CyFi-MAP: an interactive pathway-based resource for cystic fibrosis

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Cystic fibrosis (CF) is a life-threatening autosomal recessive disease caused by more than 2100 mutations in the CF transmembrane conductance regulator (CFTR) gene, generating variability in disease severity among individuals with CF sharing the same CFTR genotype. Systems biology can assist in the collection and visualization of CF data to extract additional biological significance and find novel therapeutic targets. Here, we present the CyFi-MAP—a disease map repository of CFTR molecular mechanisms and pathways involved in CF. Specifically, we represented the wild-type (wt-CFTR) and the F508del associated processes (F508del-CFTR) in separate submaps, with pathways related to protein biosynthesis, endoplasmic reticulum retention, export, activation/inactivation of channel function, and recycling/degradation after endocytosis. CyFi-MAP is an open-access resource with specific, curated and continuously updated information on CFTR-related pathways available online at https://cysticfibrosismap.github.io/. This tool was developed as a reference CF pathway data repository to be continuously updated and used worldwide in CF research.

Omics technologies revolutionized the way researchers generate data1. The integration of disease-specific data allows not only to capture knowledge at its multiple levels of organization but also to unbiasedly identify molecular features, such as phenotypes associated with complex and heterogeneous diseases—which would remain otherwise unnoticed2,3. The visual representation of molecular mechanisms has emerged as a powerful tool for a better understanding and analysis of specific disease-causing features. This led to the development of several pathway-based resources, such as KEGG (Kyoto Encyclopedia of Genes and Genomes)4, Wikipathways5, Reactome6 and MetaCore™7–10.

The need for dedicated knowledge maps as tools for the representation of mechanisms involved in specific diseases in a holistic form gave origin to the concept of a disease map. A disease map consists of a representation of disease mechanisms illustrating the major signalling, metabolic and regulatory pathways known to be involved in a specific disorder in an interchangeable format. The major feature of the disease map is to allow comparisons among different maps using a high-quality representation in a standardised format11. Furthermore, it is computer-readable, user-friendly and can be transformed into mathematical models for predictive analysis and new hypothesis generation11. This resource can store and display multiple layers of information—from subcellular, cellular, tissue, organ to whole organism systems—besides providing customizable levels of detail that can be used in the illustration of the biological mechanisms12. In the past, to further understand their mechanisms and potentially find therapeutic targets, several disease maps for complex disorders have been developed such as Parkinson’s13, Alzheimer’s14, asthma15, several types/forms of cancer16 and rheumatoid arthritis17.

Cystic Fibrosis (CF) is a life-shortening rare genetic disorder affecting 90,000 to 100,000 individuals worldwide, that results from over 2,100 variants in the CF transmembrane conductance regulator (CFTR) gene18–20. The CFTR protein functions as a chloride/bicarbonate channel activated through cAMP-induced phosphorylation at the apical plasma membrane (PM) of epithelial cells. This protein works in a dynamic network, interacting with multiple components and regulating a significant number of other channels21–23. This channel loss of function...
causes the organs in which it is expressed to be impacted. Specifically, in the lung CFTR malfunction causes severe airway dehydration and thickening of the lung mucus, leading to impaired mucociliary clearance (MCC) and, subsequently, clogging of the airways. This causes progressive loss of lung function, which is the major cause of morbidity and mortality.42,5. Furthermore, it is important to note that CFTR loss of function does not occur only in CF, but also in chronic obstructive pulmonary disease (COPD)—the third main cause of death worldwide—revealing the crucial role this protein plays in the airway epithelium.

The most common CF-causing mutation on the CFTR gene is the F508del-CFTR which is a misfolded unstable protein, which is mostly retained in the endoplasmic reticulum (ER) and targeted for ER-associated degradation (ERAD). However, if F508del-CFTR bypasses this pathway and is rescued to the PM, the protein still presents defective channel gating and low PM stability, being rapidly endocytosed and degraded.5,20–22. The biogenesis and intracellular trafficking of F508del-CFTR protein has been extensively studied to better understand the pathways that contribute to and drive CF progression. Notwithstanding, and owing to the inherent complexity of the mechanisms and pathways, the CFTR dynamic network is still only partially understood.

Several databases and software tools have been used in efforts to represent the known CF mechanisms through the representation of pathways covering different processes—such as MetaCore™ from Clarivate Analytics (i.e., CFTR folding and maturation), Reactome (i.e., RHO GTPases regulate CFTR trafficking), KEGG (i.e., ABC-family proteins mediated transport)5,23–26. Notwithstanding, these pathway-based databases present CFTR data incorporated with non-related processes and proteins such as other ABC-family proteins, which although allow comparison between some members of this family, may interfere with the reading and interpretation of CFTR specific data. Furthermore, most are not detailed and/or updated with recent disease features, besides not being freely accessible, nor containing CFTR variants information or allowing comparison between a wt and mutant protein interactions, essential to understanding the progression of this disease. yFi-MAP is developed and publicly available as a complementary resource to the existing tools. Moreover, CyFi-MAP content benefits from being discussed and approved by domain experts (available on the GitHub page, section Team). The incorporation of CFTR molecular mechanisms in a single resource represented in a standardized and inter-exchangeable manner represents one of the main advantages of CyFi-MAP. This tool allows the user to: (i) follow CFTR pathways; (ii) acquire its functional context and sub-cellular localization through the visualization of the cell compartments represented in the map; (iii) interpret differences between wt-CFTR and F508del-CFTR where unique interactions with CFTR variant are highlighted.

With this goal in mind, in this work, we aimed to build a repository of the available CFTR-related knowledge as a disease map named CyFi-MAP. CyFi-MAP is distinctive from other CFTR databases by being the first CF disease map and aiming to archive molecular mechanisms and biological pathways reported to be relevant to CFTR in a standard and consistent way. This repository is continuously updated with careful manual annotations and is expected to expand in both molecular mechanisms and in CFTR mutations representation and to be continuously updated based on the published literature. Through the map, it is possible to visualize CFTR pathways in protein cycle processes described in a sequential network, resulting in a more global biological interpretation and a faster understanding of CF pathophysiology.

Results

Concept and features. CyFi-MAP offers a resource that accurately (i.e., adequately confirmed in the literature) and graphically illustrates CFTR molecular pathways in an easy-to-read manner by the scientific community. Given that CF is caused exclusively by mutations in the CFTR gene that alter multiple cellular functions, the CyFi-MAP information was organized according to the CFTR life cycle, from its biogenesis to degradation, where sub-maps were developed: (i) one representative of wt-CFTR and (ii) one of F508del-CFTR.

The main differences between these submaps are given by the representation of some of the key processes side-by-side (available in the Map section), in order to facilitate submap comparison at molecular mechanism level. Additionally, a scheme is also available with CFTR traffic pathways inside the cell where the major physical alterations (e.g., traffic impairment and mucus clogging) in wild-type vs the mutant in airway epithelial cells are depicted (Fig. S1). An overview of the key CFTR processes/modules included in CyFi-MAP is given to inform and guide the user to the map content (Fig. 1).

Pathway inclusion strategy. We included information on CFTR interactors that was confirmed in a minimum of two published references. We focused on those that studied airway epithelial cells and used methodologies that allowed the detection of physical interaction between components (such as immunoprecipitation or nuclear magnetic resonance spectroscopy (NMR)). We also captured information on confidence and accuracy to each interaction in the map. Specifically, this step resulted in 296 research papers providing physical evidence between proteins and more than 1000 papers reviewed from PubMed (for more details on map curation see “Methods” section).

CyFi-MAP presents features that allow retrieving information visually such as (1) different types of interactions between the entities (i.e., activation, trafficking and inhibition—more details in Fig. S2 in Supplementary material), (2) the glycosylation (i.e., form B or C) and folding status (N-glycosylation) of CFTR, (3) the proteins that bind uniquely to the F508del-CFTR protein, (4) identification of each life cycle steps included—additional information in Supplementary material, and (5) cell organelles specific interactions represented through images, with the entities (i.e., proteins, complexes, ions, and others) adequately located in the biological compartments, differentiating between organelle lumen and the cytosol.
CyFi-MAP navigation. Map availability. The source of these schemes are available at the map online repository (https://cysticfibrosismap.github.io/) and the CyFi-MAP can be accessed online and explored interactively via the Molecular Interaction NetwoRks VisuAlization (MINERVA)37.

Online and interactive navigation. In the form of an interactive diagram via the MINERVA platform (Fig. 2), CyFi-MAP provides the capacity to easily follow CFTR interactions starting with its folding until degradation, with every intermediate step described (see supplementary material). The MINERVA platform allows easy navigation and exploration of the CF related molecular pathways available in CyFi-MAP. The user can also zoom in and find various details about proteins of interest (such as location of their interactions with other biological elements inside a cell, information on the protein name/alternative names and identification names in several databases such as Ensembl, human gene nomenclature, UniProt etc.; the annotation information is linked to the biological resources via direct urls). The user can also filter and extract information regarding the type of interaction through the edge colour (see Fig. S2 in Supplementary Material).

Content of the CyFi-MAP. The progression of CF disease is driven by the deregulation of multiple cell processes due to the loss of CFTR function. Hence, CyFi-MAP has greatly focused on the CFTR proteostasis network, as it encompasses alterations occurring on all those processes. Along with the map it is possible to observe several numbers which represent each step on the CFTR life cycle (a brief description of these is given in supplementary material) although the order indicated is not deterministic, serving only as an indication of all the processes in which CFTR is involved. The content included in the CyFi-MAP can be divided into four mains aspects: (i) CFTR synthesis and production, (ii) maintenance at the PM in the functional state, (iii) traffic and (iv) degradation.

CFTR synthesis and production. The folding of CFTR is highly regulated in the ER before the protein is allowed to proceed along the secretory pathway to the PM. This information is represented in the Folding module at both submaps hence, it is possible to visualize it side-by-side in the GitHub link.

Beginning with the wt-CFTR, folding starts with the nascent CFTR polypeptide chain being translocated to the ER membrane (step [1]), after which the N-glycosylation occurs (step [2]), a posttranslational modification during protein synthesis in the ER that is critical for PM expression and function (Fig. 3A)38–40. In CyFi-MAP, the different glycosylation states of wt-CFTR are represented and possible to follow. N-glycosylation starts with the addition of oligosaccharide residues (glucose3-mannose9-N-acetylglucosamine2) to CFTR (Fig. 3A, step [2]). At this step, several chaperones and co-chaperones bind to wt-CFTR to assist with folding. The folding process includes at least four ER quality control (ERQC) checkpoints that are involved in assessing CFTR correct folding41. As the process consists of the subsequent trimming of glucose, they are initially identified with three glucose (G3), the first checkpoint, before binding to calnexin (CANX) with only one (G1), the second checkpoint (Fig. 3A, steps [3–6])42.

F508del-CFTRs mostly targeted to degradation in the first checkpoint (Fig. 3B, step [2]), hence the other ERQC checkpoints are only represented at the wt-CFTR submap. There, is possible to follow the second checkpoint, where the protein enters the CANX cycle for additional rounds of refolding, the third checkpoint, where
Figure 2. CyFi-MAP Github and MINERVA platform. In the image (A) is possible to see the GitHub page with the objective to contextualize this work and provide access to the interactive MINERVA platform. In image (B) and (C) are represented both submaps, wt-CFTR and F508del-CFTR respectively, where it is possible to navigate and explore the pathways and interactions included in CyFi-MAP, providing generic information about the proteins and components.
specific signals when exposed lead to ER retention, or the fourth checkpoint, with the recognition of an export motif to leave ER (Fig. 3A, step [7] and step [8]). During these checkpoints, wt-CFTR can be recognized as
misfolded and move to the degradation or achieves an incomplete glycosylated state known as form B, which allows it to proceed to Golgi. A more detailed description of these checkpoints is in the degradation chapter.

Although insertion in the membrane of ER and ERQC checkpoints that lead to degradation are identified as different steps, there is evidence that co-translational folding and degradation occur.

After CFTR folding has been successfully achieved, the protein is ready to proceed along the secretory pathway to the Golgi apparatus, where its oligosaccharide structure is further modified by multiple glycosylation events generating its mature form, known as C form, which will be transported to the PM\(^44\). F508del-CFTR, because is highly degraded at ER, from the moment is depicted outside this organelle acquires a dark red colour, representing a rescue protein rF508del-CFTR.

**Maintenance at the plasma membrane in the functional state.** After delivery to the PM, wt-CFTR is regulated at multiple levels namely: (1) PM stabilization, at specific PM sites; (2) activation/channel shut down, where phosphorylation/dephosphorylation cycles activate/inactivate the channels; (3) ion channels and transporters regulation, concerning the regulation of and by other PM proteins; and (4) endocytosis, in either CCVs or caveolae vesicles.

In contrast, rF508del-CFTR is characterized by (1) PM stabilization; (2) channel shut-down; and (3) endocytosis, with consequent degradation. The reduced number of modules at PM is representative of the instability, loss of function and accelerated endocytosis that characterize this mutated protein.

In the PM stabilization module, Postsynaptic density 95, disks large, zonula occludens-1 (PDZ) domain-containing proteins are the main characters, responsible for anchoring CFTR to the PM\(^45\). Mechanisms such as cytoskeletal activation are represented, which enable PM anchoring and tethering of wt-CFTR to the PM (Fig. 4A, step [11]). TherF508del-CFTR stabilization module includes a lower number of interactions with PDZ proteins and acquisition of new interactions when compared to the wt-CFTR module (Fig. 4B, step [5]).

PDZK1 (CAP70) is illustrated at PM in the wt-CFTR submap to be able to potentiate the CFTR chloride channel activity by cluster two CFTR molecules (wt-CFTR submap, step [12])\(^60\) (Fig. 4A, step [12]).

Gating and channel shut down depict variants involved in CFTR activation/inactivation (Fig. 5). CFTR is regulated through CAM and phosphorylated by proteins such as protein kinase A (PKA) and Protein Kinase C Epsilon (PKCE)\(^47,48\) (Fig. 5 step [13]). This includes proteins such as β2-adrenergic receptor (ADRB2)\(^49\), A2B receptor (ADORA2B)\(^50\) and adenylyl cyclase I (ADCY)\(^51\) which participate in cAMP/PKA signalling therefore activating wt-CFTR and the transport of chloride.

The Channel shut-down present in both submaps includes dephosphorylation of CFTR and the proteins involved in its triggering—such as protein phosphatases, receptors, phospholipases, and others\(^23\).

The transporters and ion channel regulation module is only present at the wt-CFTR submap, and besides directly binding to wt-CFTR also allows to visualize PDZ proteins role as intermediates between them, maintaining the proteins in close proximity and leading to changes in their respective functions (Fig. 5B/C, steps [16], [17] and [18]). Proteins such as Solute Carrier Family 26 Member 6 (SLC26A3, also known as DRA), Solute Carrier Family 26 Member 6 (SLC26A6, also known as PAT1), Anoctamin 1 (ANO1, also known as TMEM16A) and epithelial sodium channel (ENaC) were included in this module\(^48\).

**Traffic.** In the secretory pathway, traffic is essential for all processes since folding-processing and function to degradation. CFTR traffic processes start by Coat Protein complex II (COPII) vesicles, responsible for its transport between ER and Golgi, from where wt- and rF508del-CFTR reach the PM (wt-CFTR submap, step [9])\(^62\). F508del-CFTR traffic between ER and Golgi is depicted through the COPII vesicles module with the same mechanisms as wt-CFTR (rF508del-CFTR submap, step [3])\(^33,34,57\). Although this information is only supported by high-throughput research articles and with only one paper for each interaction with the mutated protein, it was one of the exceptional cases that were selected to CyFi-MAP given that is explained by the functional context provided in wt-CFTR. A list with the exceptions is in the additional information section of the supplementary material.

From this point, the endocytosis module depicts a wt-CFTR association with endocytic adaptors undergoing CCV-mediated and caveolar endocytosis (Fig. 6A, steps [19, 20] respectively), whereas in rF508del-CFTR only the last one is depicted (Fig. 6B, step [9]).

CFTR is endocytosed and arrives at the sorting endosome from which wt-CFTR moves back to the PM, either directly—Recycling Module (Fig. 7A, step [22], [23])—or through the Golgi—Golgi Module (Fig. 7A, step [24])—or to degradation—Degradation Module (Fig. 7A, step [26]). In these processes, several proteins of the Rab family are represented as they are essential for CFTR traffic\(^58\). In the case of rF508del-CFTR, at the sorting endosome, it is sent to degradation (Fig. 7B, step [10]).

**Degradation.** During folding and processing, several quality control proteins target misfolded CFTR to degradation, therefore, in each of the organelles—ER and Golgi—there is a module called Degradation.

The four ERQC checkpoints can lead to wt-CFTR degradation at the proteasome. In the case of F508del-CFTR, only the first ERQC checkpoint, called chaperone trap, is represented with the binding of four chaperones that strongly attach to the misfolded protein (i.e., Heat Shock Protein Family A (Hsp70) Member 4 (HSPA4), Heat Shock Protein Family A (Hsp70) Member 8 (HSPA8), DnaJ Heat Shock Protein Family (Hsp40) Member A1 (DNAJA1) and DnaJ Heat Shock Protein Family (Hsp40) Member B1 (DNAJB1))\(^50,58\). Furthermore, the F508del-CFTR presents a degradation pathway in contrast to wt-CFTR: the sumoylation (Fig. 3B)\(^60\). At Golgi, CFTR degradation depicts its targeting to degradation via the lysosome. In the PM, the rF508del-CFTRs targeted to degradation by ubiquitination a consequence of the destabilization of the protein (F508del-CFTR submap, steps [7], [8] and [9]). There, proteins involved in its degradation at ER attach to itin the PM (i.e., HSPA8, STIP1 Homology And U-Box Containing Protein 1 (STUB1/CHIP), Stress-Induced
Figure 4. PM stabilization in wt-CFTR (A) and F508del-CFTR (B) submaps. Wt-CFTR represented in blue is delivered to the apical PM in the C form, as indicated at the protein, and several proteins responsible for the anchoring to the PM bind. The lines show different types of interactions, where orange lines indicate traffic, being possible to see in chloride (Cl⁻) transport across the membrane in orange. The black lines indicate binding/stimulation and the red one’s inhibition. F-actin and Arp2/3 are represented as complexes, with several proteins with a grey line separating them from the rest. In blue are represented 1-phosphatidylinositol-4-phosphate (PIP) which by the action of Phosphoinositide Kinase, FYVE-Type Zinc Finger Containing (PIKFYVE) is converted to 1-phosphatidylinositol-4,5-bisphosphate (PIP2) with a blue line depicting synthesis. Although rF508del-CFTR reaches the PM, the binding to new proteins highlighted in yellow diminishes its stabilization and anchoring therefore accelerating its endocytosis and consequent degradation.
Figure 5. PM pathways in wt-CFTR submap. (A) Gating where is possible to see adenosine 3,5-cyclic monophosphate (cAMP) role in Protein Kinase CAMP-Activated Catalytic Subunit Alpha (PRKACA) phosphorylation that leads to CFTR activation and transport of chloride across the membrane. Other proteins are involved such as Adrenoceptor Beta 2 (ADRB2), Protein kinase CK2 (formerly known as casein kinase II), Adenosine A2b Receptor (ADORA2B) and Protein Kinase C Epsilon (PRKCE). (B) Shut-down with the action of Lysophosphatidic Acid Receptor 2 (LPAR2), serine/threonine protein kinase complex (AMPK), Phospholipase C beta ½ (PLC beta ½) and Protein phosphatase 2 (PP2A) where the chloride transport is inhibited. (C) Transporters and Ion channel regulation, in which the proteins that regulate and are regulated by CFTR at the PM are represented with the interactions detected.
Phosphoprotein 1 (STIP1) and others. At the sorting endosome, in both cells, the protein is targeted to degradation by lysosome/proteasome although with different interactors assisting (i.e., Tumor Susceptibility 101 (TSG101), Hepatocyte Growth Factor-Regulated Tyrosine Kinase Substrate (HGS), Charged Multivesicular Body Protein 4B (CHMP4B), and others) bind uniquely to rF508del-CFTR (Fig. 7B, step [10]).

**Discussion**

CF is the most common life-threatening autosomal recessive disease in the Caucasian population. Caused by an absent/dysfunctional CFTR channel that leads to an impairing balance of ions across the membrane, CF is characterized by affecting several organs, especially the lung. CFTR seems to be involved not only in the transport of ions but also in the regulation of other channels, working in a dynamic network that modulates its activity. There have now been more than 30 years of scientific discoveries with new milestones achieved every year in our understanding of intracellular interactions after CFTR loss of function that control the progression of this disease. The knowledge obtained over this research has enabled diagnosis and discovery of therapies that increased life expectancy. Yet no final curative treatment has yet been developed for CF disease.

The increasing data available on public databases lead to the improvement of tools to filter and extract relevant knowledge required for the discovery of therapeutic targets. With this need, disease maps were developed as a multilayer-readable network that allows representing increasingly complex and extensive information in

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**Figure 6.** wt-CFTR (A) and rF508del-CFTR (B) endocytosis mechanisms of internalization from the PM. In the image A two types of internalization of wt-CFTR are present, through clathrin coated vesicles [19] and through caveolae vesicles [20]. As is represented, in the first, several proteins are involved, since the clathrin triskelion complex to cytoskeletal F-actin MYO6 complex and other proteins assisting the process. In rF508del-CFTR, only the complex Caveolin 1 (CAV1)/ Caveolin 2 (CAV2) was found in this protein endocytosis [9], with assistance of Flotillin 2 (FLOT2).
CyFi-MAP included data: wt-CFTR versus F508del-CFTR submap.  

CFTR life cycle can be impacted at several steps, with the most common mutation, F508del-CFTR, subjected to ER retention and degradation when not rescued through low temperature or chemical compounds. In fact, rF508del-CFTR is characterized by barely reaching the PM under physiological conditions, presenting reduced chloride/bicarbonate transport after being rescued, as well as enhanced endocytosis and degradation, with consequent dysregulation of other PM proteins [24,67]. Due to F508del-CFTR incapability to achieve a competent intermediate, the protein becomes
CyFi-MAP expansion and future work. Given the fact that new data are generated continuously, and CF aspects are yet to be included, the CyFi-MAP is constantly developed with support from the community and funding agencies. CyFi-MAP benefits from major features of the MINERVA platform via its online distribution: comments and suggestions from users with regard to changes in the map content (addition, removal, update) can be analysed directly by curators and addressed potentially in the map after further refinements. In this way, users can promote active discussions and knowledge exchange to build an increasingly accurate and continuously updated CF disease map. Notwithstanding, of specific interest as a future direction, is to include in CyFi-MAP the specific steps involved in the trafficking of CFTR processes targeted by compounds, depicting this way the specific target/mechanism where each of them acts. Additionally, we anticipate including a diagram focusing on the process where unique proteins interact with rF508del-CFTR in the ER to provide a broader view of the possible interactions and they may appear in a future version of CyFi-MAP with less stringent criteria for inclusion.

Methods

CyFi-MAP construction. The development of the CyFi-MAP follows the disease map development protocol, using primarily Kondratova et al. and Mazein et al.11,24. Specifically, three main steps entail the construction of CyFi-CFTR (Fig. 8):

I. The first step consisted of searching relevant CFTR-related information, selecting a total of 297 research papers and more than 1000 reviewed articles. A complete list of publications consulted for the CyFi-MAP development is available in https://cysticfibrosismap.github.io/. The CF disease hallmarks were obtained from peer-reviewed research papers, domain experts’ suggestions and advice, previously documented and validated pathways, and curated up-to-date database (including Reactome9/KEGG4/MetaCore from Clarivate Analytics. Please see Content curation subsection for details). This task also involved the analysis of the collected pool of data, followed by the curation of the most relevant CFTR-related knowledge (Fig. 9).

II. The second step comprised the effective diagram building, assuring the correct level of detail and the most appropriate and aesthetically output, to guarantee that the resulting map is as readable and user-friendly as possible. The biological mechanisms representation follows the Systems Biology Graphical Notation (SBGN) notation and was built in the yEd Graph Editor using the SBGN Palette (https://yed.yworks.com/support/manual/layout_sbgn.html). The yEd Graph Editor is a freely available graph editor providing functionality to manage large-scale graphs including: (i) features that considerably facilitate the diagram drawing process such as friendly user interface, drawing guides, zooming on the diagram and easy application of specific aesthetics (e.g. same colour for nodes/edges, curved connectors) to an individual or multiple elements; (ii) algorithms for automatic layout (details on using yEd to automatically layout SBGN-related diagrams are given in e.g.75). The yEd Editor also incorporates the SBGN Palette that permits the direct representation of the SBGN-specific elements into the yEd inner GraphML format. After the CyFi-MAP was developed in yEd, we converted it into the SBGN standard format by
using the ySBGN converter, (a bi-directional converter between the SBGN and yEd GraphML formats, available at https://github.com/sbgn/ySBGN). Further, the CyFi-Map SBGN diagram was loaded to the MINERVA online platform. The organelle images (developed manually and expert-revised) aim to facilitate visualization of the mechanisms at the top level; thus, special attention was given to the localization of the interactions in each organelle.

III. The third step in the construction of the CyFi-MAP was the map exploration via the MINERVA platform. In a first approach, the construction focused on the creation of small organelle-specific maps, illustrating $\text{CFTR}$-relevant processes on those locations. The maps included $\text{CFTR}$ interactions covering its intracellular and intraorganellar traffic. Later, these were improved upon by the addition of other, more widespread $\text{CFTR}$ processes and pathways, which allowed a more effective integration of the existing data. The resulting cell-wide map is expected to be continuously evolving with user input and consistent expert curation. The map is available through the web platform MINERVA that provides interactive and exploratory features.

The current version of CyFi-MAP has been manually curated by CF domain experts. To ensure a continuous updating of this resource, both regular expert verification of new information, as well as regular user input are deemed essential to achieve an accurate representation of current CF data. Constant feedback from cell biologists, biochemists, physiologists and bioinformaticians contributed to a comprehensive representation of the various layers of information.

Inside CyFi-MAP, each process comprises pathways that include proteins (as individual entities or as complexes) and different types of chemical species (ions and lipids) interacting among themselves. Nodes represent entities (i.e., proteins, ions or complexes) and edge colours correspond to processes (i.e., activation, inhibition, synthesis, or in some cases movement of entities inside the cell).

CyFi-MAP currently comprises 618 nodes and 420 edges, with 426 nodes and 307 edges in $\text{wt-CFTR}$ and 216 nodes and 117 edges in $\text{F508del-CFTR}$. In total, entities presented at both submaps are classified into 193 proteins, 25 complexes, 5 ions and 5 simple molecules in $\text{wt-CFTR}$ and 98 proteins, 12 complexes, 1 ion, and 2 simple molecules in $\text{F508del-CFTR}$.

**Content curation.** The data used in CyFi-MAP was obtained by manual human search, curation and validation with domain experts in three main sources:

**Pathway databases.** The curation process started by reviewing previous attempts to summarize CF information in these signalling networks. Pathways from MetaCore (Clarivate Analytics), Reactome and KEGG were
reviewed in order to analyse the pathway availability for the CF disease\textsuperscript{67}. Major CF-related pathways were retrieved from these databases and confirmed in the literature for their accuracy.

**Literature.** The main hallmarks of CF were extensively searched in PubMed. As \textit{CFTR} is the protein that plays a central role in the map, direct interactions with it were very carefully selected, following strict criteria. Considering that the lung is the most affected organ, the focus was on human airway epithelial cells, although other cell types were included when described on review publications. In both levels, if the study do not agree with the criteria is rejected. The 3rd level consist on finding the location of the interaction inside the cell (e.g. ER, Golgi, cytosol, PM) followed by the type of interaction (binding or inhibition). The 4th level confirms confidence to the interaction, consisting on the search for publications that support the information. The 5th level confirms information related to the protein after being selected (e.g. does it belong to a complex? Which pathway does it belong to?). The protein is manually added to the yEd Graph Editor used to build CyFi-MAP with the name accordingly to the HGNC nomenclature.

**Databases.** Among the web resources used for data gathering, the most significant were GeneCards\textsuperscript{78}, Stringdb\textsuperscript{79}, Biogrid\textsuperscript{80}, UniProt\textsuperscript{81} and HGNC (HUGO Gene Nomenclature Comittee)\textsuperscript{82} which were used to confirm the correct names of proteins/genes, their known function and their interactors. For each protein, the name was checked in HGNC. UniProt and Genecards were used to search for alternative names for the same protein so as to find the correct HGNC designation\textsuperscript{79,81}\textsuperscript{82}. Often, although a protein complex is known to interact/participate in a \textit{CFTR} process, the specific proteins that constitute that complex are not described in the original literature report.
Accordingly, proteins reported in the literature to interact with CFTR as part of larger complexes were searched for in databases to find the protein components of the complex.

During this step, name disambiguation must be considered in order to find all data related to that protein and also to not repeat proteins. For instance, by looking for syntaxin 5, names such as Syn5 and STX5 are also available for the same protein. The same happens for Golgi Associated PDZ And Coiled-Coil Motif Containing protein, known as GOPC although other names such as CAL and FIG are referred to on research papers and used to retrieve as much information as possible.

**Diagram building.** CyFi-MAP was built using yEd graph editor (https://www.yworks.com/) using the SBGN Palette, and the data was represented based on the SBGN standard. This notation provides a knowledge representation language used in the illustration of molecular pathways and protein interactions as the standard notation for disease maps. Presenting three languages that provide different types of knowledge illustration allow to adapt on the level of detail intended to be highlighted on the map, including the following layers: Activity Flow to depict interactions with process direction, Process Description detailed specific mechanisms, and Entity Relationships which describe the mechanisms without a sequential process.

CyFi-MAP was implemented following the SBGN Activity Flow, in order to provide a compact, sequential and easy-read format or involving signalling pathways. This language is useful to represent the flow of information in biological sequences/pathways in a way that the information can still be captured for underlying mechanisms of unknown influence.

Each subcellular organelle (ER, Golgi, endosome, etc.) were drawn manually and added to the map as a background image for graphical representation of the different subcellular compartments. Additionally, each interaction provides information on itself. Depending on the selected edge different types of interactions can be found on CyFi-MAP, namely (see Fig. S1 on supplementary material for more details): (1) activation, representing a normal binding, (2) synthesis, when an altered product is released, (3) trafficking, representing movement inside the map, and (4) inhibition, when the interaction inhibits a function.

**Map exploration via the MINERVA platform.** CyFi-MAP diagrams are available in the platform MINERVA accessible through GitHub (https://cysticfibrosismap.github.io/). The project description and key processes (shown side-by-side, represented through images to allow the comparison between wt-CFTR and F508del-CFTR submaps on the CyFi-MAP) are given on the website. Starting at the cell level, it is possible to identify the main differences between the submaps (Fig. 1). This view is relevant in order to compare the cells in presence of the two proteins since wt-CFTR is transported across the secretory pathway to the PM and endocytosed to be either degraded or recycled back to the PM, whereas most F508del-CFTR is retained at the ER from where it is sent for degradation. This impairment leads to the so-called ‘CF pathogenesis cascade’, which does not occur for wt-CFTR. In the cell level view, it is possible to observe these features, allowing the extraction of relevant knowledge.

Additionally, wt-CFTR and F508del-CFTR submaps were divided into modules, each representing key processes of its life cycle in order to guide the navigation through the map. To compare information in both submaps, images placing the modules folding, stabilization and sorting side-by-side are available. Mutation-specific proteins are highlighted in a different colour to emphasise differences between wild-type and mutated phenotypes.

The interactive web platform MINERVA allows accessing an interactive CyFi-MAP to navigate and explore its molecular networks. This tool provides automated content annotation, direct feedback to content curators and SBGN-compliant format. Navigation in CyFi-MAP is similar to navigation in Google Maps being possible to display differences between submaps (Fig. 1). This view is relevant in order to compare the cells in presence of the two proteins since wt-CFTR is transported across the secretory pathway to the PM and endocytosed to be either degraded or recycled back to the PM, whereas most F508del-CFTR is retained at the ER from where it is sent for degradation. This impairment leads to the so-called ‘CF pathogenesis cascade’, which does not occur for wt-CFTR. In the cell level view, it is possible to observe these features, allowing the extraction of relevant knowledge.

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All suggestions will be analysed by curators and CF domain experts to maintain CyFi-MAP quality and accuracy.

The user can contribute to CyFi-MAP by adding comments with questions, corrections or additions to the map. These will be visible to other users and developers. To add a comment to CyFi-MAP during navigation, right-click on the specific location and choose to add a comment. It is possible to choose a specific identity to link the comment, such as the protein or reaction, or to remain ‘general’, which will link the comment to the location the user chooses. The remaining fields allow the user to fill in the name and email in order to facilitate communication with the developers and to clarify any questions that may emerge. Last, there will be a box where comments can be added (Fig. S2). Any supporting information provided will be helpful to incorporate the changes into the map. After sending the comment, it will not be possible to correct it and it will be visible on the map publicly. Details on adding user’s comments in the underlying MINERVA platform are given at https://minerva.pages.uni.lu/doc/user_manual/v15.0/index/#add-comment.

The CyFi-MAP allow exploring the map with and without seeing the comments provided by the users by clicking on the checkbox Comments above in the map toolbar. These comments will allow the map users to benefit from the domain knowledge and expertise of researchers and to collect valuable information for the research community. All suggestions will be analysed by curators and CF domain experts in agreement with the pre-established curation process to maintain CyFi-MAP quality and accuracy.
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Author contributions
C.P.—conceptualization; data curation; formal analysis; investigation; methodology; original draft preparation; review and editing; validation. A.M.—conceptualization; methodology; review and editing; supervision. C.F.—review and editing. M.G.—review and editing. K.K.—review and editing. M.O.—MINERV A online version. I.B.—MINERV A online version. M.D.A.—conceptualization; funding acquisition; resources; review and editing supervision. A.O.F.—conceptualization; data curation; methodology; resources; review and editing; supervision; validation. All authors have read and agreed to the published version of the manuscript.

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The authors declare no competing interests.

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