this implies that inhibition of hyperactive ENaC in the apical membrane of airway epithelial cells may not only block ENaC-mediated Na\(^+\)/fluid absorption, but will also hyperpolarise the apical cell membrane and thus increase the driving force for Cl\(^-\)/fluid secretion via mutant CFTR channels that have been rescued and inserted into the apical plasma membrane by CFTR modulators. Whether this is reflected in synergistic effects on airway surface hydration, MCC and pulmonary outcomes in patients with CF and other muco-obstructive lung diseases needs to be assessed in clinical trials.
Clinical development of new ENaC inhibitors

Several new compounds designed for ENaC inhibition are currently in active preclinical development [44, 45, 53-57] (Table 1). These new compounds employ different modes of action ranging from highly potent and durable inhibition of the ENaC channel pore, inhibition of ENaC-activating proteases, to inhibition of ENaC expression by antisense oligonucleotides or small interfering RNA. Of these, the new small-molecule ENaC inhibitor BI 1265162 is the only compound currently in Phase II development [58]. It has demonstrated efficacy in preclinical investigations [59-63], with a markedly higher potency than amiloride (a 30–70-fold lower half maximal inhibitory concentration) [63], and no effects on serum K⁺ and plasma electrolytes have been observed [60, 62]. In addition, BI 1265162 has shown safety in Phase I volunteer studies [64]. Importantly, preclinical work supports the potential mutation-agnostic property of BI 1265162 [63]. Using highly differentiated human airway epithelial cell cultures grown at an air–liquid interface, it was found that transepithelial fluid absorption from the apical surface to the basolateral compartment was reduced by BI 1265162, with or without CFTR modulators, in both CF and normal airway cultures [63]. This work also supports the hypothesis of a potential synergistic effect with CFTR modulators. Addition of lumacaftor/ivacaftor alone to F508del/F508del CF cultures partially restored mucus transport velocity to approximately 50% of that observed in normal airway cultures; however, addition of BI 1265162 to lumacaftor/ivacaftor further improved mucus transport velocity in CF cultures to a level similar to normal airway cultures [63]. Taken together, the above evidence suggests that BI 1265162 is a promising candidate as a monotherapy for CF patients for whom CFTR modulators are ineffective and in combination with CFTR modulators providing synergistic effects. However, no ENaC inhibitor development, so far, has translated into clinical success [44] and several clinical development programmes of ENaC inhibitors for patients with CF (VX-371; QBW276; Novartis and SPX-101; Spyryx [an indirect inhibitor]) have recently been terminated (Table 1). The clinical development of VX-371 for primary ciliary dyskinesia (NCT02871778) has also recently been terminated (clinicaltrials.gov NCT02871778). Failure of ENaC inhibitors to progress in clinical development may be due to inadequate dosing and/or deposition by inhalation in CF patients with chronic lung disease characterised by heterogeneous airway mucus plugging. These ENaC inhibitors demonstrated improvement of MCC in the sheep model, but lack of translation of this preclinical model to patients with CF may be related to the fact that the sheep do not have airway mucus plugging and structural lung damage commonly present in patients with CF. Therefore, the dose of inhaled ENaC inhibitor that improves MCC in the sheep model may not be sufficient to achieve therapeutically active ENaC inhibition in CF patients. To address this issue in more detail, more recent CF animal models such as the CF pig and CF ferret, which feature CF-like lung disease
including heterogeneous mucus plugging, chronic airway infection, inflammation and bronchiectasis, may be utilised to better understand issues related to target engagement and the potential role of ENaC inhibitors as a therapeutic option in CF lung disease [65, 66]. Whether the hurdle of sufficient delivery of inhaled BI 1265162 to mucus-obstructed airways can be overcome in patients with CF remains to be seen in clinical trials. In addition, systemic side effects such as hyperkalaemia, short study duration, non-study-related exacerbations and lack of sensitivity of traditional endpoints such as ppFEV₁ to detect treatment benefits may impede the clinical development of an inhaled ENaC inhibitor therapy.

How could these hurdles be overcome in the clinical development of a new ENaC inhibitor therapy for CF? First, a successful ENaC inhibitor will have to be highly potent, delivered to the airways in a sufficient dose and have a long duration of action to provide effective ENaC inhibition and maximal treatment effect. Second, the new ENaC blocker should result in minimal off-target effects and systemic exposure. In this context, the following translational challenges that have ended the clinical programmes of earlier ENaC inhibitors have been addressed in order to minimise BI 1265162’s risk of failure in Phase II studies. The risk of underdosing was minimised by basing the dose for the Phase II trial in CF patients on fluid absorption data in the rat model (in which BI 1265162 was instilled into the trachea) [60] in addition to MCC data in the sheep model (in which BI 1265162 was nebulised), factoring in the expected lung deposition using the Respimat device in humans [67]. Of note, the improvements in MCC observed with BI 1265162 in the sheep model [62] were comparable to the improvement obtained in CF patients with the G551D gating mutation after starting CFTR modulator therapy with ivacaftor [68]. These data suggest that inhaled BI 1265162, if delivered successfully to the CF lung, has the potential to provide similar benefits on MCC in CF patients with genotypes that do not respond to CFTR modulators. In the subsequent Phase I studies [64], BI 1265162 did not result in drug-related hyperkalaemia and was well tolerated when administered as single or multiple doses up to 1200 µg daily. Moving into Phase II, lung clearance index (LCI) derived from multiple breath washout has been included as an additional endpoint [58]; LCI reflects ventilation inhomogeneity in the lung [69], correlates with mucus plugging and other morphological changes of the airways [70] and is more sensitive than spirometry (ppFEV₁) to detect functional abnormalities in the small airways typically affected in CF [69]. Crucially, BI 1265162 is an ENaC inhibitor that is considerably more potent than amiloride and has, at least preclinically, been shown to work as a mutation-agnostic monotherapy and in synergy with CFTR modulation [63].

Conclusion
Despite the exciting breakthroughs in the development of CFTR modulator therapy, this therapy is not effective for all patients with CF, as only patients with at least one F508del allele can benefit [2].
Additionally, functional restoration of the underlying ion/fluid transport defect and clinical outcomes are still suboptimal in those patients who are eligible for CFTR modulator therapy. Whereas systemic delivery of CFTR modulators can improve CFTR function in multiple affected organs, potential benefits of inhaled ENaC blockers are likely limited to the lungs. However, ENaC inhibition may become a viable option for patients for whom existing CFTR modulator therapy is ineffective, and it has the potential to act synergistically with CFTR modulation in those patients for whom it is. ENaC inhibition is therefore a promising strategy to optimise therapeutic benefit. Clinical trials with novel, long-acting ENaC inhibitors such as BI 1265162 will now have to demonstrate that ENaC inhibition is safe and translates into clinical benefit in people with CF.
Table 1. Preclinical and clinical development of ENaC inhibitors for cystic fibrosis

<table>
<thead>
<tr>
<th>Drug</th>
<th>Company</th>
<th>Type</th>
<th>MoA</th>
<th>Comments</th>
<th>References</th>
</tr>
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<td><strong>Preclinical</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Benzamil/Phenamil (second-generation amiloride derivatives)</td>
<td>–</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>Discontinued due to rapid clearance, short half-life and epithelial permeability in lungs in sheep</td>
<td>[71]</td>
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<tr>
<td>NVP-QBE 170</td>
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<td>Small molecule</td>
<td>Direct inhibition</td>
<td>–</td>
<td></td>
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<tr>
<td>QUB-TL1</td>
<td>–</td>
<td>Small molecule</td>
<td>Channel-activating protease inhibitor</td>
<td>–</td>
<td>[54]</td>
</tr>
<tr>
<td>MK 104</td>
<td>Mucokinetics</td>
<td>Small molecule</td>
<td>Channel-activating protease inhibitor</td>
<td>–</td>
<td>[55, 56]</td>
</tr>
<tr>
<td>ENaC inhibitory ASO</td>
<td>Ionis</td>
<td>Molecular inhibition</td>
<td>ASO</td>
<td>–</td>
<td>[57]</td>
</tr>
<tr>
<td>ETD001</td>
<td>Enterprise Therapeutics</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>–</td>
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<tr>
<td><strong>Phase I</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>GS-94111 (third-generation amiloride derivative)</td>
<td>Parion</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>Discontinued due to SAE of acute hyperkalaemia</td>
<td>[43]</td>
</tr>
<tr>
<td>BI 443651</td>
<td>Boehringer Ingelheim</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>Discontinued due to palatability issues</td>
<td>clinicaltrials.gov NCT02706925</td>
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<tr>
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<td>AstraZeneca</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>In Phase Ib</td>
<td>clinicaltrials.gov NCT02950805</td>
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<tr>
<td>ARO-ENaC</td>
<td>Arrowhead Pharmaceuticals</td>
<td>Molecular inhibition</td>
<td>Small interfering RNA</td>
<td>In Phase I/IIa</td>
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<td>Amiloride</td>
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<td>Camostat</td>
<td>Novartis</td>
<td>Small molecule</td>
<td>Prostatin inhibitor (channel-activating protease; major regulator of ENaC)</td>
<td>Discontinued due to adverse events/tolerability issues</td>
<td>[72]</td>
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<td>Vertex/Parion</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>Discontinued in combination with ivacaftor/lumacaftor due to lack of efficacy</td>
<td>clinicaltrials.gov NCT02709109</td>
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<tr>
<td>SPX-101</td>
<td>Spyryx</td>
<td>Peptide analogue</td>
<td>SPLUNC-1 analogue, promoting ENaC channel internalisation</td>
<td>Discontinued due to lack of efficacy</td>
<td>clinicaltrials.gov NCT03229252</td>
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<tr>
<td>QBW276</td>
<td>Novartis</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>Development terminated due to strategic reasons</td>
<td>clinicaltrials.gov NCT02566044</td>
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<tr>
<td>BI 1265162</td>
<td>Boehringer Ingelheim</td>
<td>Small molecule</td>
<td>Direct inhibition</td>
<td>–</td>
<td>clinicaltrials.gov NCT04059094</td>
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2 ASO, antisense oligonucleotide; ENaC, epithelial sodium channel; MoA, mode of action; SAE, serious adverse event; SPLUNC-1, short palate, lung, and nasal epithelium clone protein.
Figure 1. The relationship between clinical phenotype and CFTR function, and levels of functional restoration of mutant CFTR by current CFTR modulator therapies in patients with CF.


Figure 2. In healthy airways, a balance between CFTR-mediated secretion and ENaC-mediated absorption of NaCl and H2O facilitates proper hydration of airway surfaces essential for effective mucociliary clearance (a). In CF airways, deficient CFTR-mediated Cl-/fluid secretion and increased ENaC-mediated Na+/fluid absorption leads to airway surface dehydration (reduced PCL and hyperconcentrated mucus), flattened cilia, impaired mucociliary clearance, bacterial colonisation and neutrophilic inflammation (b). Partial restoration of these pathological features by rescue of mutant CFTR function with CFTR modulators (correctors and potentiators) (c) or ENaC inhibition (d). Hypothesised synergy between CFTR modulation and ENaC inhibition, resulting in further improvement in airway surface hydration and reduction in pathological features in CF airways (e). Arrows indicate direction and magnitude of ion and water movement.

-ve, negative regulation; CFTR, cystic fibrosis transmembrane regulator; ENaC, epithelial sodium channel; PCL, periciliary layer.
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Healthy individuals

100%

Obligate heterozygotes

Emergence of symptoms (e.g. chronic cough)

CFTR-related disease
Including acute recurrent pancreatitis, CBAVD, and sinusopulmonary disease

Cystic fibrosis
Pancreatic insufficiency

Pancreatic insufficiency

Clinical phenotype

Ultimate goal to achieve optimal benefit

Ivacaftor (G551D/other) and
Elexacaftor/Tezacaftor/
Ivacaftor (F508del/IVF)

Lumacaftor/Ivacaftor
(F508del/F508del)