



Belgisch Mucoviscidose Register Registre Belge de la Mucoviscidose The Belgian Cystic Fibrosis Registry (BMR-RBM-BCFR)

# ANNUAL REPORT BELGIAN CYSTIC FIBROSIS REGISTRY 2014

# Annual report Belgian Cystic Fibrosis Registry 2014

AUTHORS Simeon Situma WANYAMA Muriel THOMAS Anne MALFROOT





























#### Scientific Institute of Public Health (WIV-ISP)

Operational directorate Public health and Surveillance Unit Healthcare Services Research Rue Juliette Wytsmanstraat 14 | 1050 Brussels | Belgium

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#### LIST OF ABBREVIATIONS AND DEFINITIONS

- ABPA Allergic bronchopulmonary aspergillosis
- **BCFR** The Belgian Cystic Fibrosis registry
- **BCFA** Belgian Cystic Fibrosis patients' Association
- **BMI** Body mass index is a measure of relative weight based on an individual's mass and height. It is defined as the individual's body mass divided by the square of their height with the value universally being given in units of kg/m2.
- **BMR-RBM** Belgisch Mucoviscidose Register Registre Belge de la Mucoviscidose
- **Bronchoalveolar lavage** is a medical procedure in which a bronchoscope is passed through the mouth or nose into the lungs and fluid is squirted into a small part of the lung and then collected for examination.
- **CDC** The Centers for Disease Control and Prevention is the leading national public health institute of the United States. Its main goal is to protect public health and safety through the control and prevention of disease, injury, and disability.
- **CDC growth charts** These consist of a series of percentile curves that illustrate the distribution of selected body measurements in children.
- **CF** Cystic Fibrosis
- **CFRD** Cystic Fibrosis Related Diabetes refers to a form of diabetes as a direct consequence of having CF.
- **ConsultRN** a module of the eHealth platform that allows to obtain the demographic data of patients from the national registry database.
- **Dual energy X-ray absorptiometry** (DXA, previously DEXA) is a means of measuring bone mineral density (BMD).
- **ECFSPR** European Cystic Fibrosis Society Patient Registry
- **eHealth** As public institution, the eHealth platform promotes and supports the exchange of electronic information between all stakeholders in health care. eHealth also acts as a Trusted Third party for coding and anonymizing personal health-related data.
- **FEV**<sub>1</sub> Forced Expiratory Volume in one second is the volume of air that can forcibly be blown out in one second, after full inspiration
- **FVC** Forced Vital Capacity is the volume of air that can forcibly be blown out after full inspiration, measured in litres.
- INAMI Institut national d'assurance maladie-invalidité
- **Pseudomonas aeruginosa** is the most important pathogen in the CF airway. *P. aeruginosa* is acquired from environmental reservoirs and can cause both acute and chronic infections, depending on the clinical context.

**rhDNase** - Recombinant human deoxyribonuclease is an enzyme that breaks down DNA strands in airway secretions, hydrolyzes the DNA present in sputum/mucus of CF patients, reducing viscosity in the lungs and promoting clearance of secretions.

**RIZIV**- Rijksinstituut voor ziekte- en invaliditeitsverzekering

**TEPD** - Transepithelial potential difference is the voltage across an epithelium, and is the sum of the membrane potentials for the outer and inner cell membranes used in CF diagnosis

**VUB** - Vrije Universiteit Brussel

**WIV - ISP - IPH -** Wetenschappelijk Instituut Volksgezondheid / Institut Scientifique de Santé Publique / Scientific Institute of Public Health

#### CONTRIBUTORS, MEMBERS OF THE BOARD OF THE BCFR

D. Baran (Hôpital Erasme, Bruxelles)

H. Boboli (CHR de la Citadelle, Liège)

F. De Baets (UZ - Gent, Gent)

K. De Boeck (UZ - Gasthuisberg, Leuven)

E. De Wachter (UZ - Brussel, Brussel)

C. Knoop (Hôpital Erasme, Bruxelles)

P. Lebecque (Cliniques Universitaires St-Luc, Bruxelles)

M. Lequesne (UZ - Antwerpen, Antwerpen)

A. Malfroot (UZ - Brussel, Brussel)

V. Van Casteren (WIV-ISP, Brussel)

S. Van Daele (UZ - Gent, Gent)

S. Verhulst (UZ - Antwerpen, Antwerpen)

F. Vermeulen (UZ - Gasthuisberg, Leuven)

#### **REGISTRY MANAGEMENT**

#### Scientific Institute of Public Health (WIV - ISP)

OD Public Health and Surveillance Rue J. Wytsmanstraat, 14 1050 Brussels

V. Van Casteren: Head of unit

Tel.: 02/642.50.30 Fax: 02/642.54.10

M. Thomas : Project leader

Tel.: 02/642.50.23

S. S. Wanyama: Biostatistician

Tel.: 02/642.57.67

Y. Pirson : Administrative

Tel.: 02/642.57.41

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https://www.wiv-isp.be/en/search/biblio/cystic%252Bfibrosis/ss\_biblio\_type/Sci%252E%20report%2C%20recommendat%C2%B0%2C%20guidance%20doc%252E%2C%20directive%2C%20monograph

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#### ACCREDITED BELGIAN CF REFERENCE CENTRES

#### Mucoviscidose referentiecentrum UZ Brussel

UZ Brussel Laarbeeklaan 101 1090 Brussel

A. Malfroot, E. De Wachter, E. Vanderhelst, S. Vincken

#### Muco-Referentiecentrum Antwerpen

St.-VincentiusziekenhuisUZ AntwerpenSt.-Vincentiusstraat 20Wilrijkstraat 102018 Antwerpen2650 Antwerpen

V. Nowé, I. Stappaerts M. Lequesne, S. Verhulst, K. Van Hoorenbeeck

#### Referentiecentrum voor Mucoviscidose UZ Gent

UZ Gent De Pintelaan 185 9000 Gent

F. De Baets, S. Van Daele, E. Van Braeckel

#### Muco-Referentiecentrum Gasthuisberg Leuven

UZ Leuven – Campus Gasthuisberg Herestraat 49 3000 Leuven

K. De Boeck, F. Vermeulen, L. Dupont, M. Moens

#### Centre de référence de la Mucoviscidose UCL

Cliniques Universitaires St-Luc Avenue Hippocrate, 10 1200 Bruxelles

P. Lebecque

#### Centre Liégeois de rééducation fonctionnelle pour la Mucoviscidose

CHR La Citadelle Boulevard du 12ème de ligne, 1 4000 Liège

H. Boboli

Clinique de l'Espérance Rue St-Nicolas, 447-449 4420 Montegnée

F. Piérart

#### Institut de Mucoviscidose ULB

Hôpital Universitaire des Enfants Reine Fabiola Avenue J.J. Crocq, 15 1020 Bruxelles

G. Casimir, L. Hanssens

Hôpital Erasme Route de Lennik, 808 1070 Bruxelles

C. Knoop

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Last year was a transition year with a lot of challenges to have the registry migrated and integrated in the new Healthdata.be platform and organize training on the new data collection system. Thank you all for your efforts and patience. It is foreseen that this system will be in use for a longer period and hence are confident of less delays in future.

We would like to most sincerely thank all the partners involved in the provision of various services that go toward better care of the people with CF. We single out the National Institute for Health and Disability Insurance (RIZIV-INAMI) for their continued financial support and the CF reference centres and their multidisciplinary teams of health care professionals for their continued support to the registry and care to the patients. Finally, we also thank the patients and their parents or care givers for their willingness to participate in the annual data collection processes. This report, and the outcomes of various research questions, enhances the continued monitoring of CF disease by providing a better understanding of disease progression to researchers while stimulating ideas for research into novel treatments and disease management strategies.

The registry team

#### SUMMARY

This report presents the data collected in 2014. It is our hope that the analysis of the registry data will provide readers with information on various aspects of CF and continue to provide an important tool for monitoring the patient's quality of care and trends.

Since its establishment in 1998, the Belgium CF Registry (BCFR) has grown steadily and had 1230 patients registered in 2014. This number excludes two whose diagnosis for CF was revoked and seven without a confirmed diagnosis. There were 36 newly diagnosed patients in 2014, among them six adults, with a median age at diagnosis of 2.5 months with a range from birth to 60.0 years. Most of the newly diagnosed patients were genotyped, while 29 had sweat chloride values > 60 mmol/L.

Among the patients in follow-up in 2014, 52.0% were male and 58.4% adults with a median age of 21.3 years. This can be compared to the start of the registry 15 years ago when 39.0% were adults with a median age of 14.9 years. 45.4% of the patients are homozygous for the F508del mutation while 40.0% are F508del heterozygous. The main reasons for diagnosis of CF are acute or recurrent respiratory problems (42.7%) and failure to thrive (24.2%). About 17.0% were diagnosed via neonatal screening even though there is no national neonatal screening program in Belgium so far. Within the year, 10 deaths were reported (4 of them in transplanted patients) with age at death ranging from 11.5 – 76.9 years while 6 patients benefitted from a lung transplant. About 12.3% of the patients in the registry are living with a transplant.

Among the adults, the proportion of patients with BMI < 18.0 kg/m<sup>2</sup> continues to decline from about 36.3% in 1998 to 17.4% in 2010 and 13.6% in 2014; this decline was noted even amongst the F508del homozygous patients. Amongst the patients up to 20 years, the proportion with BMI below the tenth percentile has also been declining over the years. The above suggests better nutritional management in the patients. The patient population continues to record an improvement in lung function expressed as the mean percentage of predicted FEV<sub>1</sub>. Among the F508del homozygous patients, 38.0% of the children and 5.1% of the adults had FEV<sub>1</sub>  $\geq$  90.0% of predicted in 1998 compared to 52.9% and 7.0% in 2010 and 57.1% and 12.6% respectively among the children and adults in 2014.

The overall annual prevalence of *Pseudomonas aeruginosa* reported in 2014 was 41.3% and has been declining compared to a prevalence of 42.4% in 2012. The prevalence of the *Burkholderia cepacia* complex on the other hand had remained lower than 3.0% over the years till 2010. In 2011 the prevalence increased to 3.6%, (not statistically significant) while in 2013 and 2014 it was 4.5% and 3.7% respectively both statistically significant over the 2010 prevalence of 2.4%. There has also been a steady increase in

the prevalence of *Achromobacter xylosoxidans* from 5.9% in 2009 to 10.0% in 2014.

Thanks to improved disease management practises and novel treatments, the life expectancy and the quality of life of patients with CF has improved significantly when compared to CF cohorts a decade or two ago. The proportion of adult CF patients aged 18 years and above increases each year. But this progress is also accompanied by different challenges, expectations and disease related complications. CF related diabetes had a prevalence of 26.6% and 27.8% in non-transplanted adults in 2013 and 2014 respectively. Other complications include early osteoporosis, CF related arthritis /arthropathy. This requires specific care for the adult CF patient.

#### **CHAPTER 1: BACKGROUND**

This section briefly describes CF care in Belgium, the history, role and objectives of the Belgian CF Registry (BCFR) and the important contributions of the CF patients' association (BCFA), the National Institute for Health and Disability Insurance (INAMI - RIZIV) and the CF reference centers in the provision of care and management of CF.

#### WHAT IS CYSTIC FIBROSIS?

CF is a progressive hereditary disease with autosomal recessive transmission: only subjects who have inherited two disease causing mutations – one from each parent - are affected. Parents who are both carriers for a CFTR mutation associated with classical CF have a 1 in 4 chance of having a child with CF, in each pregnancy. It is commonly found in populations of white Caucasian descent, such as those of Europe, North America and Australasia. Prevalence is however different from country to country and is 1/2850 live births in Belgium<sup>[1]</sup>. The earliest clear medical descriptions of CF date from the 1930s<sup>[2, 3]</sup>. CF obviously existed prior to this dates even though it remained largely unrecognized and so went undiagnosed. In these early times, it was even thought of to be a result of witchcraft (http://www.cfmedicine.com/history/earlyyears.htm).

The disease is caused by the alteration (mutation) of the *CFTR* (CF Transmembrane Conductance Regulator) gene which is located on the long arm of chromosome 7. More than 2000 (http://www.genet.sickkids. on.ca/SearchPage.html) mutations have been identified in the *CFTR* gene since its discovery in 1989<sup>[4]</sup>, but not all are associated with classical CF. The *CFTR* gene codes for the CFTR protein. This is an ion channel involved in the regulation of chloride ion transport across the cell membrane. It is mainly found in the cell membranes of the respiratory and digestive tract, the sweat glands and the reproductive tract. The dysfunction of the CFTR protein leads to the production of sweat with a high salt content and mucus secretions with an abnormal viscosity causing dysfunction of many organs such as the lungs, pancreas and liver.

In the respiratory tract, thick mucus production results in persistent cough caused by chronic infection and inflammation leading to severe bronchial obstruction and finally lung destruction. In the pancreas, the sticky exocrine pancreatic secretions lead to obstruction and blocking of the ducts with secondary damage to the secretory gland tissue. Diminished secretion of pancreatic enzymes leads to fat and protein malabsorption causing steatorrhea (fatty stools) and failure to thrive. Fat malabsorption also causes deficiency of fat soluble vitamins (A, D, E and K).

Most of the children with CF have a history of recurrent chest infections, steatorrhea and failure to thrive. Newborns with CF can be affected

by meconium ileus: intestinal obstruction with vomiting, abdominal distension and delay in passing the first meconium stools. The spectrum of presenting features is very wide and can vary with the age at time of clinical presentation. The diagnosis is usually made in early childhood but in some patients with late or milder symptoms it can occur later into adulthood.

Although there is no national neonatal CF screening program in Belgium yet, there are some local initiatives where patients are screened. Infants with CF can be identified in the first weeks of life by assessing their blood immunoreactive trypsin (IRT) combined with the most frequent CFTR mutations. The sweat test remains the gold standard for the diagnosis of CF. In the majority of patients with typical features, the sweat test is diagnostic. It will reveal an excessive quantity of chloride (salt) (> 60 mmol/L). In atypical forms, the sweat test chloride levels can fall into the intermediate range (30-60 mmol/L).

It is advised to perform genotyping in all patients with CF to identify the CF causing mutations. The F508del is the most common mutation not just in the Belgian CF population but also worldwide.

Today most standard treatments are symptomatic and are essentially based on respiratory management (e.g. physiotherapy, mucolytics, antibiotics, antiinflammatories), digestive and nutritional management (e.g. pancreatic enzymes and hypercalorie diet). Due to medical progress and intensification of the care for patients with CF, the quality of life and the life expectancy have increased with most of the patients living to young adulthood.

However, to further improve the life expectancy and quality of life, new and more effective treatments are needed that target molecular defects and act upstream on what the symptomatic treatment do on the cascade of events.

More than 2000 mutations of the *CFTR* gene have been reported. They are grouped in six classes according to their functional defect: Class I: defect of protein synthesis; Class II: default of protein folding with premature degradation which interferes with the protein trafficking to the cell surface; Class III (gating mutations): responsible of deficient channel opening; Class IV: decrease of CFTR conductance channel; Class V: decreased amount of CFTR protein synthesis and Class VI; decreased stability of CFTR protein at the cell membrane. Some CFTR mutations have characteristics of more than one mutation class<sup>[21]</sup> and for many mutations it is not known to what mutation class they belong.

CFTR modulating therapies are therapies that target the basic defect. They are small-molecule pharmacologic agents that correct the function of

the defective CFTR protein. They do not correct the default in the gene itself. They are specific to certain mutations or class of mutations and thus effective only in patients carrying those specific mutations. There are different types of CFTR modulators: potentiators, correctors and read-through agents.

Currently two of those CFTR modulators are approved in the USA and in Europe for clinical use: ivacaftor (Kalydeco®) for patients carrying a class III (gating) mutation (G551D, G1244E, G1349D, G178R, G551S, S1251N, S1255P, S549N, S549R) and those carrying the R117H mutation and lumacaftor in association with ivacaftor (Orkambi®) for patients homozygous for the F508del mutation.

Clinically, ivacaftor (a potentiator) has been shown to improve the lung function and the BMI, to reduce the sweat chloride concentrations and improve the quality of life in patients aged 12 years or older with G551D and non-G551D gating mutations<sup>[5,6,7,8,13,14]</sup>. The efficacy and good tolerance of ivacaftor has also been documented in children aged 6 - 11 years<sup>[15,16]</sup> and 2 - 5 years<sup>[17]</sup> with a gating mutation.

In Belgium, ivacaftor is reimbursed (temporarily) for patients older than 6 years carrying one of the nine gating mutations since February 2016 and for children aged 2 - 5 years since October 2016. Patients who received a lung transplant are not eligible for this treatment. The physicians are required to register the data related to the outcome and its evolution on the treated patients to have the temporary reimbursement prolonged.

For patients carrying two copies of the F508del mutation, the combination of ivacaftor and lumacafactor (a corrector) (Orkambi®) has been shown to induce a significant but modest improvement in the FEV<sub>1</sub> % predicted (2.6 - 4.0%)<sup>[22]</sup>. Orkambi® was approved by the European Medicine Agency (EMA) in September 2015 and in 2017 the procedures to have it reimbursed in Belgium are underway.

Other CFTR modulating therapies are currently in development and are being evaluated in several ongoing clinical trials. For more information about new therapies we refer to the articles of Fajac I, Quon B and De Boeck K [24,25,26].

#### CF PATIENT CARE IN BELGIUM

Since 1999, 7 CF reference centres have been accredited by the National Institute for Health and Disability Insurance (INAMI - RIZIV) and receive financial support. An annual care and revalidation agreement (CF convention) for patients with CF is signed between each of the 7 CF reference centres and the RIZIV-INAMI<sup>[27,28]</sup>. Each centre has specific expertise in CF care and ensures multidisciplinary follow-up of the patients

in order to provide optimal medical, paramedical, psychological and social care to the patient and their relatives. Most of the persons with CF in Belgium are followed in one of the national CF reference centres and are registered in the national CF Registry (BCFR).

#### THE BELGIAN CF REGISTRY (BCFR)

The intent of a registry is to include, in a single database, the entire population of people with a given condition (or meeting a certain criteria), say chronic illness, within a defined geographical area.

The BCFR was started in 1999 as a scientific project initiated by the Medical Committee of the Belgian CF Association<sup>(1)</sup> and the seven CF-reference centres in Belgium collecting data of 1998. It was coordinated by the Vrije Universiteit Brussel (VUB) and co-sponsored by the CF-Patient organisation and the Fund Alfonse and Jean Forton of the King Baudouin Foundation. The VUB covered the overheads. After 5 years the scientific project came to an end and new sponsors were contacted.

In 2006, the RIZIV-INAMI became the principal sponsor and the registry was transferred to the section of Public Health and Surveillance of the Scientific Institute of Public Health (WIV - ISP). Since then, the WIV - ISP ensures the collection and the management of the data under the supervision of the board of the BCFR and the guidance of a scientific steering group. The board consists of a physician from each CF centre, a representative of the patients' association and the scientific collaborators of the WIV - ISP. The scientific steering group holds all stakeholders (representatives of the INAMI-RIZIV, patients' association, CF Centers and scientific collaborators of the WIV-ISP).

The Belgian CF Registry lies since 2006 within the framework of the CF convention described above. Participation in collection of data for the CF Registry is one of the obligatory tasks of the CF reference centres.

#### **OBJECTIVES OF THE CF PATIENT REGISTRY**

The aims of the CF patient Registry are to study epidemiological aspects of the disease among people with CF in Belgium and to provide a tool for the assessment of the management and quality of care for patients with CF. It also provides a database for scientific research to CF researchers both at the Scientific Institute and the CF reference centres. The registry also participates in activities organized by and contributes to the European CF Society Patient Registry (ECFSPR)<sup>[29]</sup> and other international projects.

#### CHAPTER 2: POPULATION AND METHODOLOGY

In this section, we describe the target population and methodology used. We briefly describe the data collection procedures and its evolution over the years and give the general structure of the database by listing some of its components. We look at the reporting procedures and use a flow chart to summarize the entire process.

#### STUDY POPULATION

The target population for the registry is people with CF who are cared for in Belgium. In 2014, there were 1230 patients included in the registry with about 1300 patients also registered as members of the patients' association. At the moment, the registry is estimated to have a coverage of more than 90% of all people with CF living in Belgium.

Prior to the registration, the physicians (at the accredited CF reference centre) provide each patient and/or their parents (or legal representative) information about the objectives of the registry. The patients are only included in the registry after signing an informed consent. They are identified by their national registry number at the centre level. This number is then pseudonymised into a unique code by a trusted third party (eHealth) before the data are transmitted to the registry. The patients' names are never transferred to the registry.

#### **DATA COLLECTION**

The clinical and demographic data is collected for all patients once each year by the treating physician from medical records and consists of more than 200 recorded items.

These data are divided into two sections:

- 1. The core data which contains demographic data, age of CF diagnosis and initial symptoms, genotype (mutations), sweat test and nasal transepithelial potential difference results including information on neonatal screening. This data is collected when the patient enters the registry and is updated if necessary during follow-up years.
- 2. Yearly follow-up sheets collect clinical data (height, weight), lung function (forced expiratory volume in one second (FEV<sub>1</sub>), forced vital capacity (FVC), forced expiratory flow 25-75% (FEF<sub>25-75</sub>)), complications that occurred or are still active during the registration year, microbiology results, treatments and medications taken as well as social data.

#### **SOFTWARE**

Until 2010, the data was collected via an Access based computer application on CD-Rom. A web-based application which worked with a trusted third party (eHealth) was then used for data collection 2011 until 2013. A new data collection, validation and analysis tool was developed in collaboration with the Healthdata.be platform and used for the first time for data collection 2014. The procedures for data collection, including the digital questionnaire is provided in the appendix.

#### **FEEDBACK**

Each of the seven reference centres (ten clinics) gets a copy of the national annual report. Since 2006, they receive a centre report based on the data from patients within the individual centre. Starting 2008, a feedback report has been provided with analyses that compare the results of each centre with data from the other centres so that the quality of care provided can be improved for points that score weaker in a centre, e.g. BMI. This method to optimize the care to the patients is called benchmarking. These analyses are corrected for some known factors such as patient age and gender with further corrections planned as more confounding data are collected by the registry, including socioeconomic data. A new-interactive reporting tool is under development and will be available on the website of Healthstat.be.

Physicians from the centres and researchers can submit research questions to the BCFR, and currently a number of research questions are being analysed. While some studies are still ongoing, abstracts have been presented at national or international conferences<sup>[63,64,65,66,67]</sup> and several articles have been published<sup>[68,69,70,71]</sup>.

#### INTERNATIONAL COLLABORATION

The Belgian CF Registry participates to the European CF patient Registry (ECFSPR). A subset of variables with similar definitions among several European countries is sent each year to the ECFSPR. Their annual reports, at-a-glance reports and publications are available on their website<sup>[29]</sup>.

The Belgian CF Registry also contributes data to the CFTR2 project (https://www.cftr2.org). The objective of the CFTR2 project is to define the disease-liability of CFTR variants by means of a multistage process which involves clinical (sweat chloride average), functional (expression in cell-based systems) and epidemiological (mutation analysis in healthy obligate heterozygotes<sup>(2)</sup>) steps<sup>[30,31]</sup>. The CFTR2 website provides information for patients, researchers, and the general public about specific variants in what is commonly referred to as the CF gene.

<sup>2</sup> An individual in a family who is proven to carry one copy of a recessive allele by having had affected progeny who inherited two copies of the mutant allele, one from each parent

#### **ETHICS AND PRIVACY**

The Belgian CF Registry has been approved by the Ethics Committees of all participating CF reference centres and clinics. The BCFR has also received the authorisation of the Privacy Commission for the treatment of coded data<sup>(3)</sup>.

#### **DATA FLOW**

The development and use of a new data collection tool on the HealthData. be <sup>(4)</sup> platform, has improved the data collection and validation procedures. It has also facilitated communication between the researchers and the data providers and enhanced the data verification and error correction. The first data via this system was collected for registry year 2014. The schema in figure 1 shows various stages from data entry and processing to reporting and publication.

#### Data collection (1)

A software called HD4DP (**H**ealth**D**ata for **D**ata **P**roviders) is installed locally by the data providers in the hospital. This software contains the electronic form to fill in all the requested data. Data that is present in the primary system of the hospital (e.g. in patient electronic files) can also be extracted and uploaded via a .csv file in HD4DP. This means that the system allows for manual data entry in the electronic form or via uploading a .csv file. In the same registration year, both modalities can be used. Several validation rules (ranges, logical checks...) have been implemented in the software at this stage of the process to minimise data entry errors.

#### • Data transfer (2)

When the registration forms have been completed, they are sent to the researchers in a secure way. The identifiers of the patients (national registry number) and the encrypted medical data are put in a digitally encrypted envelope. The envelope is sent to the "eHealthbox codage" of eHealth. The envelope is then opened. eHealth pseudonymises the identifiers and can not read the encrypted medical data. After pseudonymisation of the identifiers, all the data is put again in an encrypted envelop and sent via the eHealthbox to Healthdata.be at the WIV-ISP. The reopened envelope contains identifiers that are pseudonymised and medical data that will be decrypted to make

<sup>3</sup> FR:https://www.privacycommission.be/sites/privacycommission/files/documents/d%C3%A9lib%C3%A9ration\_SS\_084\_2010bis.pdf NL:https://www.privacycommission.be/sites/privacycommission/files/documents/beraadslaging\_AG\_084\_2010\_0.pdf

<sup>4</sup> The mission of Healthdata is to facilitate the data exchange between healthcare professionals and researchers to increase public health knowledge and to adjust health care policy in Belgium, with respect for the privacy of the patient, the healthcare professional and the medical confidentiality.

them readable for the researchers. The result is a set of registrations with all necessary medical information but unrecognizable patient data.

#### • Data monitoring and validation (3-4)

The data arrives to the WIV-ISP in a software called HD4RES (HealthData for Researchers) for validation by the researchers. The data is loaded into the system and each observation gets a time-stamp of the date and time when it was actually received. During validation, if a registration needs to be sent back to the data provider for verification or correction, the process of decoding and pseudony-misation is done in the opposite direction so that the data provider knows for which patient and which data needs to be checked. The newly corrected data is then sent again to the HD4RES as separate record via the described process above and will be assigned a new time stamp when loaded. No data sent by the data provider is changed or replaced including those initially sent with errors. It is stored as it was received.

#### • Data storage (5)

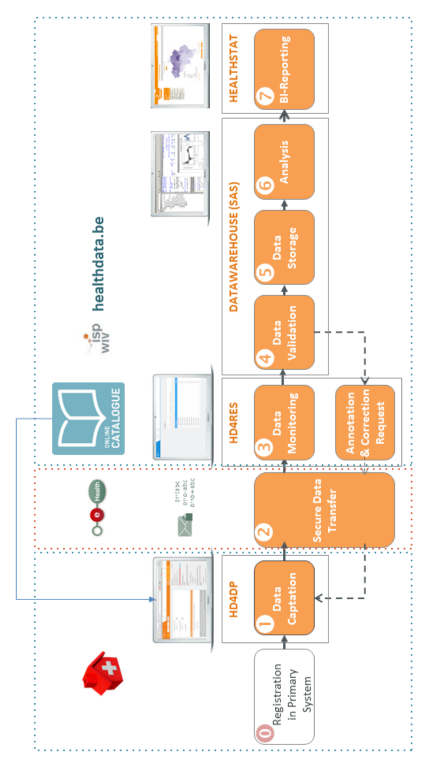
After validation all the data, including those that had errors and the new ones received from the data providers after correction, is stored in the data warehouse.

#### • Analysis (6)

The data used for the analysis is then populated based only on the valid records in the exploration environment. All old records that had been sent back for correction are excluded and only the new ones that replaced them are retained. Specific tables can then be populated for ad-hoc analyses and stored in this environment. An external analysis environment is also available for storage of the data tables sent to third parties such as the European CF Registry or used by other authorized researchers.

#### • Reporting (7)

A new reporting tool is under development and will be available on Healthstat.be. This environment will be divided in two sections: a public section with information accessible for all users and a secured section accessible only for authorized persons after identification by their electronic identification card (eID). In this section, benchmarking reports will be available.



Source: Healthdata.wiv-isp.be

CHAPTER 2: POPULATION AND METHODOLOGY

- 24
- 1. Patients without a confirmed diagnosis not included in the total number of CF patients starting 2012
- Patients with a revoked diagnosis not included in the total number of CF patients starting 2009
- 3. Patient's age at the last consultation
- 4. Patients without at least four filled-in clinical items, postulated alive or registered as deceased, and are not used in the analysis of clinical data
- 5. The new CF diagnoses are patients with the earliest diagnosis date from amongst the clinical diagnosis date, TEPD date, genotype date or the sweat test date done within the registry data year.
- 6. Prenatal diagnosis is considered without setting to zero the age at diagnosis allowing negative values since 2013. This explains the difference observed in the 2013 and 2014 values when compared to the previous years.

#### SUMMARY OF DATA REPORTS 2011 - 2014

Table 1 | A comparison of demographic data for years 2011 - 2014

	2011	2012	2013	2014
Number of CF patients	1171	1184	1186	1230
Number of CF patients with complete records	1161	1154	1153	1230
Number of CF patients without observation <sup>4</sup>	10	30	33	0
Number of CF patients with a transplant	134	141	142	151
Number of CF patients who were not seen	9	16	17	23
New CF diagnoses⁵	36	27	28	36
Number of patients without a confirmed diagnosis <sup>1</sup>	20	11	7	7
Number of patients with a revoked diagnosis <sup>2</sup>			7	2
Median patient age in years (range) <sup>3</sup>	19.7 (0.0 - 70.4)	20.3 (0.1 - 71.5)	20.7 (0.1 - 76.6)	21.3 (0.1 - 76.7)
Median patient age male (range) <sup>3</sup>	19.6 (0.2 - 64.2)	20.2 (0.1 - 65.2)	20.5 (0.1 - 66.2)	21.2 (0.2 - 67.3)
Median patient age female (range) <sup>3</sup>	19.8 (0.0 - 70.4)	20.4 (0.1 - 71.5)	20.7 (0.2 - 76.6)	21.5 (0.1 - 76.7)
Males (%)	51.8	51.4	52.0	52.0
Adults ≥ 18 years (%)	54.7	56.4	57.0	58.4
Median age at diagnosis (months) <sup>6</sup>	6.5	6.1	5.3	5.7
Age range at diagnosis (years)	0.0 - 65.0	-0.4 - 65.0	-0.4 - 74.2	-0.2 - 74.2
Median age at diagnosis, male (months)	6.5	6.2	5.3	5.9
Age range at diagnosis, male (years)	0.0 - 49.2	-0.4 - 46.9	-0.4 - 46.9	-0.2 - 59.5
Median age at diagnosis, female (months)	6.7	6.0	5.6	5.6
Age range at diagnosis, female (years)	0.0 - 65.0	-0.2 - 65.0	-0.2 - 74.2	-0.1 - 74.2
Median age at diagnosis new cases in years (range)	0.2 (0.0 - 49.2)	0.2 (-0.1 - 35.9)	0.7 (0.0 - 25.7)	0.2 (0.0 - 60.0)
Number of transplants performed	16	8	6	6
Total number of deaths reported	8	10	5	10
Median age at death in years (range)	27.7 (9.3 - 45.8)	30.2 (9.3 - 52.0)	24.9 (17.1 - 30.1)	37.4 (11.5 - 76.9)
Number of deaths among transplant patients	2	5	2	4
Overall mean FEV <sub>1</sub> % predicted <sup>[36,37]</sup>	77.8 (25.6)	77.6 (26.3)	76.0 (25.7)	76.0 (26.1)
In male	80.9 (26.3)	80.5 (26.5)	78.4 (25.2)	78.4 (25.3)
In female	74.5 (24.5)	74.4 (25.8)	73.3 (26.0)	73.3 (26.7)

#### **CHAPTER 3: DEMOGRAPHIC DATA**

In this section, the age at the end of the year by gender is presented in five-year categories. Information on the district of residence is provided, including maps of residence districts with estimated prevalence of CF based on the Belgian population in January 2014.

#### AGE ON DECEMBER 31 2014

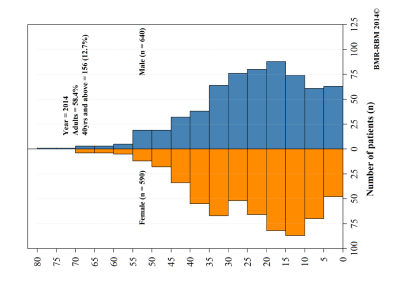
Table 2 Age on December 31 2014 by gender

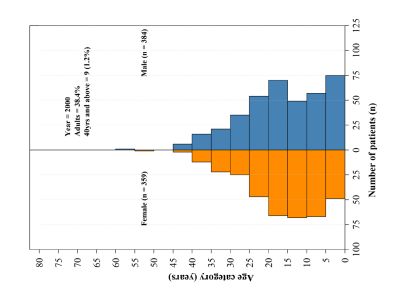
Age years	Males			Females			All Patients			
(on 31 Dec 2014)		cum n	cum %		cum n	cum %		cum n	cum %	
0 -< 5	56	56	8.8	54	54	9.2	110	110	8.9	
5 -< 10	56	112	17.5	74	128	21.7	130	240	19.5	
10 -< 15	82	194	30.3	72	200	33.9	154	394	32.0	
15 -< 20	104	298	46.6	65	265	44.9	169	563	45.8	
20 -< 25	81	379	59.2	72	337	57.1	153	716	58.2	
25 -< 30	58	437	68.3	70	407	69.0	128	844	68.6	
30 -< 35	65	502	78.4	66	473	80.2	131	975	79.3	
35 -< 40	56	558	87.2	39	512	86.8	95	1070	87.0	
40 -< 45	39	597	93.3	26	538	91.2	65	1135	92.3	
45 -< 50	17	614	95.9	21	559	94.7	38	1173	95.4	
≥ 50	26	640	100.0	31	590	100.0	57	1230	100.0	
Total	640			590			1230			

The overall median age on 31, December 2014 was 21.5 years; 21.3 for male and 21.7 for female patients respectively

Figure 2 | Age distribution by gender in 2000 (left) and 2014 (right)

The figures below compare the age distribution by gender for 2000 and 2014 data with an increase in the number of patients above age 40 years: 12.7% in 2014 compared to 1.2% in 2000.

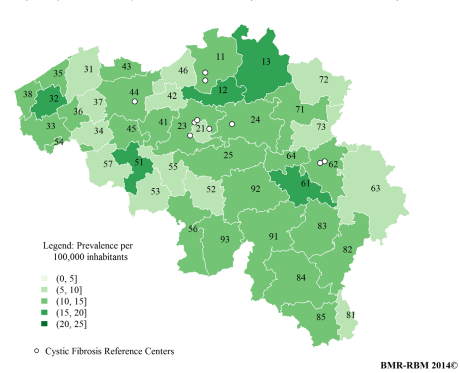




#### PREVALENCE OF CF PER DISTRICT OF RESIDENCE

The map below illustrates the prevalence of CF in each district of residence in Belgium based on the population as at the beginning of 2014. The numbers on the map refer to the district codes. A list with the district names and exact number of people with CF residing in each district is provided on page 31.

Figure 3 Prevalence per 100,000 inhabitants by district of residence in January 2014

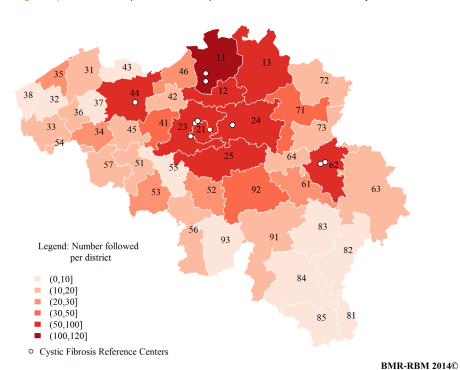


The figure 3 indicates that the highest prevalence of 19.9 in 100,000 is in Aat (Ath) District, code 51, with 17 patients in a population of 85,636 at the beginning of the year. District 55, Zinnik (Soignies) had the lowest prevalence of 5.3 in 100,000 given that there were 10 patients in a population of 187,338 during the reference period.

#### PATIENTS WITH CF UNDER FOLLOW-UP PER DISTRICT OF RESIDENCE

The figure 4 illustrates the number of CF patients resident in each district in Belgium at the beginning of 2014. The numbers on the map refer to the district codes. A list with the district names and exact number of people with CF residing in each district is provided on page 31.

Figure 4 Number of patients with CF per district of residence in January 2014



In the figure above, we see more patients concentrated in the central and northern sides of the country. Due to its populous nature though, Antwerp district (11) with 109 patients in a population of about 1.02 million had a prevalence of 10.6 in 100,000 inhabitants, while there were only 4 patients living in district 81 (Aarlen, Arlon) in a population of 60,323 inhabitants.

#### DISTRICT OF RESIDENCE

The table below shows the number of patients in the registry according to their district of residence. Most of the patients reside in Belgium. There are however some patients who reside in neighbouring countries.

Table 3 District of residence

	District/Arrondissement	n	%		District/Arrondissement	n	%
11	Antwerpen	109	8.9	61	Huy	21	1.7
12	Mechelen	62	5.0	62	Liège	76	6.2
13	Turnhout	79	6.4	63	Verviers	19	1.5
21	Brussel Hoofdstedelijk Gewest Région Bruxelles Capitale	89	7.2	64	Waremme	11	0.9
23	Halle-Vilvoorde	77	6.3	71	Hasselt	47	3.8
24	Leuven	52	4.2	72	Maaseik	16	1.3
25	Nivelles	54	4.4	73	Tongeren	11	0.9
31	Brugge	19	1.5	81	Arlon	4	0.3
32	Diksmuide	10	0.8	82	Bastogne	5	0.4
33	leper	11	0.9	83	Marche-en-Famenne	7	0.6
34	Kortrijk	27	2.2	84	Neufchâteau	9	0.7
35	Oostende	21	1.7	85	Virton	7	0.6
36	Roeselare	17	1.4	91	Dinant	11	0.9
37	Tielt	9	0.7	92	Namur	35	2.8
38	Veurne	9	0.7	93	Philippeville	7	0.6
41	Aalst	32	2.6				
42	Dendermonde	18	1.5				
43	Eeklo	9	0.7				
44	Gent	66	5.4				
45	Oudenaarde	15	1.2		Subtotal	1116	
46	Sint-Niklaas	24	1.9		Foreign country	14	1.1
51	Ath	17	1.4		Missing	-	-
52	Charleroi	29	2.4		Total	1230	
53	Mons	22	1.8				
54	Mouscron	11	0.9				
55	Soignies	10	0.8				
56	Thuin	20	1.6				
57	Tournai	12	1.0				

#### CHAPTER 4: DIAGNOSIS

In this section, we present the symptoms and clinical reasons suggesting a CF diagnosis. We also present a table of the procedure used for CF diagnosis which classifies the people with CF into two groups depending on whether or not they meet the **conditions set for inclusion** into the European CF Society Patient Registry for (ECFSPR). Also presented is the age at diagnosis and the mutations found after genotyping.

#### SYMPTOMS AND CLINICAL REASONS SUGGESTING CF

In a patient with suggestive symptoms, a family history of CF or a positive neonatal screening test, the diagnosis of CF is confirmed by an abnormal sweat test (chloride > 60 mmol/L) and/or the identification of two mutations in the *CFTR* gene. Even though Belgium has no existing national neonatal screening program implemented, some children were screened for CF. Most patients present with a combination of respiratory and/or gastrointestinal symptoms. Chronic cough, recurrent chest infections, chronic sinusitis are the most common presenting respiratory signs. Common gastrointestinal symptoms include meconium ileus (obstruction of the bowel with sticky secretions in the newborn infant), chronic diarrhea and failure to thrive due to malabsorption. Less frequently, salt loss, jaundice or a rectal prolapse are the first diagnostic signs. In some cases, the diagnosis of CF is delayed until adulthood. Most of these patients are expected to have had a milder clinical course, or to present with atypical symptoms, such as infertility.

The diagnostic signs or clinical presentation are illustrated in table 4. In the Belgian CF Registry, it is possible to report more than one diagnosis sign or symptom for the same patient. Over the years, the commonest clinical presentation of CF remains acute or recurrent respiratory problems. Other common features on presentation were failure to thrive, chronic diarrhea/steatorrhea and meconium ileus. About 17.0% of the patients were diagnosed via neonatal screening test.

Table 4 Symptoms and clinical reasons for CF diagnosis

			Newly diagnosed				
	Data 1998 – 2014		2013		2014		
	n	%	n	n %		%	
Acute or recurrent respiratory problems	509	42.7	12	46.2	12	33.3	
Failure to thrive	289	24.2	7	26.9	6	16.7	
Chronic diarrhea/steatorrhea/ malabsorption	240	20.1	5	19.2	3	8.3	
Neonatal screening test	201	16.8	6	23.1	10	27.8	
Meconium ileus	171	14.3	3	11.5	5	13.9	
Family history	125	10.5	3	11.5	5	13.9	
Nasal polyposis / chronic sinusitis	53	4.4	3	11.5	1	2.8	
Rectal prolapse	32	2.7	0	0.0	0	0.0	
Intestinal obstruction (other than meconium ileus)	26	2.2	0	0.0	1	2.8	
Prenatal diagnosis	36	3.0	1	3.8	2	5.6	
Dehydration / electrolyte imbalance	19	1.6	1	3.8	1	2.8	
Neonatal jaundice / Prolonged icterus	2	0.2	1	3.8	0	0.0	
Infertility	12	1.0	0	0.0	0	0.0	
Diagnosis other	96	8.0	1	3.8	6	16.7	
*No diagnosis reasons given	37	3.1	2		0		

<sup>\*</sup> The overall percentages are based on 1193, 37 patients did not have information on any of the above reasons given in the 2014 data and were excluded from the calculations.

There were 28 newly diagnosed in 2013, percentages are based on 26 patients. There were 36 newly diagnosed in 2014, percentages are based on 36 patients.

Note: Reasons for diagnosis are not mutually exclusive.

## **DOCUMENTATION OF CF DIAGNOSIS**

Table 5 | Documentation of CF at diagnosis

Procedure		nta - 2014	Newly diagnosed Patients 2014	
	n	%	n	%
Patients meeting inclusion criteria for the European CF S	Society Pa	tient Regi	stry	
Clinical symptoms and/or family history, sweat test and genotyping	715	58.1	20	55.6
Clinical symptoms and/or family history and sweat test	33	2.7	1	2.8
Clinical symptoms and/or family history and genotyping	214	17.4	5	13.9
Neonatal screening test, sweat test and genotyping	115	9.3	6	16.7
Clinical symptoms and/or family history, neonatal screening test, sweat test and genotyping	55	4.5	2	5.6
Sweat test and genotyping	28	2.3	٠	
Clinical symptoms and/or family history, neonatal screening test and genotyping	14	1.1	1	2.8
Subtotal	1174	95.4	35	97.2
Patients not meeting inclusion criteria for the European	CF Societ	y Patient I	Registry	
Clinical symptoms and/or family history only	14	1.1		
Genotyping only	25	2.0		
Neonatal screening test and genotyping	16	1.3	1	2.8
Clinical symptoms and/or family history and neonatal screening test	1	0.1	·	
Subtotal	56	4.6	1	2.8
Total	1230	100.0	36	100.0

Important Information: The data in the table above refers to documentation of CF at diagnosis. The clinical diagnosis was considered if at least one of a set of symptoms was identified (see page 34; excluding neonatal screening); the sweat chloride had to be greater than 60 mmol/L while genotyping was considered confirmatory if two CF mutations were reported.

According to the European CF Society Patient Registry (ECFSPR), for a patient to be included, he/she must meet at least one of the following three criteria listed on the next page. We have used those criteria to explore the documentation of the diagnosis of the patients in our registry. These data are presented in the table above.

#### Inclusion criteria for patients into the European CF Society Patient Registry (ECFSPR)

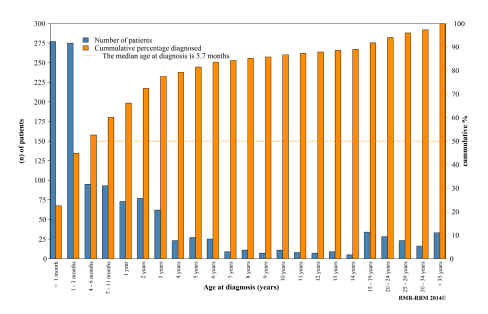
For a patient to be included in the ECFSPR, at least one of the following three criterions must be met:

- 1) two sweat tests returning results of sweat chloride > 60 mmol/L
- 2) one sweat test with chloride > 60 mmol/L and DNA Analysis/Genotyping done where two disease causing CF mutations are identified
- 3) if sweat chloride value is  $\leq$  60 mmol/L then at least two of the conditions below should be fulfilled:
  - i. A DNA Analysis/Genotyping where two disease causing CF mutations are identified ii. A Transepithelial (Nasal) Potential Difference value Consistent with a diagnosis of CF iii. Clinical Presentation at diagnosis Where typical features of CF are identified

#### **AGE AT DIAGNOSIS**

The figure gives the cumulative percentage of age at diagnosis for data 2014. The median age at diagnosis was 5.7 months; 5.9 months for male and 5.6 months for female patients respectively. At the age of 18 years 91.6% of the patients were diagnosed. The age range at diagnosis is from two months before birth, due to prenatal diagnosis, to 74.2 years. The median age at diagnosis was 3.1 months for the F508del homozygous patients, 8.4 months for the F508del heterozygous while for patients with other mutations it was 20.4 months.

Figure 5 | Age at Diagnosis



The median age at diagnosis for the 36 newly diagnosed patients in 2014 was 2.5 months; 4.1 months for male and 1.9 months for female patients respectively.

# **GENOTYPE**

All the 1230 patients have undergone a genetic analysis. However, 53 patients (4.3%) had at least one non-identified mutation. Almost half (45.4%) were homozygote for F508del (table 6) and 85.4% of the patients had this mutation on at least one of their alleles (table 7).

Table 6 General mutation pairs

Mutation pair	n	%	cumulative %
F508del Homozygous	559	45.4	45.4
F508del Heterozygous	460	37.4	82.8
F508delNI	31	2.5	85.4
OTHEROTHER	158	12.8	98.2
OTHERNI	6	0.5	98.7
NINI	16	1.3	100.0
Total	1230		

NI = Not Identified

Table 7 | Number and proportion of patients by CF allele or mutation

Marketten	Pati	ents	Alle	eles	Marketiere	Pati	ents	Alle	eles
Mutation	n	%	n	%	<ul> <li>Mutation</li> </ul>	n	%	n	%
F508del	1049	85.4	1624	66.1	E60X	6	0.5	6	0.2
G542X	65	5.3	72	2.9	G178R	6	0.5	6	0.2
N1303K	60	4.9	65	2.6	R334W	6	0.5	6	0.2
3272-26A->G	41	3.3	41	1.7	Y1092X	6	0.5	6	0.2
1717-1G->A	38	3.1	38	1.5	G85E	5	0.4	5	0.2
S1251N	32	2.6	32	1.3	G970R	5	0.4	5	0.2
A455E	28	2.3	28	1.1	L165S	5	0.4	5	0.2
2789+5G->A	25	2.0	25	1.0	Q493X	5	0.4	5	0.2
L927P	24	2.0	25	1.0	4218insT	4	0.3	4	0.2
R117H	23	1.9	24	1.0	621+1G->T	4	0.3	4	0.2
3849+10kbC->T	18	1.5	18	0.7	CFTRdele2,3	4	0.3	5	0.2
2183AA->G	17	1.4	17	0.7	G551D	4	0.3	5	0.2
R553X	17	1.4	17	0.7	L227R	4	0.3	7	0.3
W1282X	17	1.4	17	0.7	L997F	4	0.3	6	0.2
R1162X	11	0.9	15	0.6	Q1313X	4	0.3	4	0.2
I507del	10	0.8	10	0.4	R347H	4	0.3	4	0.2
5T	9	0.7	9	0.4					
306insA	8	0.7	8	0.3	Others	184	15.0	193	<i>7.8</i>
3659delC	8	0.7	8	0.3	Not identified	53	4.3	69	2.8
D1152H	8	0.7	8	0.3	Subtotal			2460	
W401X	7	0.6	7	0.3	Missing	-	-	-	-
394delTT	6	0.5	7	0.3	Total			2460	

 $The \ mutations \ detected \ in \ less \ than \ four \ patients \ were \ summarized \ into \ the \ "others" \ category \ for \ this \ purpose.$ 

# CHAPTER 5: ANTHROPOMETRY (HEIGHT, WEIGHT AND BMI)

Persons with CF are known to be prone to nutritional deficiencies. Because of thick mucus, the pancreas is unable to produce and/or carry digestive enzymes to the gut. This leads to poor absorption of proteins, fats and fat soluble vitamins resulting in poor weight gain and growth. Nutritional care is of great importance for patients with CF. Maintaining or achieving a better nutritional status has a positive impact on lung function. For this reason, close follow-up of height, weight and BMI is standard practice in all CF care centers.

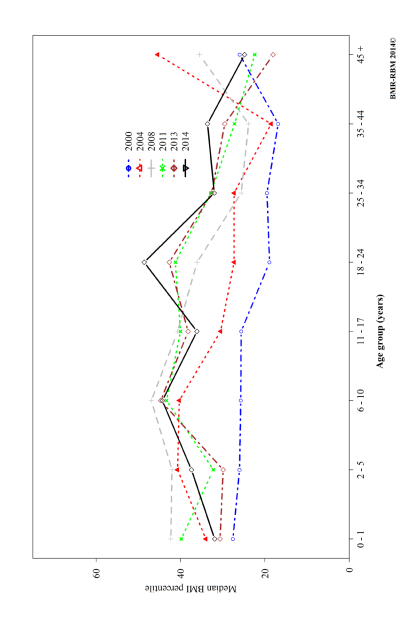
We present classifications of the patients into BMI, height and weight percentiles, compared to their healthy peers. The analysis of the height, weight and BMI z-scores is based on CDC<sup>[32]</sup> and Cachera<sup>[33]</sup> reference equations. An evolution of BMI and height data from selected years is also presented by age category.

In this section, data from 151 patients with a transplant (71 male, 80 female) were excluded from the analysis.

Figure 6 | Median BMI percentile by age group and year

## BMI PERCENTILES USING CACHERA REFERENCE VALUES

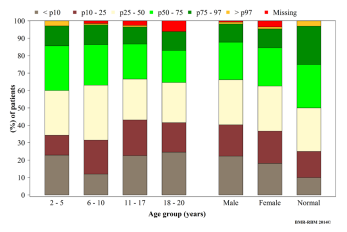
Cachera equations cover BMI z-scores for ages 0.0 - 58.0 years for male (56.0 female) patients. In 2014 data from 1001 patients was analysed. The figure below shows a general trend for better median BMI over the years with lines shifting upwards until 2008. After this, there has been a further improvement of the median BMI in the adults, and stabilization in age categories 6-17 years. No further improvement was found from 2008 onwards in the patients younger than 5 years.



### BMI PERCENTILES USING THE CDC GROWTH CHARTS

CDC growth charts cover BMI for ages 2.0 – 20.0 years. The trend depicted is quite similar to the Cachera references above. In 2014 data from 536 patients was analysed. The figure below displays the proportion in each percentile category.

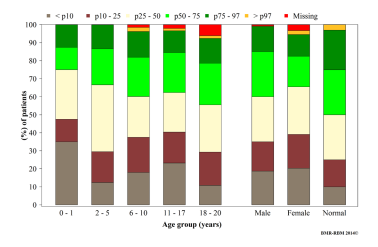
Figure 7 | CDC BMI percentiles by age



#### HEIGHT PERCENTILES USING THE CDC GROWTH CHARTS

The CDC growth charts cover height from 0.0 - 20.0 years. No extrapolation was made to get values for patients older than 20.0 years. In 2014 data from 576 patients was analysed. The figure below indicates the proportion in each percentile category.

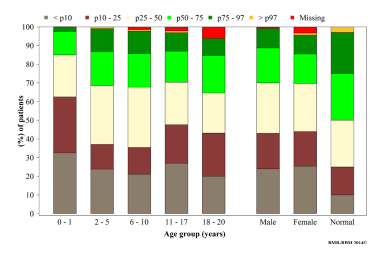
Figure 8 | CDC height percentiles by age



## WEIGHT PERCENTILES USING THE CDC GROWTH CHARTS

The CDC growth charts cover weight from 0.0-20.0 years. In 2014 data from 576 patients was analysed. The figure below indicates the proportion in each percentile category. A higher proportion with weight for age below the 10th percentile is seen in patients up to one year: this proportion declines in the years thereafter and increases again in pre-teenage children over the age 11 years. This pattern has been observed in the registry over the years.

Figure 9 | CDC Weight percentiles by age



# **CHAPTER 6: SPIROMETRY (LUNG FUNCTION)**

The forced expiratory volume in 1 second ( $FEV_1$ ) is the amount of air that a person is able to expire forcefully in one second, following full inspiration. It is expressed as a percentage of the predicted value for a reference population with same age, gender and height.

The FEV<sub>1</sub> % predicted is a clinical parameter to monitor lung function impairment. The FEV<sub>1</sub> partly determines the prognosis<sup>[34]</sup>. Because most patients with CF develop progressive pulmonary disease, measures of pulmonary involvement, in particular FEV<sub>1</sub>, are used to follow up the lung disease. However, considerable heterogeneity exists in prognosis and severity, even among patients of the same genotype<sup>[35]</sup>.

FEV<sub>1</sub> % predicted values are divided in four classes for the CF population corresponding to different degrees of lung function impairment: normal lung function ( $\geq$  90%), mild (70-89%), moderate (40-69%) and severe (< 40%) impairment.

Since lung function measurements below the age of 6 years are not reliable, data from those patients (**74 male, 71 female**) was excluded from the lung function analysis. Those with a transplant (**71 male, 80 female**) were also excluded. The values obtained at the last consultation of the year, pre- or post-bronchodilator, were analysed. An evolution of lung function from selected years by age category is also presented.

# PERCENTAGE OF PREDICTED FEV<sub>1</sub>

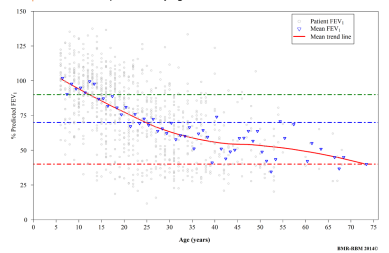
Wang's equations<sup>[36]</sup> were used for male, 6 - 17 years and female patients 6 - 15 years, while Hankinson's<sup>[37]</sup> were used for predictions for the male 18 years and above and female patients from 16 years onwards.

In 2014 data from 891 patients was analysed. The overall mean FEV<sub>1</sub> % predicted is 76.0 (SD = 26.1). The mean FEV<sub>1</sub> % predicted was 78.4 % (SD = 25.3) and 73.3 % (SD=26.7) respectively for 472 male and 419 female patients. The mean FEV<sub>1</sub> % predicted was 92.1 % (SD = 20.5) and 65.3 % (SD=23.9) respectively for 355 children and 536 adult patients.

Amongst the 393 F508del homozygous patients, the means were 75.3% (SD = 26.0) and 71.7% (SD = 26.2) respectively for the 209 male and 184 female patients. The means were 90.7% (SD = 21.5) and 62.8% (SD = 22.8) respectively for the 153 children and 240 adults homozygous for the F508del mutation.

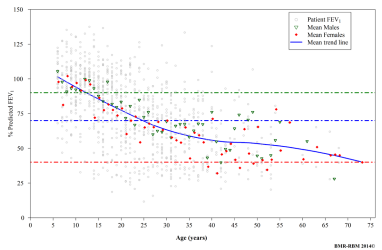
The figure below shows a scatter plot of the  $FEV_1$  % predicted with the means calculated at yearly intervals. It shows on average declining values with age, with a steeper slope up to about 35 years of age when it levels off.

Figure 10 | Mean FEV<sub>1</sub> % predicted by age



In the figure below, the scatter plot is plotted with means according to gender.

Figure 11 | Mean FEV<sub>1</sub> % predicted by age and gender



## FEV₁ CATEGORIES BY AGE GROUP

FEV<sub>1</sub>% predicted values were divided in four classes for the CF population corresponding to different degrees of lung function impairment: normal lung function ( $\geq$ 90%), mild (70-89%), moderate (40-69%) and severe (<40%) lung function impairment. The table below shows the classification for children and adults based on the data collected in 2014.

Table 8 Proportions in each FEV<sub>1</sub> severity category for children and adults

Group		dren years)		ults years)	То	tal
	n	%	n	%	n	%
Normal : ≥ 90% predicted	222	60.7	87	15.3	309	33.1
Mild: 70% - 89% predicted	84	23.0	151	26.6	235	25.2
Moderate: 40% - 69% predicted	41	11.2	195	34.3	236	25.3
Severe : < 40% predicted	8	2.2	103	18.1	111	11.9
Missing	11	3.0	32	5.6	43	4.6
Subtotal	366		568		934	
Transplants	1		150		151	
< 6 years	145		-		145	
Total	512		718		1230	

The FEV $_1$  was 70.0% of predicted or higher in 58.3% of the patients: - in 83.7% of the children (6 – 17 years) and 41.9% of the adults (18 years and above). About 5.0% had missing FEV $_1$  data.

The figure 12 represents the lung function severity groups for children and adults in 2014 using the Wang – Hankinson equations. When missing data is excluded, 12.5% of the patients had  $FEV_1$  below 40%, 34.7% had  $FEV_1$  of at least 90% in 2014. This is comparable to 2013 where 34.0% of the patients also had  $FEV_1$  of at least 90%; 60.4% of the children and 14.7% of the adults.

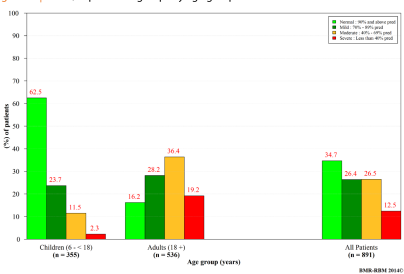
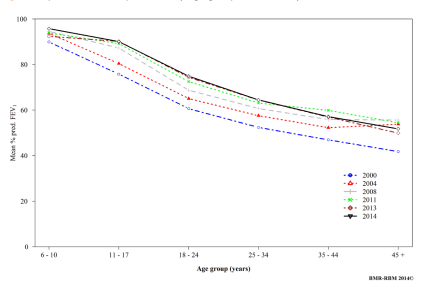


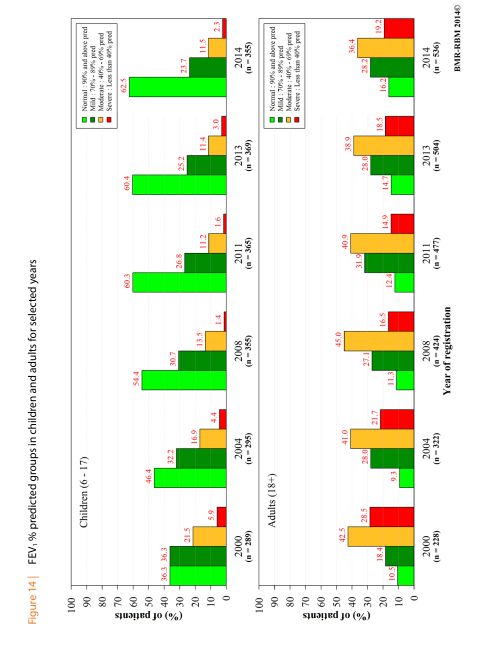
Figure 12 | FEV<sub>1</sub> % predicted groups by age group

In the figure below, the mean  $\mbox{FEV}_1$  % predicted calculated cross-sectional shows improving lung function over time in all age categories.

Figure 13 Mean FEV<sub>1</sub> % predicted by age group for selected years



In the following figure, a comparison of the proportion in each severity group over selected years is given. Missing data is excluded. There has been a general increase in the number of children with normal lung function and a reduction in the proportion of adults with severe lung function impairment, except in the two last years.



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# CHAPTER 7: MICROBIOLOGY

Decreased mucus clearance and impaired bacterial killing leads to inflammation and infection and are responsible for progressive lung damage.

Bacterial colonisation occurs very early in the natural history of the disease. In children common bacteria such as *Staphylococcus aureus* and *Haemophilus influenzae* infect the lungs<sup>[40]</sup>. Infections by *Pseudomonas aeruginosa* and sometimes *Burkholderia cepacia* complex and other gramnegative pathogens occur at a later age. The airways of patients with CF may also be chronically colonized by fungi like *Aspergillus fumigatus*<sup>[41]</sup>.

One of the main goals of CF care is to prevent or postpone by all possible means infections with the above-named pathogens and to reduce the risk of chronic infection which increases respiratory morbidity and treatment burden.

In this section, we present the annual prevalence of recorded pathogens including the prevalence of chronic infections. **Data from the transplant patients was excluded** from the respiratory microbiology analysis.

The analysis presented is based on the pathogens found at least once during the whole year (annual prevalence). The prevalence is also compared over a selected period in children and adults or by age category.

#### ANNUAL PREVALENCE OF ISOLATED PATHOGENS

During the year 2014, 1037 (96.1%) of the 1079 non-transplant patients had at least one culture done. The largest proportion (82.4%) had at least four exploitable months during the year. Sputum samples were done in 807, throat swabs in 405 while 38 patients had a broncho-alveolar lavage.

The prevalence shown in table 9 refers to pathogens ever found during the year.

Table 9 | Isolated pathogens 2011 - 2014

	20	11	20	12	20	13	20	14
Methicillin Sensitive Staphylococcus aureus (MSSA)	598	58.7	612	61.8	630	62.4	631	60.8
Haemophilus influenzae	276	27.1	294	29.7	288	28.5	279	26.9
Pseudomonas aeruginosa	426	41.8	420	42.4	426	42.2	428	41.3
Methicillin Resistant Staphylococcus aureus (MRSA)	97	9.5	87	8.8	66	6.5	77	7.4
Stenotrophomonas maltophilia	91	8.9	118	11.9	119	11.8	128	12.3
Achromobacter xylosoxidans	91	8.9	106	10.7	106	10.5	104	10.0
Burkholderia cepacia complex	37	3.6	40	4.0	45	4.5	38	3.7
Aspergillus	329	32.3	346	34.9	331	32.8	364	35.1
Scedosporium spp.	8	0.8	9	0.9	4	0.4	3	0.3
Atypical / Non – tuberculous mycobacteria (NTM)	8	0.8	11	1.1	8	0.8	11	1.1
Other pathogens	160	15.7	147	14.8	168	16.7	224	21.6

Percentages are based on 1019, 991, 1009 and 1037 patients with a culture respectively for the years 2011 through 2014

While 27.0% of the children (n = 136) and 54.7% of the adults (n = 292) had a *Pseudomonas aeruginosa*, 6.2% (n = 31) and 8.6% (n = 46) respectively had a MRSA infection. *Burkholderia cepacia* complex infection was found in 8 (1.6%) children and 30 (5.6%) adults. The prevalence of *Stenotrophomonas maltophilia* continues to rise from 8.9% in 2011 to 12.3% in 2014, a significant increase (P = 0.0121). The increase in 'Other pathogens' may need a more sensitive classification to detect if a particular new pathogen is responsible.

Table 10 | Chronic infections 2011 - 2014

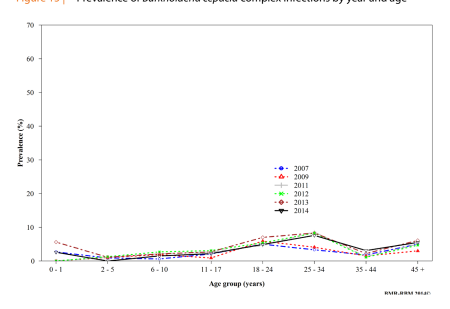
	2011		2012		2013		20	14
	n	%	n	%	n	%	n	%
Chronic Pseudomonas aeruginosa	287	28.2	276	27.9	296	29.3	294	28.4
Chronic Burkholderia cepacia complex	23	2.3	27	2.7	29	2.9	34	3.3
Chronic Stenotrophomonas maltophilia	30	2.9	27	2.7	37	3.7	41	4.0
Chronic Achromobacter xylosoxidans	39	3.8	54	5.4	64	6.3	69	6.7
Chronic MRSA	45	4.4	51	5.1	45	4.5	45	4.3

Percentages are based on 1019, 991, 1009 and 1037 patients with a culture respectively for the years 2011 through 2014

About 11.3% of the children (n = 57) and 44.4% of the adults (n = 294) had chronic *Pseudomonas aeruginosa* infection.

The figures below show the annual prevalence of infections over selected periods. Since 2007, the prevalence is based on any positive culture for a given pathogen among those collected during the year. Before 2007, the prevalence was based on a positive culture at the last consultation of the year. Therefore only data from 2007 onwards is presented.

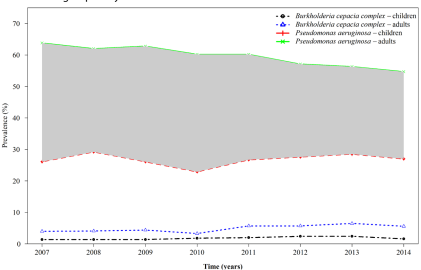
Figure 15 | Prevalence of *Burkholderia cepacia* complex infections by year and age



70 - 60 - 50 - 2007 - 2009 - 2012 - 2013 - 2014 - 2

Figure 16 Prevalence of *Pseudomonas aeruginosa* infections by year and age

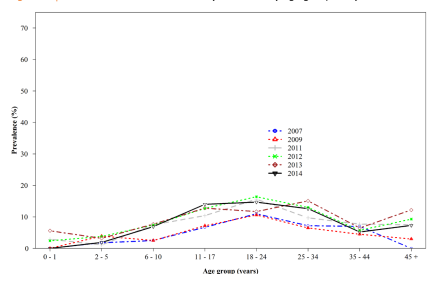
Figure 17 | Prevalence of *Pseudomonas aeruginosa* and *Burkholderia cepacia* complex by age group and year



<u>Note</u>: The shaded area is the difference in *Pseudomonas aeruginosa* prevalence between children and adults.

There has been an increase in the importance of some pathogens such as *Achromobacter xylosoxidans*, figure 18, whose prevalence has been rising over the years.

Figure 18 | Prevalence of Achromobacter xylosoxidans by age group and year



## **CHAPTER 8: COMPLICATIONS**

The defective chloride channel in CF causes a range of disturbances within the human body. Chloride channels are needed to regulate fluid exchanges at the surface of the epithelial cells. In CF, the transport through the cell wall of chloride, other ions and water are disturbed. CF affects the respiratory<sup>[42]</sup>, digestive<sup>[43,44]</sup>, and reproductive<sup>[45,46]</sup>, systems with variable degrees of severity. Complications in CF are mainly found in organs where mucus linings are needed (airways, intestines) and glands which need fluid to excrete their substances (pancreas, testicles…).

In this section we cover the most frequent complications recorded during the year by age category. The overall prevalence is compared to the prevalence reported in previous years.

**Data from transplant patients was excluded** from the analysis of complications.

### RESPIRATORY COMPLICATIONS

**Allergic bronchopulmonary aspergillosis (ABPA)**: ABPA is an allergic reaction to *Aspergillus fumigatus*, a fungus that colonizes the airway of people with CF. Diagnosis is not always obvious as many symptoms of ABPA (cough, wheezing, shortness of breath, and decline in lung function) are common symptoms of CF lung disease. Diagnosis relies on a combination of the clinical picture, blood tests, lung function and imaging.

**Nasal polyps**: In patient with CF, chronic infections of the upper airways (chronic sinusitis) may cause the formation of nasal polyps which are mucosal overgrowths. They are responsible for nasal obstruction.

**Haemoptysis**: When the damage within the bronchi reaches a blood vessel the patient with CF is coughing up blood. Haemoptysis is mild in most cases, but sometimes the bleeding is so massive (> 250 cc/24h) that a therapeutic embolization of the bleeding vessel can be indicated.

**A pneumothorax** occurs when air reaches the vacuum of the pleural space, mainly caused by destruction of the alveoli causing the leakage of air. This complication is more common in the adults who have more advanced lung disease<sup>[47,49]</sup>

Table 11 | Prevalence of respiratory complications

Complication	2011		2012		2013		2014	
Complication	n	%	n	%	n	%	n	%
Allergic bronchopulmonary aspergillosis (ABPA)	63	6.1	40	3.9	52	5.1	72	6.7
Pneumothorax	2	0.2	3	0.3	2	0.2	3	0.3
Nasal polyps	85	8.3	108	10.6	155	15.2	205	19.0
Massive haemoptysis	10	1.0	6	0.6	11	1.1	15	1.4
Massive haemoptysis requiring embolization	7	0.7	2	0.2	5	0.5	5	0.5

Percentages are based on 1030, 1018, 1017 and 1079 non transplant patients respectively for the years 2011 through 2014

ABPA is the most frequent major respiratory complication. In 2014 ABPA was reported in 29 (5.7%) children and 43 (7.6%) adults.

## **GASTRO-INTESTINAL AND ENDOCRINE COMPLICATIONS**

**Exocrine pancreatic insufficiency**: Pancreatic enzymes are needed to digest fat and proteins. Pancreatic insufficiency is the inability of the pancreas to produce and transport enough enzymes to the duodenum to digest fat and proteins resulting in malabsorption with steatorhea (fatty stools), malnutrition and a deficiency in fat-soluble vitamins (ADEK). A small proportion of patients with CF remain pancreatic sufficient (10-15%).

**Gastro-oesophageal reflux** is a condition in which contents of the stomach or small intestine repeatedly move back up into the oesophagus. When repeated it causes oesophagitis and can lead to malnutrition but also respiratory infections and it may worsen the respiratory function.

**Distal intestinal Obstruction syndrome (DIOS)**: The intestinal cells with defective chloride channels produces thick intestinal mucus which in combination with stool and undigested food residue can cause intestinal obstruction. DIOS is especially found in the terminal ileum and caecum. It can cause acute abdominal pain and, if left untreated, it can progress to a complete bowel obstruction with vomiting. DIOS usually responds to medical treatment, but in a few cases a surgical intervention may be required.

**Liver disease**: All patients with CF present a defective CFTR protein in their biliary tract. Some patients develop liver disease leading to cirrhosis (replacement of the liver tissue by fibrosis) and others do not. Cirrhosis may be complicated by portal hypertension and in rare cases by hepatic insufficiency. In some cases a liver transplant is required.

**CF-related diabetes (CFRD)**: Insulin is produced in the endocrine part of the pancreas. It is a hormone which maintains the balance of sugar in blood. The malfunctioning of the endocrine part of the pancreas by fibrosis leads to an insufficiency of the secretion of insulin leading to diabetes.

Table 12 | Prevalence of gastro-intestinal and endocrine complications

Compliantion	20	11	2012		2013		2014	
Complication								%
Exocrine pancreatic insufficiency	855	83.0	830	81.5	838	82.4	867	80.4
Acute pancreatitis	11	1.1	6	0.6	8	0.8	14	1.3
CF related diabetes (CFRD)	143	13.9	128	12.6	159	15.6	176	16.3
Impaired Glucose Tolerance (IGT)	68	6.6	59	5.8	66	6.5	77	7.1
Gastro-oesophageal reflux	179	17.4	198	19.4	188	18.5	265	24.6
Cirrhosis with portal hypertension	32	3.1	33	3.2	43	4.2	43	4.0
Gallstones	28	2.7	32	3.1	21	2.1	36	3.3
Intestinal obstruction (surgery)	5	0.5	4	0.4	5	0.5	2	0.2
Intestinal obstruction (no surgery)	55	5.3	74	7.3	52	5.1	37	3.4
Gastroparesis	2	0.2	7	0.7	3	0.3	2	0.2
Clostridium infection (treatment needed)	10	1.0	4	0.4	5	0.5	15	1.4

Percentages are based on 1030, 1018, 1017 and 1079 non transplant patients respectively for the years 2011 through 2014

In non-transplant patients, the data shows that 427 (83.6%) of the children and 440 (77.5%) of the adults are pancreatic insufficient. CFRD was reported in 18 (3.5%) of the children and 158 (27.8%) of the adults.

### MISCELLANEOUS COMPLICATIONS

**Reproductive system complications**: Most men (95 – 99%) with CF are infertile because of congenital bilateral absence of the vas deferens (which allows the transport of the spermatozoids). However, as the production of spermatozoids is being preserved, techniques of assisted procreation are possible. Although women with CF may be less fertile than other women, it is possible for them to conceive and to have successful pregnancies. Those pregnancies require a higher surveillance.

**Osteopenia and osteoporosis**: Osteopenia and osteoporosis which are the result of a progressive loss of the bone mass are more frequent and earlier in CF due to various risk factors like malabsorption of vitamins D, use of glucocorticoids, chronic inflammation...

**CF-related arthritis/arthropathy**: Patients with advanced lung disease sometimes develop painful inflammation of joints. The exact cause is not known and regression of joint symptoms is usually seen when respiratory disease is stabilized.

**Psychiatric disease**: It is difficult to define and quantify the extend of psychological repercussions in CF. In this report, the analysis is based on patients who visited or were treated by a specialist (psychiatrist or neurologist).

Table 13 Other complications reported

Complianting	20	11	20	12	2013		2014	
Complication	n	%	n	%	n	%	n	%
CF-related arthritis / arthropathy	98	9.5	72	7.1	102	10.0	23	2.1
Cancer	3	0.3	2	0.2	3	0.3	4	0.4
Surgery	98	9.5	66	6.5	72	7.1	97	9.0
General anaesthesia	86	8.3	73	7.2	102	10.0	109	10.1
Psychiatric disease	23	2.2	24	2.4	36	3.5	41	3.8
Osteopenia	107	10.4	112	11.0	117	11.5	113	10.5
Osteoporosis	24	2.3	24	2.4	23	2.3	21	1.9
Hypertension requiring treatment	16	1.6	14	1.4	31	3.0	14	1.3
Others	126	12.2	143	14.0	161	15.8	170	15.8

Percentages are based on 1030, 1018, 1017 and 1079 non transplant patients respectively for the years 2011 through 2014

\*Since 2011, data on the Osteopenia (the lowest z-score on Dual X-ray absorptiometry (DXA) between -1.0 and -2.5) and Osteoporosis (the lowest z-score on DXA < -2.5) complications are recorded separately.

# CHAPTER 9: THERAPY, MEDICATION AND HOSPITALIZATION

Till today no definitive cure for CF exists. The problems of all CF patients are related to defective epithelial function with impaired production of mucus and fluids leading to complications described in the previous section. Treatment of the disease is therefore mostly based on preventing or reducing symptoms in order to avoid complications or to stabilize them. Thanks to those preventive and symptomatic treatments, a steady improvement of the health status is obtained together with a better life expectancy.

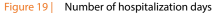
In this section, the most common treatments and therapy used in 2014 are presented. An evolution of the proportion of patients hospitalized by age category for selected years is also presented. This analysis excludes data from the transplant patients.

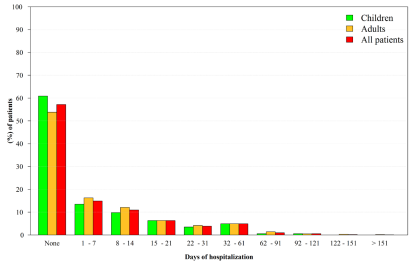
Except for anti-conceptive therapy, the percentages are based on **511** children and **568** adults from a total of **1079 non transplant patients**.

#### VISITS TO CF CARE CENTERS AND HOSPITALIZATION

In 2014, most of the patients (n = 902) 83.6% had a minimum of the four recommended visits to a CF reference centre during the year.

The figures below concern the number of days of hospitalization. In 2014, 57.2% of the patients were not hospitalized: 60.9% of the children (n = 311) and 53.9% of the adults (n = 306) were not hospitalized. A small proportion, (nine patients, 0.9%) was hospitalized for over three months.





Children ---- Adults 50 40 (%) at patients 20 10 2002 2004 2006 2008 2010 2012 2014 2000 Time (years) BMR-RBM 2014©

Figure 20 Proportion of patients hospitalized for more than two weeks by age and year

#### RESPIRATORY THERAPIES

**To prevent and treat respiratory infections**: Because mucus is stuck within the bronchi, evacuating mucus is one the most important interventions. This is the reason why a patient with CF has regular chest physiotherapy sessions. Most of the patients also learn the technique of autogenic drainage where the patients, through breathing techniques, perform drainage by themselves.

Different types of inhaled medication are used to treat the symptoms of CF. Maintenance inhaled medications include mucolytics that thin the sticky airway secretions such as RhDNase or hypertonic saline. Bronchodilators are given to dilate the bronchi. Inhaled antibiotics are used to treat infection, prevent or postpone colonization. In advanced lung disease oxygen is needed<sup>[50]</sup>.

Every year, more than half of the patients with CF are hospitalized, mostly for the treatment of a worsening pulmonary status (pulmonary exacerbations) caused by an infection requiring intravenous antibiotics.

Inflammation caused by repeated infections plays an important role in the progression of lung injury. This explains the interest in the use of anti-inflammatories such as azithromycin, inhaled corticoids or other anti-inflammatory drugs as complementary treatment.

Table 14 Physiotherapy, inhalation therapy, oral anti-inflammatory and antibiotics

	Chil	dren	Adı	ults	То	tal
Treatment						
Regular chest physiotherapy	504	98.6	542	95.4	1046	96.9
Antibiotics	450	88.1	496	87.3	946	87.7
Oral only	282	55.2	196	34.5	478	44.3
IV only	3	0.6	10	1.8	13	1.2
Oral and IV	147	28.8	255	44.9	402	37.3
Inhaled antibiotics	256	50.1	353	62.1	609	56.4
Inhalation therapy (excluding antibiotics)	484	94.7	519	91.4	1003	93.0
RhDnase	378	74.0	430	75.7	808	74.9
Other mucolytics	103	20.2	112	19.7	215	19.9
Hypertonic saline	296	57.9	336	59.2	632	58.6
Bronchodilators	399	78.1	421	74.1	820	76.0
Corticosteroids	203	39.7	344	60.6	547	50.7
Intranasal steroids	251	49.1	286	50.4	537	49.8
Oral anti-inflammatories	184	36.0	348	61.3	532	49.3
Azithromycin	171	33.5	327	57.6	498	46.2
Systemic corticosteroids	26	5.1	32	5.6	58	5.4
NSAID	6	1.2	57	10.0	63	5.8
Oxygen therapy	8	1.6	28	4.9	36	3.3

#### GASTRO-INTESTINAL AND NUTRITIONAL THERAPIES

**Optimizing the nutritional status:** The nutritional status of a patient is correlated with the disease severity therefore every person with CF should take a well-balanced high-calorie and high-fat diet. Most individuals with CF are pancreatic insufficient<sup>[43]</sup> and must take pancreatic enzymes at every meal to digest food correctly. Also supplements of vitamins ADEK are administered routinely. Some people with CF can only obtain a correct nutritional status by receiving supplemental feedings given overnight by a tube placed into the stomach (enteral feeding) or given intravenously (parenteral feeding).

Medications that inhibit the gastric acid secretion (proton pump inhibitors or H2 blockers) are given to treat symptoms of gastro-oesophageal reflux or to improve the efficacy of pancreatic enzymes in a higher pH environment. Prokinetics are medications that increase the tone of the lower oesophagus sphincter and enhance the gastrointestinal motility resulting in an acceleration of gastric emptying.

Table 15 | Digestive and nutritional therapies

Toronto	Chil	dren	Adı	ults	То	tal
Treatment	n	%	n	%	n	%
Pancreatic enzymes	433	84.7	440	77.5	873	80.9
Fat soluble vitamins (A,D,E and K)	446	87.3	430	75.7	876	81.2
Proton pump inhibitor and/or H2 receptor blocker	227	44.4	288	50.7	515	47.7
Ursodeoxycholic acid	136	26.6	122	21.5	258	23.9
Enteral feeding	21	4.1	4	0.7	25	2.3
Parenteral feeding	9	1.8	15	2.6	24	2.2
Gastrostomy tube	27	5.3	12	2.1	39	3.6
Prokinetics	14	2.7	25	4.4	39	3.6

## **OTHER TREATMENTS**

Monitoring the onset of other complications followed by appropriate therapeutic interventions: Possible complications of the disease need to be regularly monitored. When clinical, biological or imaging finding point towards liver disease, ursodeoxycholic acid is started. This hydrophilic bile acid normally present in human bile stimulates the biliary secretion so that the bile is less thick and would prevent liver damage. Depending on complications, other medications are prescribed such as insulin therapy when a patient develops CF related diabetes or bisphosphonates for osteoporosis. The treatment burden for CF patients is high. Most CF patients spend a lot of time every day performing therapies. This imposes also a substantial burden on their family<sup>[51]</sup>.

Table 16 Other treatments

T	Chil	dren	Adı	ults	Total	
Treatment		%		%		%
Insulin therapy	15	2.9	111	19.5	126	11.7
Oral therapy for diabetes	2	0.4	26	4.6	28	2.6
Bisphosphonates			11	1.9	11	1.0
Anti-conceptive therapy (females aged 12 and over)	14	18.9	117	57.1	131	47.0
Use of Psychopharmaca	11	2.2	54	9.5	65	6.0
CFTR Modulating Therapy	19	3.7	33	5.8	52	4.8

<u>Note</u>: Out of 1079 non transplant patients, 176 had CFRD. Among these, 116 used insulin therapy only, 19 used only oral therapy for diabetes while five patients used both oral therapy for diabetes and insulin therapy. However, in 36 patients with CFRD no use of either of the two treatments was reported.

## **INTRAVENOUS ANTIBIOTICS**

In the year 2014, 38.4% (n = 414) of the patients received IV antibiotics; 29.4% (n = 150) amongst the children and 46.5% (n = 264) among the adults. The figures below show the proportion that received IV antibiotics at home and/or in hospital in 2013 (top) compared to 2014 (bottom). There are few patients below six years taking IV antibiotics at home. Most of the adults have IV antibiotics both at home and in hospital.

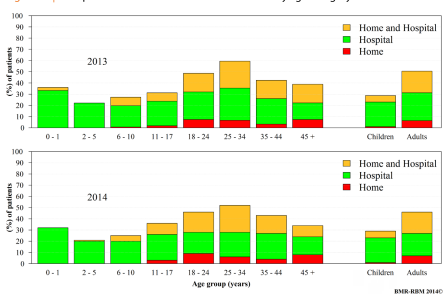


Figure 21 Proportion that used IV antibiotic treatment by age category

#### **ORAL ANTIBIOTICS**

The figures below show the days of oral antibiotics used in 2013 (top) and compared to 2014 (bottom). Unknown days or missing data was excluded from this analysis. In 2014, about 15.4% of the patients did not take any oral antibiotics; 14.6% of the children and 16.1% of the adults. More than a quarter of both the children (27.5%) and adults (30.6%) used oral antibiotics for over six months.

Figure 22 | Days of oral antibiotics Children 60 Adults 2013 50 40 30 20 All patients 10 0 None < 1 month 1 - < 3 months > 6 months 70 Children 60 Adults 50 40 30 20 2014 All patients 30.6 10 0 1 - < 3 months None < 1 month 3 - 6 months > 6 months Days of antibiotics (per os) BMR-RBM 2014©

## CHAPTER 10: TRANSPLANTS AND CF

When a patient with CF develops severe and progressive lung disease, lung transplantation may become an option. However, like other major surgeries, lung transplantation involves significant risks. A lung transplant is indicated only for patients who have a severe disease and who have exhausted all other forms of conventional medical treatment. For these patients, lung transplantation may offer prolonged survival and an improved quality of life; in some cases even a 'new' life.

The success rate of lung transplantation for CF patients is steadily improving. At present the actuarial survival rate is more than 70 % at 3 years and more than 45% at 10 years after primary transplant<sup>[52]</sup>. The longest surviving patients had their transplant operations now more than 20 years ago.

The first (heart)-lung transplant in a Belgian patient with CF was performed in 1988. Since this time about 230 patients<sup>(5)</sup> with CF<sup>[54]</sup> (208 reported in the CF Registry) have received a (heart)-lung transplant in Belgium and approximately 10 lung transplants per year are now performed for CF. Some patients with CF will need other types of transplantations such as liver transplantation for end-stage CF-related liver cirrhosis or renal transplantation for end-stage renal disease because of diabetes, antibiotic toxicity or the toxicity of immunosuppressive drugs required after lung transplantation.

This is a short analysis of data concerning the transplant patients. It includes a table with all transplants ever reported in the registry. This table shows the type and year of transplant and also the number of patients presumed still alive by the time of data collection 2014, categorized by the year of first transplant.

## TRANSPLANT STATUS

The registry records show that since inception of the registry in 1998, at least 208 patients, 101 male and 107 female, have benefitted from transplantation; either single or multiple. There are at least 26 patients who had a transplant on more than one occasion. In 2014, 13 patients were on the waiting list, 29 had been evaluated but were not on the list with one patient each refusing the transplant or being declined by the transplant centre.

The age at the first transplant was estimated using the last consultation in the year of transplant where available or the last day of the year if the former was missing. The data shows that the mean (SD) and median

<sup>5</sup> Source: Communication from Eurotransplant International Foundation

(range) age of the transplant patients in the year of the first transplant was 29.2 (9.7) and 28.3 years (5.3 - 60.7) respectively. The oldest patient at first transplant was 60.7 years. About 9.6% (20 patients) had the first transplant done before age 18 years.

In 2014, there were 151 patients with a recorded transplant. The mean (SD) and median (range) age of the transplant patients reported as alive in 2014 was 36.5 (9.9) and 35.3 (17.0 – 65.8) years respectively at the last consultation in 2014. 71 were male while 80 were female, while 99.3% of the transplant patients in the 2014 data were adults. Four transplant patients died in 2014.

The data presented in table 17 concerns all recorded transplants in the registry and also data of patients reported alive by the time of data collection for year 2014 (the numbers in the far right column). Slight differences may be noted with previous reports as we seek to consolidate the transplant data yearly. In the table, the numbers of transplants performed do not add up to the number of patients: - a patient can have several transplantations.

Multiple transplants are counted as separate transplant occasions. A total of 236 transplants on 208 patients are so far reported in the registry. There are 26 patients who have had a transplant on more than one occasion.

## **TYPE OF TRANSPLANT**

The most frequent transplant done is the Lung transplant, either single or with a liver or heart. So far, 189 isolated lung transplants have been carried out.

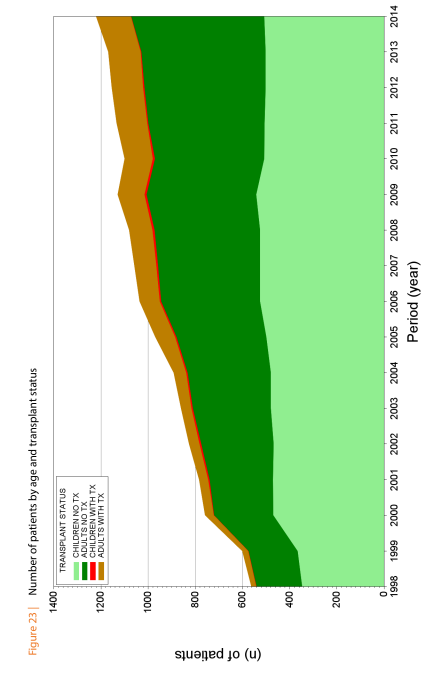
Table 17 | Type of transplant by year

		Totals							
Year	Lung	Lung-Heart	Lung-liver	Liver	Kidney	Liver-Kidney	Heart	Transplants performed	Patients alive
1991		2						2	1
1992								-	-
1993	•	2						2	1
1994		3						3	1
1995	2	3		1			•	6	1
1996		1						1	-
1997	1	3					•	4	4
1998	9	1						10	3
1999	5							5	2
2000	10							10	5
2001	13		1	4				18	13
2002	10			2				12	8
2003	9			1				10	8
2004	12					•		12	9
2005	10		1	3				14	8
2006	14			1	1		1	17	10
2007	14			1	1			16	9
2008	15				3			18	12
2009	10			1		1		12	9
2010	12		2	1				15	12
2011	15				2			17	12
2012	12							12	11
2013	11				3			14	8
2014	5				1			6	4
Total	189	15	4	15	11	1	1	236	-
Alive**	132	6	4	8	1	-	-	-	151*

<sup>\*</sup> Total number of patients alive in 2014 by transplant year considering the first transplant

<sup>\*\*</sup>Number of patients alive considering type of transplant

In the figure 23, the total number of patients recorded each year is presented as a proportion of children and adults, with and without a transplant. The figure shows that while the proportion of adults is increasing, so is the number of adult patients living with a transplant.



# CHAPTER 11: REPORTED DEATHS

This section has a summary of data on the deaths reported to the registry since inception in 1998. The number of deaths is also classified into age groups with information on the primary cause of death given in the latter part. Note that the stated causes of death are not mutually exclusive.

The data is updated each year from center reports with delays of up to two years noted in the confirmation of the data. Since 2013, this background data is automatically updated using the ConsultRN module by linking the registry data collection to the national registry database. The numbers may thus differ slightly from previously reported due to this automatic update.

### AGE AT DEATH

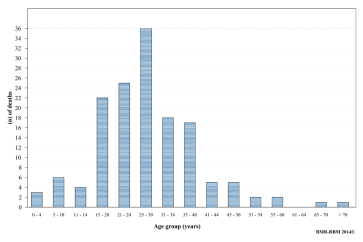
The data in the registry data shows that there have been 147 reported deaths, 79 male and 68 female. Eighteen (12.2%) of the deaths were in children below 18 years while 8 (5.4) of the deaths were in children younger than ten years. The mean (SD) and median (IQR) age at death for all reported cases are 28.5 (11.8) and 27.2 (13.4) respectively with the youngest at 0.6 years and the oldest case at 76.9 years. The table below shows the year and age category at death for confirmed cases.

Table 18 | Categorized age at death

	Age at death														
Year	0 - 4	5 - 10	11 - 14	15-20	21 - 24	25 - 30	31 - 34	35 - 40	41 - 44	45 - 50	21 - 54	55 - 60	65 - 70	75 - 80	Total
1998	1			2		1	1				•				5
1999				5	2	4		1							12
2000				1	2	1	3	2							9
2001		2		1	3	2		2							10
2002					1	3	4	2				1			11
2003		1	1	3	4	3	1	1		1					15
2004				2	2	3	1		1						9
2005	1				1	1									3
2006	1			2				2	1						6
2007		1	1	1		2		1	1	1					8
2008				2		3				1					6
2009			1	1	2	2	2		1	1					10
2010					3		2				1	1			7
2011		1			2	3	1		1	1					9
2012		1		1	1	2	2	2			1				10
2013				1	2	3									6
2014			1			3	1	4					1	1	11
Total	3	6	4	22	25	36	18	17	5	5	2	2	1	1	147

The figure below shows the number of confirmed deceased patients by age category. Most of the deceased patients were in the age category 25 - 30 years.

Figure 24 | Reported deaths since 1998 by age category



#### PRIMARY CAUSE OF DEATH

Most of the reported deaths are attributable to multiple causes, some not listed in the table below. Other causes of death thus include and are not limited to: - septic shock, multi-organ failure, terminal renal insufficiency, intoxication, hypoglycaemic coma and massive haemoptysis.

Table 19 | Primary causes of death for reported cases

Cause of death <sup>1</sup>	n	% *
Respiratory	62	42.2
Transplant	36	24.5
Other	30	20.4
Cancer	6	4.1
Cardiac	5	3.4
Liver	5	3.4
Suicide	3	2
Trauma	2	1.4
Unknown + missing	20	13.6

<sup>\*</sup>based on the total reported deaths

<u>Note</u>: The most common primary causes of death are associated with the respiratory system or are as a result of post-transplant complications.

<sup>&</sup>lt;sup>1</sup>Causes of death are not mutually exclusive and these percentages are attributable to the specific cause of death

#### CHAPTER 12: EDUCATION AND EMPLOYMENT

People with CF are living longer. According to the registry data the median patient age has increased from 14.9 in 1998 to about 21.4 in 2014 suggesting better life expectancy. This introduces new challenges and expectations alike. Despite their therapy burden, they are now studying, graduating and taking up a career either part-time or even full-time. They are thus taking up roles in society that were previously less attainable by cohorts decades back.

This section contains a summary of social and economic data, including education level of the people with CF registered in 2014. It details information on social allowances and employment status. All patients with substantial data, including those with a transplant are included in this analysis.

#### **EDUCATION**

Table 20 | Education level

	Chil	dren	Ad	ilts Total		tal
Education level						%
No school	44	8.7	61	9.4	105	9.1
Regular school / education attendance	454	90.1	126	19.5	580	50.4
Has finished school/education	4	0.8	455	70.4	459	39.9
Unknown	2	0.4	4	0.6	6	0.5
Subtotal	494		646		1150	
Missing	8		72		80	
Total	512		718		1230	

#### SOCIAL ALLOWANCES AND EMPLOYMENT

Table 21 | Social allowances or benefits and employment

Description	Chil	dren	Adults		
Description					
Additional child allowance	480	93.8	87	12.1	
Income support (in adults)			158	22.0	
Disability allowance	1	0.2	92	12.8	
Preferential tariff (in adults)			426	59.3	
Pension allowance (in adults)			11	1.5	
Integration support (in adults)			239	33.3	
Employment*			204	34.5	

<sup>\*</sup>data excludes 126 adults still actively in school

Amongst the 204 patients who said they were employed, 108 (52.9%) worked full time, 86 (42.2%) part-time while for 10 patients, this was unknown. Nine children and 34 adults had no information on social allowances.

#### CHAPTER 13: CF AND FERTILITY

With the improving trend in life-expectancy for people with CF, it is increasingly clear that social life and expectations other than education and employment come into play. Infertility in men with CF mainly arises from the failure of the vas deferens to develop properly<sup>[57]</sup>. A very small number (2 – 3%) are fertile<sup>[55, 56]</sup>. Many men with CF have moved into adolescence without being counselled on the possibility of being infertile<sup>[58]</sup>. This issue, including other aspects of their sexual functioning could be included and discussed openly in a transition care program as they move from the paediatric clinic to the adult clinic. Regular reproductive health education and a sperm analysis should be offered once they are at the adult clinic. Men can have children via assisted reproduction techniques including Percutaneous Epididymal Sperm Aspiration (PESA) or Testicular Sperm Aspiration (TESA) which are then used in either Intracytoplasmic Sperm Injection (ICSI) or in in-vitro fertilisation (IVF).

Despite most children with CF achieving near normal to normal growth, puberty and onset of other related physiological developments such as periods is often delayed in girls by one to two years. This is often a reflection of disease severity especially poor weight<sup>[60]</sup>, or general ill health or CFRD. Most girls with CF have normal menstrual cycles but with a higher incidence of missed (amenorrhea) or irregular periods. When the egg is released, the thick mucus in the cervix may act as a barrier to sperm penetration thereby reducing fertility. Most women with CF have become pregnant without any difficulty. The fertility issues in women with CF should however be dealt with like any other normal women.

Mothers with CF are more likely to have pre-gestational diabetes, deliver preterm and have a primary cesarean delivery<sup>[61]</sup>. Issues of persistent decrement in lung function have also been documented. The use of contraception is thus advised to prevent unplanned pregnancies. Preconception care aimed at improving pulmonary function, eliminating pulmonary infections and exacerbations, improving the nutritional status as well as routine screening for signs of CFRD, including during gestation, will improve fertility before conception and the pregnancy outcome<sup>[62]</sup>.

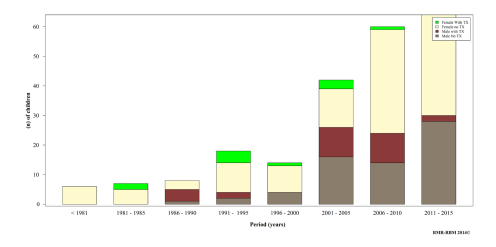
#### NUMBER OF BIRTHS REPORTED

Data from the registry shows that by 2014, there were at least 229 biological children reported from 142 parents. More than half of those who have children are female (n = 82, 57.8%). Among those with children, 25 (17.6%) are transplant patients. There were 52 (36.6%) parents homozygous for the F508del mutation. Most of the patients are alive except four who died in 2014, among them two transplant patients. The mean age at which they had the first child was 28.8 years (median 29.0 years) with a range 16.0 - 49.0 years. The mean and medians were 31.5 (32.0) and 26.8 (27.0) years respectively for the male and female parents.

The mean number of children was 1.6 (median 1.0, range 1.0 - 5.0). 74 patients (52.1%) have one child while 54 (38.0%) have two children and 14 (9.9%) have three children or more. 20 parents had twins of which five were transplant patients (one female). There are three patients who have had a child before their 18th birthday. In 2014, 13 patients had a child (16 children).

Consistently more children have been born each year since 2009 when compared to the years before. The figure below shows the total number of births among CF parents.

Figure 25 | Number of reported births by period



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# APPENDIX

## SUMMARY OF ITEMS COLLECTED IN THE REGISTRY BMR-RBM-BCFR

### **BMR-RBM 2014**

1. Background				
Year of collection Treating physician	::			
1.1 - Identification				
Patient ID (pseudo NISS) Date of birth Gender Place of residence Deceased? Date of death Order in the family Country of origin father Country of origin mother Height father (cm) Height mother (cm) Received transplant Diagnosis confirmed	: / /	//ale	□ Unknown	□ Missing data
1.2 - Diagnosis				
Date of clinical diagnosis Symptoms Respiratory problems Nasal polyposis/chronic s Chronic dia-steatorrhea/m Meconium ileus Intestinal obstruction (other Rectal prolapse Dehydration/electrolyte imt Failure to thrive Prenatal diagnosis Neonatal screening test Prolonged icterus Family history Infertility Other Specify other Missing data	inusitis alabsorption or than mecon. ileus) palance			
Neonatal screening test	☐ unknown	☐ not done		☐ performed positive

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1.3 - Sweat test			
Date of sweat test Type of sweat test Chloride Sodium	: /	□ Conductivity	☐ Missing data
1.4 - Genotype			
Legacy name Date of genotype Chromosome 1 Chromosome 1 other T status 1 Chromosome 2 Chromosome 2 other T status 2	: / /	········ 'T □ 9T □ Missing ·······	
cDNA name Date of genotype Chromosome 1 Chromosome 1 other T status 1 Chromosome 2 Chromosome 2 other T status 2	: / /		
	: / /		
Nasal transepithelial r	otential difference :   Not execut	ed □ Normal □	Evocative/Abnormal
Date	☐ Inconclusiv		
Dale	/ /		

# APPENDIX

# Non-transplant patients

2. Observation	on				
2.1 - Patient st	atus				
Patient status	☐ First registration☐ Revoked diagno		☐ In follow-up ☐ Not seen patiend ☐ Moved to other center ☐ Missing data		
2.2 - Cause of	death				
(type)	organ transplant ☐ Chronic Lung A ☐ Infection ☐ Other		ı (CLAD)		
Other cause (type)					
Cause unknow					
2.2 Last cons	ultation of the y	oar			
Date consultatio  Anthropometry  Weight (kg)  Height (cm)  Lung function	n :/. <b>y</b> :	. /			
Executed FVC (L) FEV <sub>1</sub> (L) FEF25-75 (L	:	□ No □ Ye		☐ Missing data	
	ung function of	-			
Date of best FVC (L) FEV <sub>1</sub> (L) FEF25-75 (L Weight (kg) Height (cm)	: :				

3. Microbiology				
3.1 - Microbiology: all cultures of the registra	ition ye	ear		
Microbiology executed Swabs Sputum Broncho-alveolar lavage (BAL) Missing values	□ Exec	cuted	□ Not executed □	I Missing data
3.2 - Pathogen ever found during the registra	ition ye	ear		
Pseudomonas aeruginosa Burkholderia cepacia complex Stenotrophomonas maltophilia Achromobacter xylosoxidans (Alcaligenes) Methicillin resistant Staphylococcus aureus (MRSA) Methicillin sensible Staphylococcus aureus (MSSA) Haemophilus influenzae Aspergillus Scedosporium prolificans Atypical Mycobacterium (NTM) Other No pathogens Missing values				
3.3 - Colonisation				
Number of exploitable months  Pseudomonas colonisation  Burkholderia cepacia complex colonisation  Stenotrophomonas colonisation  Achromobacter xylosoxidans colonisation  MRSA colonisation	□ No □ No □ No □ No	□ Yes □ Yes □ Yes	□ >=4 □ Unknown statu:	Missing data Missing data Missing data Missing data

4. Complications	
4.1 - Respiratory causes	
Allergic bronchopulmonary aspergillosis ABPA Treated Treatment for ABPA	□ No □ Yes □ Missing data □ No □ Yes □ Missing data □ Oral steroids and antifungal □ Omalizumab and antifungal □ Other □ Unknown □ Missing data
Pneumothorax	□ No □ Yes □ Missing data
Nasal polyps (having required/requiring therapy Massive haemoptysis Requiring embolization Bronchiectasis   No CT scan this year	3
4.0. Discotton constant	
4.2 - Digestive causes	DOEDD WILLIAM I
CF diabetes □ No □ IGT □ CFRD with fasting hypergl OGTT done this year	□ CFRD □ CFRD without fasting hyperglycemia □ Missing data □ No □ Yes □ Missing data
Acute pancreatitis Gastro-oesophageal reflux	□ No □ Yes □ Missing data □ No □ Yes □ Missing data
Cirrhosis with portal hypertension Gallstones	□ No □ Yes □ Missing data □ No □ Yes □ Missing data
Intestinal obstruction : requiring surgery Intestinal obstruction : not requiring surgery	□ No □ Yes □ Missing data □ No □ Yes □ Missing data
Gastroparesis Clostridium	□ No □ Yes □ Missing data □ No □ Yes □ Missing data
4.3 - Other complications	
CF related arthritis / arthropathy  Osteopenia / Osteoporosis □ z-score≥-1 □ z-score≤-2.5	□ No □ Yes □ Missing data □ z-score<-1 and >-2.5 □ Not done □ Missing data
Date of most recent DEXA Psychiatric disease	/ / □ Missing data
Cancer Type:	□ No □ Yes □ Missing data
Hypertension treated Other complications Type:	□ No □ Yes □ Missing data □ No □ Yes □ Missing data
4.4 - Surgery	
Surgery Type:	□ No □ Yes □ Missing data
General anaesthesia	□ No □ Yes □ Missing data
4.5 - New	
Renal Function (last consultation of the year Date	ar)
Creatinine (mg/dl)	

g data
g data
g data
g data

6. Transplantation	n					
Transplant 1 Transplant status	☐ Not evaluated ☐ Received transpla ☐ Missing data			ansplant cer t not on waiti		Waiting list Refused by patient
7. Social data						
·	•	Child 1 Child 2 Child 3 Child 4 Child 5	: / : / : / : /	☐ Missing of (month/yo (month/yo (month/yo (month/yo (month/yo (month/yo (month/yo	ear) ear) ear) ear) ear)	
School						
School status				□ Regular : lool/educatio		ation attendance  No school
Employment data			3			
Patient works Percentage Financial benefits		□ No □ Unkr	☐ Yes nown	☐ Missing o	data □Parttime	☐ Missing data
Additional child all Integration suppor Disability allowand Preferential tariff Pension Income support	t	No No No No No No No	☐ Yes	☐ Missing ( ☐ Missing ( ☐ Missing ( ☐ Missing ( ☐ Missing (	data data data data	
Family composition Household compo	sition	☐ Unch	•	☐ Changed☐ Missing of		
Number of siblings	s including the patient s with CF s decreased from CF					

# **Transplant patients**

2. Observation	on		
2.1 - Patient sta	atus		
Patient status	<ul><li>□ First registration in this cen</li><li>□ Revoked diagnosis</li><li>□ No information</li></ul>	ter □ In follow-up □ Moved to other ce □ Missing data	☐ Not seen patient enter
2.2 - Cause of	death		
Cardiac Respiratory Hepatic Trauma Suicide Associated with (type)	cancer		
Associated with	organ transplant		
	<ul><li>□ Chronic Lung Allograft Dysfu</li><li>□ Infection</li><li>□ Other</li></ul>	ntion (CLAD)	
Other cause			
Cause unknown			
2.3 - Last cons	ultation of the year		
Date consultation Anthropometry Weight (kg) Height (cm)			
Lung function			
Executed FVC (L) FEV <sub>1</sub> (L) FEF25-75 (L	:		☐ Missing data
2.4 - The best le	ung function of the year		
Date of best FVC (L) FEV <sub>1</sub> (L) FEF25-75 (L Weight (kg) Height (cm)	: :	  	
2.5 - Best post-	transplantation lung funct	ion (time interval between t	the 2 functions ≥ 3 weeks)
-	ction 1 : / /	Date lung function 2: Weight (kg) : Height (cm) : FVC (L) :	
		1	2014—1X-V2

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4.	Com	plica	tions

### 4.1 - Respiratory causes

4.2 - Digestive ca	iuses					
		☐ IGT asting hyperglyd			☐ CFRD without fas☐ Missing data☐ Missing data☐ Missing data☐ Control of the con	ting hyperglycemia
4.3 - Other comp	lications					
Cancer Type :					☐ Missing data	
4.4 - New						
Renal Function ( Date Creatinine (mg/dl)  Renal Function ( Date Creatinine (mg/dl)  Post-transplanta Chronic lung allog	(1 month post	-transplantatio				
☐ Bronchiolitis	Obliterans Synd	drome (BOS) :	_ _	BOS 0-p BOS 1 BOS 2 BOS 3		
☐ Restrictive al ☐ Not possible		ne (RAS)				
Post-transplant lyr	mpho-proliferat	ive disease	□ No	□ Yes	☐ Missing data	

5.1 - Therapy received during the	o region an	on year		
Number of consultations	<b>0</b>	□<4	□ ≥4	Missing data
Days in hospital				
5.2 - Respiratory system				
Home O2-therapy	□ No	•	■ Day and Night	Missing data
Intranasal steroids	□ No	☐ Yes		
Antiinflammatories p.o.	□ No	☐ Yes	Missing data	
Azithromycine	☐ No	☐ Yes		
5.3 - Digestive system				
Pancreatic sufficient	□ No	☐ Yes	☐ Missing data	
Pancreatic enzymes	☐ No	☐ Yes	☐ Missing data	
Fat soluble vitamins (ADEK)	□ No	□ Yes	☐ Missing data	
Ursodeoxycholic acid	☐ No	□ Yes	☐ Missing data	
Tube feeding	☐ No	□ Yes	☐ Missing data	
Gastrostomy	□ No	Yes	Missing data	
Parenteral feeding	☐ No	□ Yes	Missing data	
i.4 - Miscellaneous				
Oral therapy for diabetes	☐ No	☐ Yes	Missing data	
Insulin therapy	☐ No	Yes	Missing data	
Prokinetics	□ No	Yes	Missing data	
PPI + H2 receptor blocker	□ No	Yes	Missing data	
Anticonceptive therapy	☐ No	Yes	Missing data	
Psychopharmaca	☐ No	Yes	Missing data	
Biphosphonates	☐ No	Yes	Missing data	
Randomised drug trial	☐ No	Yes	Missing data	
CFTR modulating therapy	☐ No	□ Yes	Missing data	
Regular chest physiotherapy	□ No	☐ Yes	■ Missing data	
5.5 - Immunosuppressive treatm	ents (> 3 m	onths) (opt	ional)	
Calcineurin inhibitors				
Cyclosporine	☐ No	□ Yes	Missing data	
Tacrolimus	☐ No	□ Yes	■ Missing data	
Cell cycle inhibitors				
Azathioprine	☐ No	□ Yes	Missing data	
Mycophenolate mofetil (MMF)	□ No	□ Yes	■ Missing data	
Other immunosuppressive treatme	ents			
Steroids	□ No	□ Yes	■ Missing data	
Everolimus	□ No	☐ Yes	☐ Missing data	
Other immunosuppressive therapy, s			•	

6. Transplantatio	n							
Transplant 1								
Transplant status		ansplant		ransplant center t not on waiting list	☐ Waiting list☐ Refused by patient			
Type of transplant	<ul><li>□ Lung</li><li>□ Lung-liver</li></ul>		Lung-Heart Kidney	☐ Liver☐ Liver-Kidney	<ul><li>☐ Heart</li><li>☐ Missing data</li></ul>			
Year of transplant			:	_	_			
Precise date of entering the Tx waiting list :/								
Precise date of Tx		ŭ	://					
CMV status								
Recipient	pos.	☐ neg.	☐ unknown					
Donor	□ pos.	•	□ unknown					
Induction therapy	∕ □ No	☐ Yes	☐ Missing data					
Product			ŭ		Duration (days):			
Transplant 2								
Transplant status		ansplant	•	ransplant center t not on waiting list	<ul><li>□ Waiting list</li><li>□ Refused by patient</li></ul>			
Type of transplant	•		Lung-Heart Kidney	☐ Liver☐ Liver-Kidney	<ul><li>☐ Heart</li><li>☐ Missing data</li></ul>			
Year of transplant			:					
Precise date of en	tering the Tx w	aiting list	://					
Precise date of Tx			://					
CMV status								
Recipient	pos.	□ neg.	□ unknown					
Donor	pos.	neg.	□ unknown					
Induction therapy	y □ No	☐ Yes	■ Missing data					
Product					Duration (days):			
Transplant 3								
Transplant status		ansplant		ransplant center t not on waiting list	<ul><li>□ Waiting list</li><li>□ Refused by patient</li></ul>			
Type of transplant	<ul><li>□ Lung</li><li>□ Lung-liver</li></ul>		Lung-Heart Kidney	<ul><li>□ Liver</li><li>□ Liver-Kidney</li></ul>	<ul><li>☐ Heart</li><li>☐ Missing data</li></ul>			
Year of transplant			:					
Precise date of en	tering the Tx w	aiting list	://					
Precise date of Tx			://					
CMV status								
Recipient	pos.	☐ neg.	□ unknown					
Donor	pos.	☐ neg.	□ unknown					
Induction therapy	y □ No	☐ Yes	Missing data					
Product					Duration (days):			

7. Social data						
Pregnancy / Paternity						
Parenthood this year	☐ No	☐ Yes	☐ Missing d	ata		
Birthdays of the biological children for th	is patient					
	Child 1:	/	(month/ye	ear)		
	Child 2: / (month/year)					
	Child 3:					
	Child 4:	/	ar)			
	Child 5:					
	Child 6: / (month/year)					
School						
School status	☐ Unkno	own	□ Regular s	school/educa	tion attendance	
	☐ Has finished school/education			า	■ No school	
	■ Missir	ng data				
Employment data						
Patient works	☐ No	☐ Yes	■ Missing d			
Percentage	■ Unknown		□ Fulltime	□ Parttime	Missing data	
Financial benefits						
Additional child allowance	☐ No	☐ Yes	■ Missing d			
Integration support	☐ No	☐ Yes	Missing d			
Disability allowance	☐ No	☐ Yes	■ Missing d			
Preferential tariff	☐ No	☐ Yes	■ Missing d			
Pension	☐ No	☐ Yes	■ Missing d			
Income support	☐ No	☐ Yes	■ Missing d	ata		
Family composition						
Household composition	Unchanged		□ Changed			
	☐ First registration ☐ Missing data			ata		
Number of siblings including the patient						
Number of siblings with CF						
Number of siblings decreased from CF						
General remark						

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OPERATIONAL DIRECTORATE PUBLIC HEALTH AND SURVEILLANCE

Unit Health Services Research Juliette Wytsmanstreet 14 | 1050 Brussels | Belgium www.wiv-isp.be



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