MISSION OF THE CYSTIC FIBROSIS FOUNDATION
The mission of the Cystic Fibrosis Foundation is to cure cystic fibrosis and to provide all people with the disease the opportunity to lead full, productive lives by funding research and drug development, promoting individualized treatment, and ensuring access to high-quality, specialized care.

SOURCE OF DATA
Cystic fibrosis patients under care at CF Foundation-accredited care centers in the United States, who consented to have their data entered.

SUGGESTED CITATION
Cystic Fibrosis Foundation Patient Registry
2020 Annual Data Report
Bethesda, Maryland
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FIGURE PERMISSIONS
To request use of charts and data provided in this report, contact the CF Foundation Patient Registry team by email at reghelp@cff.org.

PHOTOGRAPHY BY
Dennis O’Clair
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Courtesy of the Russell family

SPECIAL ACKNOWLEDGMENTS
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Samar Rizvi
Thomas O’Neil
September 2021

Dear Friends and Colleagues:

We are pleased to share the 2020 Patient Registry Annual Data Report. The impact of two significant events is visible in this report: first, the impact of the COVID-19 pandemic and, second, the availability of elexacaftor/tezacaftor/ivacaftor (ETI) to many more people with cystic fibrosis (CF).

The delivery of CF care was dramatically impacted by the pandemic as indicated by the dramatic shift to telehealth in March 2020 and the continuation of telehealth visits throughout the year. On a positive note, “lost to follow-up” remained low at 3.2 percent, demonstrating the resilience of the CF community. One caveat — the pandemic resulted in less registry data than in past years, and that may affect some of our findings. For example, fewer microbiology cultures were collected, and that may have contributed to the lower annualized prevalence of potential microbial pathogens. Also, fewer pulmonary function measurements were collected, and some were from home spirometers. These factors may have impacted the precision of the annualized values of FEV\(_1\). I strongly encourage you to read the “About This Report” chapter in which some of these considerations are described.

Many of the positive trends in clinical metrics are almost certainly in part attributable to the second significant event in 2020, the prescription of ETI to many more people with CF. For example, the data showed a decrease in annual mortality rate (0.8 per 100 people in 2020 as compared to 1.6 per 100 people in 2010). We also observed significant decreases in the proportion of people with CF treated with IV antibiotics for a pulmonary exacerbation and in the number of people with CF who underwent a lung transplantation. One surprising observation is the marked increase in the number of pregnancies in 2020. While ETI likely contributed to many of these trends, it is difficult to tease out potential confounding by the pandemic and other factors. Further research will be needed.

Many thanks to each and every one of you who contribute to the success of the Registry — most importantly, people with CF and their families who generously agree to share their data and the Registry coordinators and care team members who collect and enter the data. It would not be possible without your vital contributions.

Thank you all for your hard work throughout the year and your commitment to the CF Foundation’s mission.

Bruce C. Marshall, MD
Executive Vice President and
Chief Medical Officer
Cystic Fibrosis Foundation
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ABOUT THIS REPORT

The Annual Data Report is based on data entered in the CF Foundation Patient Registry through our online portal, PortCF®. Data are entered by teams of dedicated health professionals in our nationwide network of more than 120 CF Foundation-accredited Care Centers.

Inclusion Criteria

This Annual Data Report contains data from individuals diagnosed with CF who (a) have consented to participate in the Registry, and (b) were seen in a CF Care Center during the 2020 calendar year, including those who were born, diagnosed, or died in the year. Data from individuals who have received a lung transplant were only included in the chapters on Demographics, Diagnosis, CFTR Gene Variants, Transplantation, and Survival.

The included populations represented in the figures vary and are based on the eligibility criteria described in the title and/or footnotes for the figure. Figures titled in gray reflect patient-level data that include individuals who have received a transplant. Figures titled in purple reflect patient-level data that do NOT include individuals who have received a lung transplant. Figures titled in blue present data on center-level variation across the CF Care Network. To reduce outliers, the majority of the center-level variation figures exclude centers with fewer than 10 individuals for that measure. Some center-level variation figures, such as cystic fibrosis-related diabetes (CFRD), infant, and pulmonary exacerbation measures, include centers with five or more eligible individuals for sufficient comparison.

Exclusion Criteria

Data from individuals with a diagnosis of CFSPID/CRMS (CF screen positive, inconclusive diagnosis/CFTR-related metabolic syndrome) or CFTR-related disorder were excluded from all figures except for the one on new diagnoses in the reporting year.

Since 2017, data from individuals who have received a lung transplant have been excluded from any chapters not specified in the inclusion criteria. In the past, data from individuals who received a lung transplant were included in many of the figures. This impacted reporting of prevalent complications such as CFRD and osteoporosis. Exclusion of data from individuals who received a lung transplant has resulted in a lower prevalence of these complications as compared to previous years.

More information about data included in the Annual Data Report and interpretation of the tables and figures can be found in the online Technical Supplement on cff.org.
INTRODUCTION

The CF Foundation Patient Registry (CFFPR) is composed of data collected via the CF Foundation Care Center Network, including participant demographic characteristics, routine clinical measurements, therapeutic history, hospitalizations, transplant, and vital status. Data reported to the CFFPR are obtained exclusively through participant contact with the CF Care Center, which is based on an interdisciplinary approach to monitor aspects of CF-related disease. Information is reported by CF care teams following encounters with physicians, respiratory therapists, social workers, dietitians, and other team members as well as hospitalizations and home-based IV treatment. Disruptions to the delivery of care have the potential to impact CFFPR data collection due to reduced participation in CF care and potential incomplete data entry.

The onset of the COVID-19 pandemic in early 2020 posed challenges to the delivery of CF care, including, but not limited to, temporary CF Care Center restrictions to in-person health care delivery and reduced staff availability due to deployment to the COVID-19 response or furlough. The impact of these disruptions on data reported to the CFFPR varies depending on the component of the care model. Interpretation of temporal trends comparing 2020 to prior years will require consideration of the completeness of reporting, as well as changes to the availability of CF care throughout the year and modifications to data collection via Port CF®.

The picture below shows a timeline of important events that should be considered when interpreting 2020 registry data.

Changes to PortCF in 2020

Changes to the CFFPR data fields available for data entry in 2020 included additional options to care location and a new case report form to document COVID-19 diagnoses. The COVID-19 pandemic was declared by the World Health Organization (WHO) on March 11, 2020. In response to the increase in telehealth utilization in the first half of 2020, the CFFPR data entry process was modified on May 28, 2020, to include a “By Phone” or “By Phone/Computer with Video” as care location options. Prior to May 28, encounters that occurred via telehealth were specified as “other” location. To minimize data entry burden, entries recorded as “other” were not updated. In the interval between March and June, it is likely that encounters reported as location “other” were, in fact, telehealth visits. The COVID-19 case report form was added to the registry data capture platform on July 9, 2020.
Use of Telehealth in Delivery of CF Care

The chart below shows the monthly total number of in-person clinic and telehealth/other encounters from January 2019 through December 2020. During this time, most encounters between people with CF and their care teams occurred in-person in the clinic. With the onset of the COVID-19 pandemic quarantines in March 2020, the number of in-person clinic encounters substantially decreased, and by April, they fell to 895 encounters. Conversely, the total number of encounters taking place outside the clinic setting (i.e., telehealth/other) rose to nearly 9,000 encounters in April. However, by December, the number of in-person clinic and telehealth/other encounters were nearly identical with a total number of encounters of 11,073, comparable to the monthly totals observed throughout 2019. The total number of encounter records in 2020 (128,038), was 7.5 percent lower than in 2019, with total number of encounters in clinics 44.0 percent lower than in 2019.

Coverage of Care in 2020

In addition to a reduction in the overall number of encounters reported, there was also a reduction in the completeness of measurements taken at each encounter. Based on established guidelines for CF care, encounters reported to the CFFPR typically include measurements of nutritional status, lung function, infection status, prescribed medications, and CF-related complications. In the chart below, the proportion of each encounter that included these data elements in 2020 is compared to 2019, by quarter. In the first quarter of 2020, encounters reported fewer measurements across all indicators but were within 10 percent of 2019 reporting. In the second quarter of 2020, a substantial number of encounters were missing FEV₁ measurements as well as microbiology culture results. The differences in reporting were likely due to several factors at both the individual and CF Care Center level. For example, some individuals with CF may have preferred telehealth visits in lieu of in-person care to maintain social distancing, and some CF Care Centers limited access to services such as pulmonary function testing or collection of culture specimens due to COVID-19 infection control protocols. By the end of 2020, these components of care improved but did not reach the level reported in 2019. The percentages of encounters with prescribed medications and complications in 2020 were lower than in 2019, but were consistently reported throughout the year.
While most data elements are reported at each encounter, data from all encounters during the calendar year are aggregated for each person (as an average for continuous measures and as ever/never for binary indicators). Below, the percentage of people with at least one measure is displayed to show the specific impact on the results shown in this report. Among CFFPR participants, there were fewer individuals with any microbiology culture results reported in 2020. Overall, the proportion of CFFPR participants with at least one height and weight measurement, as well as those age 7 years and older with at least one pulmonary function test (PFT) measurement, were similar to the proportions reported from 2017 to 2019. Reporting of medications and complications was also high at the individual level (at least one report per year). The proportion of individuals reporting at least one culture in 2020 was lower than in 2019.

The overall proportion of individuals with at least one data measurement reported in 2020 was largely consistent with the data in 2019. However, the number of measurements provided per individual were notably lower in 2020 compared to 2010. This is important because annualized measures of lung function, weight, and height are reported as an average of the maximum value from each quarter. Through 2019, the average number of PFT measurements reported per individual was nearly 5.0 measurements per year, which fell to an average of 2.5 measurements per individual in 2020, among those age 7 years and older. In addition, prior to 2020, microbiology cultures reported per individual were between 3.5 to 4.0 average cultures per year, which dropped to approximately 2.0 cultures per individual in 2020. It should be noted that the number of microbiology cultures sampled can impact the likelihood of detection of microorganisms.
Summary

The year 2020 presented many challenges that impacted delivery of care to individuals with CF and the reporting of data to the CFFPR. The information presented above is intended to provide some context for the differences in metrics and results observed in this year’s report as compared to previous years. We encourage readers of this report to interpret prevalence or incidence reported in 2020 in the context of reduced access to in-person care, which primarily impacted availability of culture surveillance, pulmonary function testing, and height and weight measurements.
### Summary of the Cystic Fibrosis Foundation Patient Registry, 2005–2020

#### Demographics

<table>
<thead>
<tr>
<th></th>
<th>2005</th>
<th>2010</th>
<th>2015</th>
<th>2019</th>
<th>2020</th>
</tr>
</thead>
<tbody>
<tr>
<td>People with CF (n)</td>
<td>23,091</td>
<td>26,354</td>
<td>29,044</td>
<td>31,293</td>
<td>31,411</td>
</tr>
<tr>
<td>Newly diagnosed individuals (n)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Detected by newborn screening (%)</td>
<td>18.8</td>
<td>56.2</td>
<td>58.7</td>
<td>60.9</td>
<td>63.1</td>
</tr>
<tr>
<td>Median age at diagnosis for all people with CF (months)</td>
<td>6</td>
<td>5</td>
<td>4</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Mean age (years)</td>
<td>17.8</td>
<td>19.2</td>
<td>20.9</td>
<td>22.7</td>
<td>23.3</td>
</tr>
<tr>
<td>Median age (years)</td>
<td>15.8</td>
<td>17.2</td>
<td>18.6</td>
<td>20.2</td>
<td>20.8</td>
</tr>
<tr>
<td>Adults ≥18 years (%)</td>
<td>42.6</td>
<td>47.5</td>
<td>51.6</td>
<td>55.8</td>
<td>57.2</td>
</tr>
<tr>
<td><strong>Race (not mutually exclusive)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White (%)</td>
<td>95.1</td>
<td>94.4</td>
<td>93.8</td>
<td>93.4</td>
<td>93.4</td>
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<tr>
<td>African American (%)</td>
<td>4.0</td>
<td>4.3</td>
<td>4.6</td>
<td>4.7</td>
<td>4.7</td>
</tr>
<tr>
<td>Other race (%)</td>
<td>2.1</td>
<td>2.8</td>
<td>3.3</td>
<td>3.8</td>
<td>3.9</td>
</tr>
<tr>
<td>Hispanic (any race) (%)</td>
<td>6.3</td>
<td>7.3</td>
<td>8.7</td>
<td>9.5</td>
<td>9.6</td>
</tr>
<tr>
<td>Males (%)</td>
<td>52.1</td>
<td>51.7</td>
<td>51.6</td>
<td>51.8</td>
<td>51.8</td>
</tr>
</tbody>
</table>

#### Mortality

<table>
<thead>
<tr>
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<th>2010</th>
<th>2015</th>
<th>2019</th>
<th>2020</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total deaths (n)</td>
<td>357</td>
<td>421</td>
<td>451</td>
<td>377</td>
<td>252</td>
</tr>
<tr>
<td>Annual mortality rate (per 100) (%)</td>
<td>1.5</td>
<td>1.6</td>
<td>1.6</td>
<td>1.2</td>
<td>0.8</td>
</tr>
<tr>
<td>Predicted median survival (five-year increments)</td>
<td>35.3</td>
<td>38.0</td>
<td>40.9</td>
<td>46.2</td>
<td>50.0</td>
</tr>
<tr>
<td>95% confidence interval (five-year increments)</td>
<td>34.2 - 36.5</td>
<td>36.9 - 39.1</td>
<td>39.2 - 42.0</td>
<td>45.1 - 47.6</td>
<td>48.5 - 51.3</td>
</tr>
<tr>
<td>Median age at death (years)</td>
<td>25.3</td>
<td>26.3</td>
<td>30.1</td>
<td>32.4</td>
<td>34.1</td>
</tr>
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</table>

#### GI/Nutrition

<table>
<thead>
<tr>
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<th>2010</th>
<th>2015</th>
<th>2019</th>
<th>2020</th>
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</thead>
<tbody>
<tr>
<td>BMI percentile in individuals 2 to 19 years (median)</td>
<td>46.4</td>
<td>50.9</td>
<td>54.7</td>
<td>58.3</td>
<td>61.3</td>
</tr>
<tr>
<td>Weight &lt;10th Centers for Disease and Control Prevention (CDC) percentile (%)</td>
<td>19.1</td>
<td>14.8</td>
<td>12.0</td>
<td>10.0</td>
<td>9.1</td>
</tr>
<tr>
<td>Height &lt;5th CDC percentile (%)</td>
<td>14.1</td>
<td>11.7</td>
<td>10.2</td>
<td>9.5</td>
<td>9.5</td>
</tr>
<tr>
<td>BMI in individuals 20 to 40 years (median)</td>
<td>21.2</td>
<td>21.7</td>
<td>22.1</td>
<td>22.5</td>
<td>23.1</td>
</tr>
<tr>
<td>Pancreatic enzyme replacement therapy (%)</td>
<td>80.0</td>
<td>86.9</td>
<td>86.6</td>
<td>84.4</td>
<td>83.8</td>
</tr>
<tr>
<td>Supplemental feeding - tube (%)</td>
<td>9.5</td>
<td>10.3</td>
<td>11.0</td>
<td>10.2</td>
<td>9.0</td>
</tr>
<tr>
<td>Supplemental feeding - oral only (%)</td>
<td>36.0</td>
<td>38.3</td>
<td>42.3</td>
<td>44.3</td>
<td>39.4</td>
</tr>
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</table>

#### Pulmonary

<table>
<thead>
<tr>
<th></th>
<th>2005</th>
<th>2010</th>
<th>2015</th>
<th>2019</th>
<th>2020</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVC % predicted (mean)</td>
<td>84.5</td>
<td>87.1</td>
<td>87.9</td>
<td>89.9</td>
<td>93.0</td>
</tr>
<tr>
<td>FEV, % predicted (mean)</td>
<td>73.7</td>
<td>75.7</td>
<td>76.5</td>
<td>78.7</td>
<td>82.8</td>
</tr>
<tr>
<td>FEV/FVC ratio (mean)</td>
<td>75.1</td>
<td>74.7</td>
<td>74.3</td>
<td>74.5</td>
<td>75.8</td>
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#### Respiratory Microbiology

<table>
<thead>
<tr>
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<th>2015</th>
<th>2019</th>
<th>2020</th>
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<tbody>
<tr>
<td>Number of Microbiology cultures per patient (mean)</td>
<td>2.9</td>
<td>3.5</td>
<td>3.8</td>
<td>3.8</td>
<td>2.2</td>
</tr>
<tr>
<td>P. aeruginosa (P aeruginosa or PA) (%)</td>
<td>56.6</td>
<td>51.5</td>
<td>47.6</td>
<td>43.2</td>
<td>32.0</td>
</tr>
<tr>
<td>B. cepacia complex (%)</td>
<td>3.1</td>
<td>2.5</td>
<td>2.5</td>
<td>2.6</td>
<td>1.6</td>
</tr>
<tr>
<td>S. aureus (%)</td>
<td>63.7</td>
<td>67.1</td>
<td>70.7</td>
<td>70.2</td>
<td>63.3</td>
</tr>
<tr>
<td>MSSA (%)</td>
<td>51.9</td>
<td>50.5</td>
<td>54.4</td>
<td>55.3</td>
<td>48.9</td>
</tr>
<tr>
<td>MRSA (%)</td>
<td>17.4</td>
<td>25.8</td>
<td>26.1</td>
<td>24.6</td>
<td>19.6</td>
</tr>
<tr>
<td>S. maltophilia (%)</td>
<td>12.5</td>
<td>13.9</td>
<td>13.6</td>
<td>11.9</td>
<td>6.3</td>
</tr>
<tr>
<td>Mycobacterial species (%)</td>
<td>-</td>
<td>10.1</td>
<td>11.9</td>
<td>13.9</td>
<td>10.0</td>
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Table continues on the next page
### Summary of the Cystic Fibrosis Foundation Patient Registry, 2005-2020

#### Health Care Utilization and Pulmonary Exacerbations (PEX)

<table>
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<tr>
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<tbody>
<tr>
<td>Outpatient visits to CF centers reported per year (mean)</td>
<td>4.2</td>
<td>4.7</td>
<td>4.4</td>
<td>4.3</td>
<td>2.4</td>
</tr>
<tr>
<td>Treated with IV antibiotics for a pulmonary exacerbation (%)</td>
<td>35.2</td>
<td>34.5</td>
<td>35.3</td>
<td>31.5</td>
<td>14.0</td>
</tr>
<tr>
<td>Number of pulmonary exacerbations per year (mean)</td>
<td>0.65</td>
<td>0.65</td>
<td>0.70</td>
<td>0.60</td>
<td>0.22</td>
</tr>
<tr>
<td>Number of days of treatment for all PEX per year (mean)</td>
<td>29.4</td>
<td>29.5</td>
<td>29.1</td>
<td>27.7</td>
<td>21.1</td>
</tr>
<tr>
<td>Number of days of home IV treatment for all PEX per year (mean)</td>
<td>13.5</td>
<td>11.9</td>
<td>10.9</td>
<td>9.3</td>
<td>6.9</td>
</tr>
<tr>
<td>Number of days of hospitalization for all PEX per year (mean)</td>
<td>15.8</td>
<td>17.6</td>
<td>18.2</td>
<td>18.3</td>
<td>14.3</td>
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#### Pulmonary Therapies

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<thead>
<tr>
<th>Therapy</th>
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<th>2015</th>
<th>2019</th>
<th>2020</th>
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</thead>
<tbody>
<tr>
<td>Dornase alfa (≥6 years) (%)</td>
<td>76.0</td>
<td>85.6</td>
<td>90.6</td>
<td>92.4</td>
<td>91.5</td>
</tr>
<tr>
<td>Inhaled tobramycin (PA+ and ≥6 years) (%)</td>
<td>69.5</td>
<td>70.6</td>
<td>70.2</td>
<td>68.3</td>
<td>65.3</td>
</tr>
<tr>
<td>Inhaled aztreonam (PA+ and ≥6 years) (%)</td>
<td>-</td>
<td>22.4</td>
<td>42.7</td>
<td>43.5</td>
<td>42.7</td>
</tr>
<tr>
<td>Azithromycin (PA+ and ≥6 years) (%)</td>
<td>53.0</td>
<td>70.0</td>
<td>67.0</td>
<td>64.3</td>
<td>61.9</td>
</tr>
<tr>
<td>Hypertonic saline (≥6 years) (%)</td>
<td>-</td>
<td>52.0</td>
<td>68.7</td>
<td>74.6</td>
<td>73.5</td>
</tr>
<tr>
<td>Oxygen (%)</td>
<td>5.6</td>
<td>10.8</td>
<td>11.1</td>
<td>10.9</td>
<td>7.4</td>
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<tr>
<td>Non-invasive ventilation (%)</td>
<td>-</td>
<td>2.3</td>
<td>2.9</td>
<td>3.0</td>
<td>2.4</td>
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#### CFTR Modulators

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<tr>
<th>Age Group</th>
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</thead>
<tbody>
<tr>
<td>Individuals &lt;2 years prescribed a modulator (%)</td>
<td>-</td>
<td>-</td>
<td>0.1</td>
<td>7.4</td>
<td>6.6</td>
</tr>
<tr>
<td>Individuals 2 to 5 years prescribed a modulator (%)</td>
<td>-</td>
<td>-</td>
<td>5.1</td>
<td>42.2</td>
<td>44.5</td>
</tr>
<tr>
<td>Individuals 6 to 11 years prescribed a modulator (%)</td>
<td>-</td>
<td>-</td>
<td>6.5</td>
<td>47.2</td>
<td>51.9</td>
</tr>
<tr>
<td>Individuals ≥12 years prescribed a modulator (%)</td>
<td>-</td>
<td>-</td>
<td>27.9</td>
<td>64.1</td>
<td>82.3</td>
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#### Transplants

<table>
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<tr>
<th>Transplant</th>
<th>2005</th>
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</thead>
<tbody>
<tr>
<td>Lung (all procedures) (n)</td>
<td>153</td>
<td>191</td>
<td>226</td>
<td>244</td>
<td>91</td>
</tr>
<tr>
<td>Liver (n)</td>
<td>14</td>
<td>17</td>
<td>17</td>
<td>20</td>
<td>21</td>
</tr>
<tr>
<td>Kidney (n)</td>
<td>3</td>
<td>7</td>
<td>10</td>
<td>14</td>
<td>10</td>
</tr>
</tbody>
</table>

#### Lost to Follow Up

<table>
<thead>
<tr>
<th>Year</th>
<th>2020</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lost to follow up (%)</td>
<td>3.7</td>
</tr>
</tbody>
</table>

---

**A** Includes data from transplant recipients.

**B** We anticipate that additional 2020 diagnoses will be entered into the Registry in 2021.

**C** Pulmonary function data throughout this report reflect the use of Global Lung Initiative (GLI) equations.

**D** Includes PA and multidrug-resistant PA found in any culture during the year.

**E** Includes MSSA and MRSA and reflects the prevalence of S. aureus among individuals who had a bacterial culture during the year. The percentages for MSSA and MRSA individually are greater than the total S. aureus percentage because MSSA and MRSA are not mutually exclusive.

**F** Percentage of people with CF with one or more mycobacterial species isolated out of those who had a mycobacterial culture during the year.

**G** Defined as a period of treatment with IV antibiotics in the hospital and/or at home.

**H** Among those with one or more pulmonary exacerbations in the year.

**I** Percentage of people with CF on therapy at any clinical visit in the year. All individuals noted as intolerant or having an allergy to a specific therapy were excluded.

**J** Includes Tobramycin solution for inhalation (i.e., TOBI®), TOBI™ Podhaler® (Tobramycin Inhalation Powder), and Bethkis® since 2013. In prior years, only TOBI® was available.

**K** Individuals were considered eligible if they met the selection criteria used in the first U.S. azithromycin trial.

**L** Includes continuous, nocturnal, or with exertion.

**M** Defined as patients seen in the previous reporting year (2019) but not the current reporting year (2020), and not known to have died.
DEMOGRAPHICS

The Registry contains data on people with CF from 1986 to 2020. During that time, substantial changes in specialized CF care have led to improved survival. This section shows the current and longitudinal distribution of demographic characteristics of individuals with CF in the Registry.

In 2020, there were 31,411 individuals with CF in the Registry. The number of adults with CF continues to increase, while the number of children remains relatively stable. In 2020, adults were 57.2 percent of the CF population, compared with 32.1 percent in 1990.

The decrease in the number of individuals reported in 2003 is due to a delay in obtaining informed consent forms before the close of the calendar year at some CF Care Centers.

In 2020, 9.6 percent of the individuals in the Registry identify as Hispanic. There has been a steady increase over the past 15 years, reflecting national population trends. Hispanics with CF tend to be younger than the overall CF population, with a median age of 14.41 years. In 2020, 3.5 percent of registry participants identified as Black or African-American.
In 2020, the median age of people with CF in the Registry was 20.3 years. The range is from birth to 89.7 years. Despite gains in survival, the age distribution remains markedly skewed toward younger ages.

The percentages in this chart differ from those presented in the summary table. This graph includes individuals in only one race category whereas the summary table includes individuals in more than one race category.

In 2020, the median age of people with CF in the Registry was 20.3 years. The range is from birth to 89.7 years. Despite gains in survival, the age distribution remains markedly skewed toward younger ages.

The percentages in this chart differ from those presented in the summary table. This graph includes individuals in only one race category whereas the summary table includes individuals in more than one race category.

In 2020, the median age of people with CF in the Registry was 20.3 years. The range is from birth to 89.7 years. Despite gains in survival, the age distribution remains markedly skewed toward younger ages.
Characteristics of Adults with CF

As a growing number of individuals with CF enter adulthood, it is encouraging to note that many are pursuing higher education and employment, are in committed relationships, and are having children of their own. Nearly three-fourths of adults with CF are either studying or working. However, these charts also highlight the approximately one-quarter of adults who report being disabled or unemployed.

**Education**
- College Graduate 30.7%
- Some College 29.3%
- High School Diploma or Equivalent 24.1%
- Master’s/Doctoral level degree 8.4%
- Less than High School 7.4%

**Marital Status**
- Married/Living Together 44.9%
- Single 49.7%
- Separated/Divorced 5.0%
- Widowed 0.4%

**Employment**
- Full-time 40.0%
- Part-time 12.1%
- Student 17.2%
- Retired 3.4%
- Unemployed 8.2%
- Homemaker 3.5%
- Disabled 15.7%
Over the last twenty years, there has been almost a tripling of the number of people who are married or living together and nearly four times the number of people with a college degree.

The number of pregnancies among women with CF has increased steadily since the 2000s. Registry data show that 619 women with CF were pregnant in 2020. The overall pregnancy rate among women with CF remained relatively constant between 2000-2019, with a marked increase in 2020. This stands in contrast to the pregnancy rate in the general U.S. population, which declined during this time.4

![Number of Pregnancies in Women 14 to 45 Years with CF, 2000, 2005, 2010, 2015, 2020](chart)

![Characteristics of Adults with CF, 2000, 2010, 2020](chart)
Health Insurance Information

Barriers to access insurance coverage for specialized care and treatments exist for some individuals with CF. Across all age groups, about half of the individuals in the Registry receive at least some component of their health insurance through federal or state-funded programs. Registry data show that in 2020, a majority of individuals with CF who were age 18 to 25 were covered under their parents’ health insurance plan. Despite the economic upheaval brought on by the COVID-19 pandemic, the percentage of people on private insurance was similar to 2019.

### Insurance Coverage in 2020

<table>
<thead>
<tr>
<th></th>
<th>Age &lt; 18 (%)</th>
<th>Age 18 - 25 (%)</th>
<th>Age ≥ 26 (%)</th>
<th>All (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Individuals (n)</td>
<td>13,228</td>
<td>5,890</td>
<td>11,604</td>
<td>30,722</td>
</tr>
<tr>
<td>Health insurance (e.g., private insurance)</td>
<td>50.6</td>
<td>63.8</td>
<td>65.3</td>
<td>58.7</td>
</tr>
<tr>
<td>Medicare/Indian Health Services</td>
<td>0.8</td>
<td>5.3</td>
<td>24.3</td>
<td>10.6</td>
</tr>
<tr>
<td>Medicaid/state programs</td>
<td>55.4</td>
<td>41.6</td>
<td>25.9</td>
<td>41.6</td>
</tr>
<tr>
<td>TriCare or other military health plan</td>
<td>3.1</td>
<td>2.1</td>
<td>1.8</td>
<td>2.4</td>
</tr>
<tr>
<td>Other</td>
<td>1.5</td>
<td>1.4</td>
<td>1.2</td>
<td>1.4</td>
</tr>
<tr>
<td>No health insurance</td>
<td>0.4</td>
<td>1.4</td>
<td>1.2</td>
<td>0.9</td>
</tr>
</tbody>
</table>

“The insurance coverage” reflects coverage at any point during the year; thus, these categories are not mutually exclusive (except for the “no health insurance” option).

### Additional Insurance Information in 2020

<table>
<thead>
<tr>
<th></th>
<th>46.8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individuals who participated in a patient assistance program (%)</td>
<td></td>
</tr>
<tr>
<td>Individuals 18 to 25 years covered under parents’ insurance (%)</td>
<td>52.6</td>
</tr>
</tbody>
</table>

“Patient assistance program” refers to any program that provides free medication or co-pay assistance.

A large proportion of children with CF use Medicaid or state programs, including 55.6 percent of children younger than 10 years of age. Though the overall prevalence of Medicare use is low among the entire population, it increases with age. Among adults aged 30 to 35 years, 19.7 percent report Medicare coverage. This increases to 26.9 percent among adults aged 40 to 64 years. Individuals aged younger than 65 years who receive Medicare have qualified based on the federal criteria for disability.
DIAGNOSIS

Diagnostic Characteristics of Individuals with CF

This section examines characteristics of individuals diagnosed with CF, as well as trends over time for two key diagnostic tests: genotyping and sweat chloride test.

In 2020, 63.1 percent of total new diagnoses and 88.5 percent of diagnoses among those less than 6 months old were reported as being detected by newborn screening (NBS). There is evidence that individuals diagnosed prior to the onset of symptoms have better lung function and nutritional outcomes later in life. Diagnosis in the newborn period also represents an important opportunity for CF Care Centers to partner with community physicians and families to ensure the best possible care and outcomes for infants with CF.

There may be a lag in reporting of newly diagnosed individuals, particularly infants born late in the year (i.e., late 2020), who were not seen at a CF Care Center before the end of the calendar year. Future reports will be adjusted to include these individuals for the 2020 diagnosis year.

With the widespread use of NBS for CF, it was necessary to develop terminology for individuals with a positive NBS who have inconclusive sweat test results and/or fewer than two CF-causing genetic variants. In the United States, this is referred to as CFTR-related metabolic syndrome (CRMS); other countries use the term CF screen positive, inconclusive diagnosis (CFSPID). Diagnosis guidelines, published in 2017, harmonized the criteria for CRMS/CFSPID. CRMS was added to the Registry as a diagnostic option in 2010. Entry of a diagnosis of CF versus CRMS into the Registry is a clinical decision; there is no requirement that individuals meet published diagnostic criteria for CF or CRMS. The percentage of CRMS/CFSPID decreased from 12.2 percent new diagnoses in 2019 to 11.4 percent in 2020.
Individuals can also be diagnosed with CFTR-related disorder. This option has been available in the Registry since 2010. Individuals with this diagnosis do not meet diagnostic criteria for CF or CRMS, are affected by CF-related conditions such as congenital bilateral absence of the vas deferens (CBAVD), and often have variants in the CFTR gene. The percentage of individuals reported to have CFTR-related disorder decreased from 9.2 percent of those diagnosed during the year in 2019 to 6.3 percent in 2020. Collection and analysis of data from individuals with CRMS and CFTR-related disorder will provide new and important information for these distinct populations.

In 2020, 494 newborn infants were diagnosed with CF. Of the 398 infants with a known gestational age at birth, 87.9 percent were born full-term, comparable with the figure for the general U.S. population. The mean birth weight for full-term infants with CF is also about the same as for the U.S. population, suggesting that babies born with CF do not initially show nutritional deficiencies.
The majority of those diagnosed in their first year via NBS are asymptomatic or minimally symptomatic at time of diagnosis. Among the 15.5 percent of infants diagnosed in 2020 before age one with meconium ileus (or other intestinal obstruction), 26.3 percent had bowel perforation. Since the number of infants with meconium ileus is small, the percentage with bowel perforation may fluctuate year to year, ranging from 15.9 percent to 34.9 percent over the last several years. Those diagnosed after age one often present with acute or persistent respiratory abnormalities.

### Symptoms Reported at CF Diagnosis

<table>
<thead>
<tr>
<th>Symptom</th>
<th>All Individuals (%)</th>
<th>Diagnosed in 2020 (%)</th>
<th>Diagnosed in 2020 Age &lt; 1 (%)</th>
<th>Diagnosed in 2020 Age ≥ 1 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Individuals (n)</td>
<td>31,411</td>
<td>708</td>
<td>522</td>
<td>186</td>
</tr>
<tr>
<td>Asymptomatic</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DNA analysis</td>
<td>13.8</td>
<td>25.1</td>
<td>23.4</td>
<td>30.2</td>
</tr>
<tr>
<td>Family history</td>
<td>14.5</td>
<td>10.5</td>
<td>9.7</td>
<td>12.5</td>
</tr>
<tr>
<td>Newborn (neonatal) screening</td>
<td>29.1</td>
<td>63.1</td>
<td>84.0</td>
<td>N/A</td>
</tr>
<tr>
<td>Prenatal screening (CVS(^A), amniocentesis)</td>
<td>2.6</td>
<td>4.7</td>
<td>6.0</td>
<td>N/A</td>
</tr>
<tr>
<td>Symptomatic</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acute or persistent respiratory abnormalities</td>
<td>34.5</td>
<td>14.7</td>
<td>1.6</td>
<td>50.0</td>
</tr>
<tr>
<td>CBAVD(^A) or infertility/GU(^C) abnormalities</td>
<td>0.7</td>
<td>3.7</td>
<td>0.0</td>
<td>13.5</td>
</tr>
<tr>
<td>Digital clubbing</td>
<td>0.5</td>
<td>1.1</td>
<td>0.0</td>
<td>4.2</td>
</tr>
<tr>
<td>Edema</td>
<td>0.5</td>
<td>0.1</td>
<td>0.2</td>
<td>0.0</td>
</tr>
<tr>
<td>Electrolyte imbalance</td>
<td>2.9</td>
<td>0.4</td>
<td>0.4</td>
<td>0.5</td>
</tr>
<tr>
<td>Failure to thrive/malnutrition</td>
<td>26.4</td>
<td>5.5</td>
<td>4.6</td>
<td>7.8</td>
</tr>
<tr>
<td>Liver problems</td>
<td>1.0</td>
<td>0.6</td>
<td>0.2</td>
<td>1.6</td>
</tr>
<tr>
<td>Meconium ileus/other intestinal obstruction</td>
<td>16.9</td>
<td>11.3</td>
<td>15.5</td>
<td>N/A</td>
</tr>
<tr>
<td>Nasal polyps/sinus disease</td>
<td>3.7</td>
<td>2.7</td>
<td>0.0</td>
<td>9.9</td>
</tr>
<tr>
<td>Rectal prolapse</td>
<td>2.5</td>
<td>0.4</td>
<td>0.2</td>
<td>1.0</td>
</tr>
<tr>
<td>Steatorrhea/abnormal stools/malabsorption</td>
<td>20.5</td>
<td>4.2</td>
<td>2.9</td>
<td>7.8</td>
</tr>
<tr>
<td>Other</td>
<td>5.1</td>
<td>4.8</td>
<td>1.6</td>
<td>13.5</td>
</tr>
</tbody>
</table>

Data are not mutually exclusive. We anticipate that additional 2020 diagnoses will be entered into the Registry in 2021.

\(^A\) Chorionic villus sampling
\(^B\) Congenital bilateral absence of the vas deferens
\(^C\) Genitourinary
Previous figures in this section refer to infants born or diagnosed in 2020; the following figure includes all individuals followed in the Registry in 2020.

### Age at Diagnosis of All Individuals with CF Seen in 2020

![Age at Diagnosis of All Individuals with CF Seen in 2020](chart)

#### Diagnostic Tests

**Sweat Chloride Testing**

Sweat chloride testing is an important diagnostic test recommended for all individuals regardless of genotype. In 2020, 90.4 percent of individuals in the Registry had a sweat chloride test result recorded. Baseline sweat chloride tests are becoming more important, as changes in sweat chloride are now viewed as indicators of the physiological effects of CFTR modulators. Fewer sweat tests were reported among individuals diagnosed in 2020. Access to sweat chloride testing may have been impacted by the pandemic.

### Percentage of Individuals with a Sweat Chloride Test Reported by Year of Diagnosis, 1990-2020

![Percentage of Individuals with a Sweat Chloride Test Reported by Year of Diagnosis, 1990-2020](chart)

Some individuals diagnosed in 2020 may not have had a sweat chloride test result entered in the Registry before the close of the reporting year.
Median sweat chloride test results have remained fairly consistent over time for individuals who are F508del homozygous. In contrast, there has been a gradual decline in median sweat chloride values among individuals who are not homozygous for F508del, suggesting that more individuals with “less severe” genotypes are being entered into the Registry.

**Genotyping**

The cystic fibrosis transmembrane conductance regulator (CFTR) gene and the most prevalent CF-causing variant (F508del) were both discovered in 1989. Since then, genotyping has become a key component of the diagnostic work-up. In addition, with the introduction of CFTR modulators, genotyping all people with CF is critical for research and clinical care. In 2020, 99.4 percent of individuals (n=31,203) in the Registry had been genotyped.
CFTR GENE VARIANTS

To date, more than 2,000 variants have been found in the CFTR gene. Some variants result in virtually no CFTR function, and others are associated with some residual function. Various strategies have been used to categorize variants with the goal of grouping individuals with CF with a similar disease course and prognosis. In previous reports, a five-variant class system was used. This classification system is helpful for understanding the impact of variants on the biosynthesis and function of the CFTR protein. However, it is increasingly recognized that this classification schema is an oversimplification, given that many variants lead to more than one defect in CFTR function.

In this era of CFTR modulators, a second approach is being evaluated for variant classification, referred to as theratyping. This system considers whether a variant responds to a specific CFTR modulator. However, theratyping is still early in its evolution and lacks a standardized format for reporting. In this section, we report data on specific variants and the difference in sweat chloride values between genotypes with little to no CFTR function versus those with residual CFTR function.

The most common CFTR variant is F508del: 85.8 percent of individuals in the Registry who have been genotyped have at least one copy of this variant. There is a substantial drop in prevalence to the next most common variants. No other variant is currently found in more than 5 percent of the population with CF.
### Prevalence of the 25 Most Common CFTR Variants in People with CF Seen in 2020

<table>
<thead>
<tr>
<th>CFTR Variant</th>
<th>Number of Individuals</th>
<th>Percentage of Individuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>F508del</td>
<td>26,791</td>
<td>85.8</td>
</tr>
<tr>
<td>G542X</td>
<td>1,417</td>
<td>4.5</td>
</tr>
<tr>
<td>G551D</td>
<td>1,345</td>
<td>4.3</td>
</tr>
<tr>
<td>R117H</td>
<td>1,025</td>
<td>3.3</td>
</tr>
<tr>
<td>N1303K</td>
<td>737</td>
<td>2.4</td>
</tr>
<tr>
<td>W1282X</td>
<td>683</td>
<td>2.2</td>
</tr>
<tr>
<td>3849+10kbC&gt;T</td>
<td>578</td>
<td>1.9</td>
</tr>
<tr>
<td>R553X</td>
<td>548</td>
<td>1.8</td>
</tr>
<tr>
<td>1717-1G&gt;A</td>
<td>494</td>
<td>1.6</td>
</tr>
<tr>
<td>621+1G&gt;T</td>
<td>490</td>
<td>1.6</td>
</tr>
<tr>
<td>2789+5G&gt;A</td>
<td>462</td>
<td>1.5</td>
</tr>
<tr>
<td>3120+1G&gt;A</td>
<td>383</td>
<td>1.2</td>
</tr>
<tr>
<td>ST</td>
<td>325</td>
<td>1.0</td>
</tr>
<tr>
<td>D1152H</td>
<td>318</td>
<td>1.0</td>
</tr>
<tr>
<td>3272-26A&gt;G</td>
<td>246</td>
<td>0.8</td>
</tr>
<tr>
<td>R1162X</td>
<td>243</td>
<td>0.8</td>
</tr>
<tr>
<td>2184insA</td>
<td>240</td>
<td>0.8</td>
</tr>
<tr>
<td>I507del</td>
<td>233</td>
<td>0.7</td>
</tr>
<tr>
<td>3659delC</td>
<td>222</td>
<td>0.7</td>
</tr>
<tr>
<td>G85E</td>
<td>214</td>
<td>0.7</td>
</tr>
<tr>
<td>1898+1G&gt;A</td>
<td>205</td>
<td>0.7</td>
</tr>
<tr>
<td>L206W</td>
<td>198</td>
<td>0.6</td>
</tr>
<tr>
<td>R334W</td>
<td>193</td>
<td>0.6</td>
</tr>
<tr>
<td>R347P</td>
<td>189</td>
<td>0.6</td>
</tr>
<tr>
<td>A455E</td>
<td>182</td>
<td>0.6</td>
</tr>
</tbody>
</table>

The number and percentage of individuals with a given variant include those with one or two copies of the mutation.

### F508del Variant Prevalence

<table>
<thead>
<tr>
<th>F508del Variant</th>
<th>Percentage of Individuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homozygous F508del</td>
<td>44.5</td>
</tr>
<tr>
<td>Heterozygous F508del</td>
<td>41.2</td>
</tr>
<tr>
<td>Neither F508del or Unknown</td>
<td>14.2</td>
</tr>
</tbody>
</table>
Among less common variants, the number of individuals with an R117H variant has increased over the years. Among those genotyped in 1993, less than 1 percent had an R117H variant, compared with 4.0 percent of those genotyped in 2020. This may be due to the inclusion of R117H in newborn screening algorithms, which has improved detection of these individuals. Of note, 107 (10.5 percent) of the 1,020 patients with an R117H variant had a sweat chloride value less than 30 mmol/L.

The clinical significance of the R117H variant depends in part on the poly-T tract variant on the chromosome. Research indicates that a shorter poly-T tract is associated with a higher likelihood of having CF. Unfortunately, the Registry has incomplete information on the poly-T tract status for most individuals (59.0 percent) who are reported as having a diagnosis of CF and an R117H variant. Of the 418 individuals with poly-T tract status recorded in the Registry, 154 (41.0 percent) are classified as having 5T.

---

**Poly-T Tract Status of Individuals with an R117H Variant, 2020**

<table>
<thead>
<tr>
<th>Status</th>
<th>Count</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Known</td>
<td>418</td>
<td>52.2%</td>
</tr>
<tr>
<td>Not 5T</td>
<td>264</td>
<td>31.6%</td>
</tr>
<tr>
<td>Unknown</td>
<td>602</td>
<td>71.9%</td>
</tr>
</tbody>
</table>

---
Individuals with variants typically associated with little or no CFTR function tend to have higher sweat test values, and are more likely to be prescribed pancreatic enzyme replacement therapy (PERT) than individuals with a variant typically associated with residual CFTR function (e.g., 96.7 percent vs. 32.8 percent).

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Median</th>
<th>5th Percentile</th>
<th>95th Percentile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individuals with variants associated with little or no</td>
<td>101.0</td>
<td>79.0</td>
<td>126.0</td>
</tr>
<tr>
<td>CFTR function N=19,348</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individuals with one or more variants associated with</td>
<td>71.0</td>
<td>28.0</td>
<td>112.0</td>
</tr>
<tr>
<td>residual CFTR function N=3,711</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individuals with one or more unclassified variants</td>
<td>91.0</td>
<td>38.0</td>
<td>121.0</td>
</tr>
<tr>
<td>N=4,688</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All individuals</td>
<td>98.0</td>
<td>50.0</td>
<td>124.5</td>
</tr>
</tbody>
</table>

*These charts use the highest sweat test value reported to the Registry. For 132 individuals, this value may reflect sweat chloride values after initiation of CFTR modulator therapy.*
GUIDELINES: CARE, SCREENING, AND PREVENTION

The CF Foundation sponsors the development of clinical practice guidelines to promote high-quality care for both physical and mental health care for individuals with CF during infancy, childhood, and adulthood.\textsuperscript{17-26} Many CF Care Centers report four clinic visits, two pulmonary function tests, and at least one microbiology culture annually for the majority of their CF patients. Similarly, among children aged 2 to 5, the majority have at least four visits and at least one culture each year.

CF Care Centers report that respiratory therapists/physical therapists, dietitians/nutritionists, and social workers evaluate most of their patients at least once per year, as recommended.\textsuperscript{21}

The influenza vaccination rate for people with CF age six months and older remains high across the CF Care Center network. Smoking and secondhand smoke exposure remain challenges, particularly for infants and young adults.

Patient Care Guidelines

The COVID-19 pandemic presented many challenges to providing care to individuals with CF including the redeployment of staff and the shift to telehealth. This required centers to be innovative in the delivery of care, the collection of PFTs (e.g., using home spirometers), and the collection of microbiology culture data. While the vast majority of individuals did not meet the recommended four clinic visits, four cultures, and two PFTs \textsuperscript{20,21} in 2020, almost all patients were seen at least once with a culture and PFT. In addition, there were other touchpoints with the clinical team through telehealth visits and transmission of home spirometry data.
The guidelines on infection prevention and control recommend that individuals with CF have quarterly microbiology cultures.20-22 In 2020, 89.9 percent of individuals received at least one culture, and 16.5 percent of individuals had four or more respiratory cultures.

The multidisciplinary care team plays an important role in CF care.20 With the exception of 2020, there has been an increase in the number of individuals with CF who receive an annual evaluation from a respiratory/physical therapist, dietitian/nutritionist, and social worker. While the percentage decreased in 2020, the multidisciplinary team continued to play a vital role in care, and each of these providers saw at least 80 percent of individuals a minimum of once during the year. In 2020, 69.8 percent of individuals were evaluated by all three specialists. In addition, to supporting individuals’ complex treatment regimens, the Registry now reports whether patients were seen by a pharmacist in clinic. In 2020, 45.8 percent of individuals were seen by a pharmacist at least once as compared to 46.7 percent in 2019 and 39.0 percent in 2018.
The CF Foundation recommends the inclusion of a mental health professional as part of the CF care team. Information on screening for anxiety and depression is included in the Complications section (page 62).

CF Foundation guidelines recommend annual measurement of fat-soluble vitamins to screen for vitamin deficiency. The CF Foundation Hepatobiliary Disease Consensus Group recommends a yearly panel of liver blood tests for all people with CF to screen for possible liver disease. Registry data suggest that these tests are being done for the majority of individuals.

### Percentage of Individuals Screened by or Monitored with Annual Labs, by Center

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individuals with Fat-Soluble Vitamins Measured</td>
<td></td>
<td></td>
<td>86.6</td>
<td>66.7</td>
<td>100.0</td>
</tr>
<tr>
<td>Individuals with Liver Enzymes Measured</td>
<td></td>
<td></td>
<td>89.0</td>
<td>66.7</td>
<td>100.0</td>
</tr>
</tbody>
</table>

### Flu Vaccine Status by Age in Years, 2020 (Stacked Bar Chart)

Influenza immunization is recommended for individuals with CF age six months and older on an annual basis. In 2020, 68.8 percent of the total population was reported to have received a flu vaccine, compared to 78.1 percent in 2019. Immunization coverage among children was higher than adults, with 75.6 percent of individuals younger than 18 years immunized compared to 63.2 percent of those 18 years or older. The proportion of individuals with an unknown vaccination status was 18.6 percent in 2020, an increase compared to 13.1 percent unknown in 2019. The change in these parameters during 2020 may reflect avoidance of health care settings/vaccine administration sites due to social distancing, as well as decreased emphasis on the importance of influenza vaccination given mask wearing, lower incidence of influenza infection, and heavy focus on COVID-19.
Infant Care Guidelines

The CF Foundation recommends that evaluation and treatment of infants detected by NBS should be done at a CF Foundation-accredited Care Center, with the goal of an initial visit within 24 to 72 hours of diagnosis. It is important to make a definitive diagnosis as quickly as possible to minimize the stress and uncertainty for families. For those diagnosed with CF, families can be educated about the disease and treatment can be started. Of infants detected by NBS in 2020, 90.1 percent had their first clinic encounter, care episode, or sweat test (first CF event) within 60 days of birth.

The CF Foundation infant care guidelines recommend monthly CF Care Center visits during the first six months of life and every one to two months in the second six months. Therefore, we expect infants with CF detected by NBS to have 9 to 12 visits in the first year of life. There is marked variation in the number of encounters for individuals in the first year of life across the CF Care Center network. Despite the COVID-19 pandemic, CF Care Centers provided a similar level of care to infants compared to previous years based on median number of clinic visits, number of cultures, and fecal elastase tests performed.
The chart shows data for children born in 2019 because a full year of data is available for these individuals. The median number of visits in the first year of life is nine.

Respiratory cultures are being collected at the majority of clinic visits for infants with CF. Infant care guidelines recommend that cultures be performed at least quarterly during the first two years of life.\textsuperscript{5}

The chart shows data for children born in 2019 because a full year of data is available for these individuals. The median number of cultures in the first year of life is five.
Fecal elastase testing, which provides an objective measure of pancreatic function, is recommended in the infant care guidelines. There is marked variation in the use of this test across the CF Care Center network. The guidelines also recommend that infants begin salt supplementation after diagnosis, and this is widely followed across the CF Care Center network.

### Infant Care Guidelines, by Center

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fecal Elastase Value Reported for Individuals Less Than 24 Months</td>
<td></td>
<td></td>
<td>71.4</td>
<td>11.1</td>
<td>100.0</td>
</tr>
<tr>
<td>Salt Supplementation in Individuals Less Than 24 Months</td>
<td></td>
<td></td>
<td>100.0</td>
<td>85.7</td>
<td>100.0</td>
</tr>
</tbody>
</table>

### Testing and Therapies for Individuals Under 24 Months of Age, 2013-2020

<table>
<thead>
<tr>
<th>Year</th>
<th>Salt Supplementation</th>
<th>Fecal Elastase</th>
</tr>
</thead>
<tbody>
<tr>
<td>2010</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2011</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2012</td>
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<td>2013</td>
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<td>2014</td>
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<td>2015</td>
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<td>2016</td>
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<td>2017</td>
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<td>2018</td>
<td></td>
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<tr>
<td>2019</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2020</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fecal elastase testing, which provides an objective measure of pancreatic function, is recommended in the infant care guidelines. There is marked variation in the use of this test across the CF Care Center network. The guidelines also recommend that infants begin salt supplementation after diagnosis, and this is widely followed across the CF Care Center network.
MICROBIOLOGY

This section provides information on trends in CF airway pathogens over time and by age group for individuals who never received a lung transplant. Infection prevention and control guidelines provide current best practices for reducing exposure to CF pathogens in the health care setting and in everyday life.27

As noted in the Introduction, the decreased number of cultures collected in 2020 is likely a contributing factor to the lower prevalence of bacterial and mycobacterial pathogens. There was a decrease from 94.7 percent to 85.8 percent (difference of 8.9 percent) of CFFPR participants reporting at least one microbiology culture and a decrease from 51.5 percent to 34.0 percent (difference of 17.5 percent) of individuals reporting at least one mycobacterial culture.

The prevalence of Pseudomonas aeruginosa (P. aeruginosa or PA) continues to decrease. This may relate in part to widespread implementation of therapy to eradicate initial acquisition.22,27 The prevalence of infection with multidrug-resistant P. aeruginosa (MDR-PA) dropped from 7.3 percent in 2019 to 4.2 percent in 2020.

Some of the increase in Staphylococcus aureus (S. aureus) through 2019 may be due to improved microbiologic practices for detection and reporting of Gram-positive organisms. From 2000 to 2010, there was a significant increase in the number of individuals with CF with a positive culture for methicillin-resistant S. aureus (MRSA). Since 2010, prevalence appears to have plateaued with the exception of 2020, which may reflect less frequent culture surveillance during the COVID-19 pandemic. The stabilization of prevalence is potentially due to increased awareness and infection prevention and control strategies.

Average Number of Cultures Per Individual, 2007-2020

![Average Number of Cultures Per Individual, 2007-2020](image-url)
The graph below shows the proportion of individuals in various age groups who cultured positive for the bacterial species indicated during 2020.
Pseudomonas aeruginosa

The percentage of individuals with a positive culture for *P. aeruginosa* has continued to decline over time, with the largest decrease observed among individuals younger than 18 years (44.5 percent had a positive culture in 2000 compared with 18.1 percent in 2020). Many individuals with CF now transition to adult care without *P. aeruginosa* in their respiratory tract.

Rates of MDR-PA infection are greatest in older adolescents and adults with CF. This finding likely reflects cumulative exposure to antibiotics. The clinical significance of this drug resistance is unclear. Among the individuals with CF who had at least one bacterial culture in 2020, 4.2 percent were reported to have MDR-PA. Among the individuals with CF with a *P. aeruginosa* infection in 2020, 13.2 percent were reported to have MDR-PA.

*Multidrug resistance is defined as resistance to all antibiotics tested in two or more antibiotic classes in a single culture.*
**Staphylococcus aureus**

Overall, more than half of individuals had at least one culture positive for methicillin-sensitive *S. aureus* (MSSA) in 2020. This chart shows that the highest prevalence of MRSA occurs in individuals between the ages of 10 and 30, whereas MSSA peaks among those younger than 15.
Nontuberculous Mycobacteria

Prevalence of nontuberculous mycobacteria (NTM) infections is increasing in the general population.\textsuperscript{28} The CF Foundation/European Cystic Fibrosis Society Guidelines Committee recommends that individuals with CF who are able to expectorate sputum be cultured for NTM infections annually.\textsuperscript{29} Individuals should also be screened before and six months after beginning azithromycin and annually thereafter.\textsuperscript{2} The data show improvement in screening rates over time, but wide variation by CF Care Center persists in these measures. Additionally, the median percentage of individuals (by Center) who produced a sputum sample in 2020 was lower than in 2019, 68.3 percent compared to 78.2 percent, respectively.

| Percentage of Individuals with a Mycobacterial Culture, by Center |
|---|---|---|---|
| Individuals Who Produced a Sputum Sample During the Year | 0 | 50 | 100 |
| Median | Min | Max |
| 68.3 | 0.0 | 100.0 |
| Individuals Taking a Chronic Macrolide Who Produced a Sputum Sample During the Year | 70.0 | 0.0 | 100.0 |

A throat swab is insufficient for a mycobacterial culture, so a patient must be able to produce sputum for this culture to be performed. A majority (67.8 percent) of the individuals who produced sputum for a bacterial culture also had a mycobacterial culture performed during the year although it decreased from 77.0 percent in 2019.
Among those cultured, the percentage of positive mycobacterial cultures increases until age 20, after which the percentage with a positive culture plateaus and remains relatively constant until age 60.

Of the 10,220 individuals who had a mycobacterial culture performed in 2020, 1,019 (10.0 percent) had a mycobacterial species isolated one or more times, a decrease from 13.9 percent in 2019. Notably, the relative proportion of *M. abscessus* isolated in 2020 was higher than reported more than a decade ago in a CF Foundation-supported multicenter prevalence study.30

### Mycobacterial Species Isolated in 2020

<table>
<thead>
<tr>
<th>Species</th>
<th>Number of Individuals</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>M. avium</em> complex (MAC)</td>
<td>498</td>
</tr>
<tr>
<td><em>M. abscessus</em>/<em>M. chelonae</em></td>
<td>369</td>
</tr>
<tr>
<td><em>M. gordonae</em></td>
<td>43</td>
</tr>
<tr>
<td><em>M. fortuitum</em></td>
<td>13</td>
</tr>
<tr>
<td><em>M. tuberculosis</em></td>
<td>4</td>
</tr>
<tr>
<td><em>M. kansasii</em></td>
<td>1</td>
</tr>
</tbody>
</table>

*Data are not mutually exclusive. Some individuals had more than one species isolated in 2020.*
NUTRITION

Nutritional outcomes are a key measure of health in people with CF. This section is divided into three age groups to report nutrition metrics: infants younger than 2 years, children 2 to 19 years, and adults 20 years and older for individuals who never received a lung transplant. Overall improvements in nutritional metrics are observed for all ages. Recent CF Foundation evidence-informed guidelines recommend enteral tube feeding as a means to improve age-dependent anthropometrics and nutrition in individuals with CF who are unable to consume adequate nutrition to meet goals.31

Goals for weight-for-length and BMI percentile in children are based on Centers for Disease Control and Prevention (CDC) growth curves. However, the CDC and the American Academy of Pediatrics recommend the use of World Health Organization (WHO) growth curves for children less than 24 months of age.32 WHO growth curves are used to report the data below.
The following charts show the population-level variation first for infants younger than 24 months using WHO weight-for-length, weight, and length percentiles by age and then for children aged 2 to 19 years using height, weight, and BMI percentiles. As would be expected for a large population, there is substantial variation observed for all three metrics in both age groups. Among infants younger than 24 months of age, the median values for weight-for-length are above the recommendation of 50th percentile. However, we still see evidence that infants’ growth is below what is expected for the U.S. population.

### All Individuals, WHO Nutritional Outcomes for Infants Under 24 Months

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>5th Percentile</th>
<th>95th Percentile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight-for-Length, N=1,504</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight Percentile, N=1,508</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length Percentile, N=1,504</td>
<td></td>
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<td></td>
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</tbody>
</table>

The goal BMI percentile established by the CF Foundation nutrition guidelines for children age 2 to 19 years is at or above 50 using CDC growth curves. In 2020, the median weight and BMI percentile was above the 50th percentile; however, height percentile remains well below that of the general population.

### All Individuals, CDC Nutritional Outcomes for Individuals 2 to 19 Years

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>5th Percentile</th>
<th>95th Percentile</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMI Percentile, N=13,462</td>
<td></td>
<td></td>
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<tr>
<td>Weight Percentile, N=13,697</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Height Percentile, N=13,619</td>
<td></td>
<td></td>
<td></td>
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</tbody>
</table>
For individuals born between 1991 and 2005, we observed improved weight and height percentiles with successive birth cohorts. Beginning with birth cohorts in 2006, there have been fewer observed increases compared to subsequent birth cohorts. For height, the most notable improvements were seen in the 2016-2020 birth cohort. More recently, there is less change between cohorts. This is potentially a result of the stabilization of improvements observed from universal early intervention due to NBS.33,34
The BMI goal established by the CF Foundation nutrition guidelines is at or above 22 for women and 23 for men age 20 years and older. Improvements in nutrition and dietary interventions have substantially decreased the percentage of adults that are malnourished, defined as a BMI less than 18.5 (4.3 percent in 2020 compared to 16.2 percent in 2000). However, 37.0 percent of adults have a BMI in the range categorized by CDC as overweight (26.7 percent) or obese (10.3 percent), with a higher prevalence in men (41.3 percent) than women (32.4 percent). The percentage of adults who are overweight or obese has more than doubled in the past 20 years (14.3 percent in 2000).

### Table: BMI Value for Adults 20 Years and Older

<table>
<thead>
<tr>
<th></th>
<th>15</th>
<th>20</th>
<th>25</th>
<th>30</th>
<th>35</th>
<th>Median</th>
<th>5th Percentile</th>
<th>95th Percentile</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Individuals</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N=14,622</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>23.6</td>
<td>18.6</td>
<td>32.9</td>
</tr>
<tr>
<td>Women</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>23.1</td>
<td>18.5</td>
<td>33.9</td>
</tr>
<tr>
<td>N=6,948</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Men</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>24.1</td>
<td>18.7</td>
<td>32.3</td>
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<td>N=7,674</td>
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<td></td>
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<td></td>
</tr>
</tbody>
</table>

Significant progress in nutritional outcomes continues for all people with CF. Aging of the CF population and a greater number of late diagnoses with genotypes associated with milder disease may be contributing to the trend in adult nutritional outcomes shown below.
For some individuals with CF, tube feeding is a tool to improve nutritional outcomes. Use of tube feeding is most common in children and adolescents with up to 20 percent of children in some age groups reporting tube feedings.

**Infant Feeding**

The majority of infants with CF receive formula feeding as the primary form of feeding or as a supplement to breastfeeding. Cow’s milk-based formula with the standard caloric density of 20 calories per ounce is the most common feeding used from birth to age 3 months. More calorie-dense formulas are used after 3 months of age. CF Foundation infant care guidelines recommend human breast milk or standard infant formula as the initial form of feeding. Fortified human breast milk, calorie-dense formulas, or complementary foods are recommended if the infant is failing to gain weight adequately.⁵

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**Form of Feeding by Age in 2020**

Infants may be included in more than one age category. They may also be counted more than once within an age category if different forms of feeding were recorded during separate clinic visits while within the same age category.
PULMONARY AND NUTRITIONAL OUTCOMES

Pulmonary and nutritional outcomes are two key measures of CF health. These metrics are the primary focus of quality improvement work within the CF Care Center network. The data show that for all people with CF, better pulmonary function, and higher BMI percentile are associated.

Pulmonary and nutritional goals\textsuperscript{25} are as follows:

- **For children**, FEV\textsubscript{1} percent predicted greater than or equal to 100, and BMI percentile meeting or exceeding the 50th percentile.
- **For adults**, FEV\textsubscript{1} percent predicted greater than or equal to 75, and BMI value greater than or equal to 22 for women and 23 for men.

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**FEV\textsubscript{1} Percent Predicted vs. BMI Percentile for Children 6 to 19 Years in 2020**

Goal: 50th percentile

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**FEV\textsubscript{1} Percent Predicted vs. BMI Value for Adults 20 to 40 Years in 2020**

Goal: 22 for Women, 23 for Men
The figures below on the left show median BMI percentile and FEV₁ percent predicted values for each center in 2020. The figures on the right show how median values for all centers have improved over the last 30 years.
PULMONARY FUNCTION

Pulmonary function is an important clinical indicator of the health of individuals with CF. This section provides information on trends in pulmonary function by age for individuals who have never had a lung transplant, as well as variations in pulmonary function across CF Care Centers. Pulmonary function is measured using the FEV₁ percent predicted and calculated using the Global Lung Initiative (GLI) reference equations.¹

Successive birth cohorts show improved pulmonary function across all ages for individuals who are old enough to reliably perform pulmonary function testing. The majority of individuals aged 18 years have an FEV₁ percent predicted approaching normal or greater than or equal to 70.

As noted in the Introduction, we captured approximately half the usual number of spirometry measurements per individual in 2020 and some of that data was derived from home spirometers. Also of note, there were fewer height measurements reported for growing children. These factors may have impacted the precision of the population, center and individual-level FEV₁ percent predicted values. Across all birth cohorts, the median FEV₁ percent predicted has historically declined as individuals age, with more recent birth cohorts reporting a higher median value compared to older birth cohorts at a given age. In contrast to trends reported in 2019, the figure below illustrates increases in median FEV₁ percent predicted by age among ages at which FEV₁ was measured in 2020. Among individuals born between 2011-2015, median FEV₁ percent predicted increased to 100 percent predicted among those who attained 8 to 9 years of age in 2020. Among individuals born 1991-1995, the median FEV₁ percent predicted increased across 25 to 29 years of age compared to the prior downward trend observed in earlier ages for that group. Overall, secular improvements in lung function continue to be observed across all birth cohorts. For example, children aged 6 to 9 years in 2020 have nearly 10 percent higher median FEV₁ percent predicted compared to median FEV₁ percent predicted measured among the 1991-1995 cohort when those individuals were 6 to 9 years of age.

Median FEV₁ Percent Predicted, by Age and Birth Cohort

![Median FEV₁ Percent Predicted, by Age and Birth Cohort](chart.png)
The proportion of people with CF aged 18 years who are in the normal/mild lung disease category (FEV\textsubscript{1} ≥70 percent predicted) more than doubled from 40.1 percent in 1990 to 87.4 percent in 2020. The proportion in the severe lung disease category aged 18 years (FEV\textsubscript{1} <40 percent predicted) decreased from 23.6 percent in 1990 to 1.8 percent in 2020.

Although much focus is placed on spirometry, it is not a sensitive measure of early lung disease in CF and may underrepresent the extent of early structural lung disease. With that in mind, the vast majority of children have normal or "mild" impairment in pulmonary function. This proportion decreases until age 35, when the population has nearly equal proportions of individuals with normal/mild or moderate lung disease.
The median FEV₁ percent predicted among individuals aged 6 to 11 is 98.6 percent and for individuals aged 18 years or older it is 77.1 percent. Indeed, half of the individuals aged 18 years or older have only mildly reduced lung function with an FEV₁ percent predicted greater than 70.0 percent. However, a wide spectrum of lung function is observed among adults with a quarter having a FEV₁ percent predicted less than 50.0 percent.

<table>
<thead>
<tr>
<th>FEV₁ Percent Predicted</th>
<th>0</th>
<th>25</th>
<th>50</th>
<th>75</th>
<th>100</th>
<th>125</th>
<th>Median</th>
<th>5th Percentile</th>
<th>95th Percentile</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Individuals 6 to 11 Years&lt;br&gt;N=4,221</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>98.6</td>
<td>69.4</td>
<td>121.2</td>
</tr>
<tr>
<td>All Individuals 12 to 18 Years&lt;br&gt;N=4,431</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>96.8</td>
<td>63.8</td>
<td>119.1</td>
</tr>
<tr>
<td>All Individuals 18 Years and Older&lt;br&gt;N=14,187</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>77.1</td>
<td>33.8</td>
<td>110.9</td>
</tr>
</tbody>
</table>
Pulmonary Exacerbations

Pulmonary exacerbations, characterized by intravenous (IV) antibiotic treatment in the hospital or at home, are associated with morbidity, mortality, and decreased quality of life. They are also a major driver of health care costs. This section displays trends in the rate of pulmonary exacerbations over time and by age group for individuals who have never received a lung transplant, as well as variation in exacerbation rates and treatment characteristics by CF Care Center.

Despite notable improvements in pulmonary function and nutritional status over the years, until recently there has not been a marked decline in the proportion of individuals with CF who are treated with IV antibiotics for pulmonary exacerbations. In 2020, a substantial decrease in number of reported exacerbations was observed, particularly in adolescents and adults. The pandemic, with less exposure to viruses, and elexacaftor/tezacaftor/ivacaftor are likely contributing factors.

Individuals with CF who are between ages 15 and 40 are more likely than those in other age groups to have a pulmonary exacerbation treated by IV antibiotics.

Pulmonary Exacerbations by Age in Years, 2020

Individuals Treated with IV Antibiotics for a Pulmonary Exacerbation, 2006–2020
When the CF Foundation developed guidelines for the treatment of pulmonary exacerbations in 2009, little published literature or data were available upon which to base recommendations. Current practice within the CF Foundation Care Center network indicates a median treatment duration of about two weeks, with adults more likely to complete some of their treatment at home. Further research is underway to develop evidence for best practices in the treatment of pulmonary exacerbations.

### Duration of Pulmonary Exacerbation Treatment in Days, by Center

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>7</th>
<th>14</th>
<th>21 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median Total Duration of IV Antibiotic Treatment for a Pulmonary Exacerbation in Individuals Less than 18 Years Old</td>
<td></td>
<td></td>
<td></td>
<td>12.7</td>
<td>4.0</td>
<td>18.0</td>
</tr>
<tr>
<td>Median Duration of Hospital Stay for Treatment of a Pulmonary Exacerbation in Individuals Less than 18 Years Old</td>
<td></td>
<td></td>
<td></td>
<td>10.5</td>
<td>2.5</td>
<td>14.5</td>
</tr>
<tr>
<td>Median Total Duration of IV Antibiotic Treatment for a Pulmonary Exacerbation in Individuals 18 Years and Older</td>
<td></td>
<td></td>
<td></td>
<td>13.0</td>
<td>6.8</td>
<td>17.5</td>
</tr>
<tr>
<td>Median Duration of Hospital Stay for Treatment of a Pulmonary Exacerbation in Individuals 18 Years and Older</td>
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<td></td>
<td></td>
<td>8.0</td>
<td>3.5</td>
<td>15.0</td>
</tr>
</tbody>
</table>

### Percentage of Total Pulmonary Exacerbation Treatment Duration in Hospital, by Center

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individuals Birth to 17 Years</td>
<td></td>
<td></td>
<td>91.2</td>
<td>64.7</td>
<td>100.0</td>
</tr>
<tr>
<td>Individuals 18 Years and Older</td>
<td></td>
<td></td>
<td>69.4</td>
<td>8.4</td>
<td>100.0</td>
</tr>
</tbody>
</table>
CFTR Modulators by Year, 2014–2020

<table>
<thead>
<tr>
<th>Year</th>
<th>Eligible, Not Prescribed</th>
<th>Total Eligible</th>
<th>% Eligible on Modulators</th>
<th>Total on Modulators</th>
</tr>
</thead>
<tbody>
<tr>
<td>2014</td>
<td>5,393</td>
<td>1,763</td>
<td>60.6%</td>
<td>1,069</td>
</tr>
<tr>
<td>2015</td>
<td>6,114</td>
<td>10,433</td>
<td>48.3%</td>
<td>5,040</td>
</tr>
<tr>
<td>2016</td>
<td>6,253</td>
<td>12,815</td>
<td>52.3%</td>
<td>6,701</td>
</tr>
<tr>
<td>2017</td>
<td>6,328</td>
<td>15,189</td>
<td>58.8%</td>
<td>8,936</td>
</tr>
<tr>
<td>2018</td>
<td>4,623</td>
<td>17,076</td>
<td>69.0%</td>
<td>11,788</td>
</tr>
<tr>
<td>2019</td>
<td>2,623</td>
<td>23,116</td>
<td>69.2%</td>
<td>16,005</td>
</tr>
<tr>
<td>2020</td>
<td>1,939</td>
<td>23,304</td>
<td>86.1%</td>
<td>20,067</td>
</tr>
</tbody>
</table>

On October 21, 2019, the US Food and Drug Administration (FDA) approved a new triple-combination therapy (elexacaftor/tezacaftor/ivacaftor) for individuals with CF who are aged 12 years and older and have at least one allele with the F508del variant. This significantly expanded eligibility of modulators for people with CF to approximately 80.0 percent of the registry population based on their age and genotype.

Approval of elexacaftor/tezacaftor/ivacaftor increased both the number of people who were eligible for CFTR modulators (23,004) and the number of people who were prescribed such therapies (20,067). In addition to the new CFTR modulator, age criteria for previously approved modulators were lowered over the last several years. Of those currently eligible, 86.1 percent of individuals were prescribed a CFTR modulator as of their last encounter of 2020. As shown in the chart below, lumacaftor/ivacaftor was prescribed for 1,895 individuals, tezacaftor/ivacaftor was prescribed for 1,480 individuals, and elexacaftor/tezacaftor/ivacaftor was prescribed for 14,753 individuals. The number of individuals eligible, but not prescribed a CFTR modulator fell by more than 50.0 percent from 7,111 in 2019 to 3,237 in 2020.
The overall proportion of individuals prescribed a CFTR modulator has expanded over time as visualized in the figure below. In 2014, only 7.0 percent of the population was eligible for CFTR modulator therapy, compared to approximately 80.0 percent in 2020. From 2015 to 2019, the proportion of the population eligible but not prescribed a CFTR modulator has ranged from 20.0 percent to 28.0 percent, and decreased to 14.0 percent in 2020. As shown in the figure below, a large proportion of individuals changed from either lumacaftor/ivacaftor or tezacaftor/ivacaftor to elexacaftor/tezacaftor/ivacaftor in 2020. Of those who were prescribed elexacaftor/tezacaftor/ivacaftor in 2020, 25.0 percent were prescribed a CFTR modulator for the first time.
Variability across centers exists with the proportion of eligible individuals prescribed a modulator ranging from a minimum of 69.2 percent to a maximum of 100 percent. There is a difference of approximately 5.0 percent in the median proportion of eligible individuals prescribed a modulator comparing pediatric and adult centers in 2020.
Pulmonary Therapies

Chronic pulmonary therapies are a major component of the treatment regimen for individuals with CF. This section provides data on uptake and trends in the prescription of pulmonary medications recommended for chronic use by the CF Foundation pulmonary guidelines committee for individuals never transplanted. Data are also provided on medications that are not recommended and on those for which the committee did not find sufficient evidence to recommend for or against chronic use.³⁷

Many of the CF therapies are used by between 60.0 percent and 80.0 percent of the eligible population. The use of most therapies has increased over time. In recent years, additional formulations of inhaled tobramycin have become available, and they are included in the chart below. Dornase alfa, which is recommended for individuals aged 6 and older, is used by the vast majority of people and its use continues to rise. Use of chronic inhaled antibiotics seems to have plateaued. The availability of multiple pulmonary therapies for CF is beneficial; however, this also contributes to treatment complexity and overall burden for individuals with CF and their caregivers. With the FDA approval of mannitol at the end of 2020, information about reported use will be provided in future years.

Tobramycin includes all available formulations of inhaled tobramycin. Dornase alfa, tobramycin solution for inhalation, and inhaled aztreonam were approved by the Food and Drug Administration in 1993, 1997, and 2010, respectively.
**Pulmonary Medication Prescriptions by Age**

There are three primary inhaled antibiotics used for treatment of *P. aeruginosa* infections. Tobramycin is used most frequently, followed by aztreonam, and then colistin. For all medications, peak use occurs during adolescence and young adulthood.

![Graphs of Any Tobramycin, Aztreonam, and Colistin use by age](images)

Dornase alfa and hypertonic saline are each prescribed for the majority of individuals with CF. Azithromycin is also widely used in individuals with *P. aeruginosa*, with peak use occurring at slightly older ages than for use of dornase alfa and hypertonic saline.

![Graphs of Dornase Alfa, Hypertonic Saline, and Azithromycin use by age](images)

A substantial proportion of individuals with CF are prescribed inhaled corticosteroids and, to a lesser degree, leukotriene modifiers. Oral corticosteroids are used infrequently.

![Graphs of Oral Corticosteroids, Inhaled Corticosteroids, and Leukotriene Modifiers use by age](images)
Bronchodilators are used extensively among individuals with CF. Almost all people with CF are prescribed beta agonists and a very small percentage are prescribed anticholinergics. The vast majority of those prescribed beta agonists are on short-acting formulations.

Multiple pulmonary therapies are available for individuals with CF. Inhaled medications are effective treatments for pulmonary disease, and they require time to prepare, administer, and clean equipment after treatment. Those with *P. aeruginosa* infection or colonization are typically prescribed inhaled antibiotics. Almost all individuals are prescribed at least one inhaled medication, and about 80.0 percent of individuals aged 15 to 40 are prescribed two or more of these therapies.

*Inhaled medication includes dornase alfa, hypertonic saline, and inhaled antibiotics. Inhaled antibiotic use includes the use of tobramycin, aztreonam, colistin, or other aminoglycosides.*
### Medications Recommended for Chronic Use

Recommended therapies are widely prescribed. However, there is considerable variation across the CF Foundation Care Center network. Some individuals with CF are using multiple inhaled antibiotics for the treatment of *P. aeruginosa* infections. CFTR modulators are discussed in a separate chapter (p. 50).

### Pulmonary Therapies Recommended for Chronic Use, by Center

<table>
<thead>
<tr>
<th>Therapy Description</th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dornase Alfa Prescription in Individuals 6 Years and Older</td>
<td></td>
<td></td>
<td>89.8</td>
<td>71.9</td>
<td>100.0</td>
</tr>
<tr>
<td>Any Inhaled Tobramycin Prescription in <em>P. aeruginosa</em>-positive Individuals 6 Years and Older</td>
<td></td>
<td></td>
<td>66.7</td>
<td>33.3</td>
<td>100.0</td>
</tr>
<tr>
<td>Azithromycin Prescription in Eligible <em>P. aeruginosa</em>-positive Individuals 6 Years and Older*</td>
<td></td>
<td></td>
<td>60.0</td>
<td>15.4</td>
<td>100.0</td>
</tr>
<tr>
<td>Hypertonic Saline Prescription in Individuals 6 Years and Older</td>
<td></td>
<td></td>
<td>75.4</td>
<td>40.0</td>
<td>100.0</td>
</tr>
<tr>
<td>Inhaled Aztreonam Prescription in <em>P. aeruginosa</em>-positive Individuals 6 Years and Older</td>
<td></td>
<td></td>
<td>41.8</td>
<td>0.0</td>
<td>80.0</td>
</tr>
</tbody>
</table>

*Individuals were considered eligible if they met the selection criteria used in the U.S. trial of azithromycin in individuals chronically infected with *Pseudomonas aeruginosa*.*

2 Individuals were considered eligible if they met the selection criteria used in the U.S. trial of azithromycin in individuals chronically infected with *Pseudomonas aeruginosa*.2
Medications with Insufficient Evidence to Recommend For or Against Chronic Use

In 2013, the CF Foundation pulmonary guidelines committee determined that there was insufficient evidence to recommend for or against the chronic use of inhaled beta agonists, inhaled anticholinergics, inhaled colistin, leukotriene modifiers, and ibuprofen for adults to improve lung function, reduce exacerbations, or improve quality of life. Use of colistin has decreased in recent years. Inhaled beta agonists are used extensively, but the other medications are used infrequently. Adult use of ibuprofen is less than two percent.

<table>
<thead>
<tr>
<th>Pulmonary Therapies with Insufficient Evidence to Recommend For or Against Chronic Use, by Center</th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inhaled Beta Agonist Prescription in Individuals 2 Years and Older</td>
<td></td>
<td></td>
<td>96.6</td>
<td>84.1</td>
<td>100.0</td>
</tr>
<tr>
<td>Inhaled Anticholinergic Prescription in Individuals 2 Years and Older</td>
<td></td>
<td></td>
<td>5.7</td>
<td>0.0</td>
<td>22.1</td>
</tr>
<tr>
<td>Leukotriene Modifier Prescription in Individuals 2 Years and Older</td>
<td></td>
<td></td>
<td>17.9</td>
<td>0.0</td>
<td>48.3</td>
</tr>
<tr>
<td>Inhaled Colistin Prescription in Individuals 6 Years and Older with <em>P. aeruginosa</em></td>
<td></td>
<td></td>
<td>8.2</td>
<td>0.0</td>
<td>31.7</td>
</tr>
</tbody>
</table>

Medications Not Recommended for Chronic Use

Inhaled steroids continue to be commonly prescribed, despite the recommendation against their chronic use in the absence of asthma or allergic bronchopulmonary aspergillosis (ABPA).

<table>
<thead>
<tr>
<th>Pulmonary Therapies Not Recommended for Chronic Use, by Center</th>
<th>0</th>
<th>50</th>
<th>100 Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inhaled Steroid Prescription in Individuals 6 Years and Older Without Asthma or ABPA</td>
<td></td>
<td></td>
<td>33.1</td>
<td>0.0</td>
<td>78.3</td>
</tr>
</tbody>
</table>
Medication Use in Young Children

In 2016, the CF Foundation released the first set of guidelines focusing on the preschool timeframe from ages two to five. Although the results of rigorous efficacy trials are not available for this age group, the guidelines recommend that dornase alfa and hypertonic saline be selectively offered to these individuals depending on individual circumstances. The chart below shows the use of medications among children younger than age six.

<table>
<thead>
<tr>
<th>Medication Use in Individuals Under 6 Years, 2020</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Age &lt; 2 Years (%)</td>
</tr>
<tr>
<td>-------------------</td>
</tr>
<tr>
<td>Number of Individuals (n)</td>
</tr>
<tr>
<td>Dornase alfa</td>
</tr>
<tr>
<td>Hypertonic saline</td>
</tr>
<tr>
<td>Inhaled bronchodilators</td>
</tr>
<tr>
<td>Inhaled corticosteroids</td>
</tr>
<tr>
<td>Inhaled tobramycin</td>
</tr>
<tr>
<td>Azithromycin</td>
</tr>
<tr>
<td>Inhaled aztreonam</td>
</tr>
</tbody>
</table>
**Airway Clearance Techniques**

The CF Foundation pulmonary guidelines recommend airway clearance for all individuals with CF. A high-frequency chest wall oscillation (HFCWO) vest is the most widely used airway clearance technique in persons with CF after infancy.

![Primary Airway Clearance Techniques by Age and Overall, 2020](chart)

*Positive Expiratory Pressure (PEP)/Oscillating Positive Expiratory Pressure (OscPEP)*

The CF Foundation pulmonary guidelines recommend aerobic exercise as an adjunct therapy for airway clearance and for its additional benefits to overall health. Many individuals with CF report exercising in addition to their primary method of airway clearance, with 31.9 percent of children and 46.8 percent of adults using exercise as one of their methods of airway clearance.

![Exercise as an Airway Clearance Technique (ACT) by Age in Years, 2020](chart)
Gastrointestinal (GI) Therapies

The CF Foundation infant clinical care guidelines recommend that pancreatic enzyme replacement therapy (PERT) be started for all infants with two CFTR variants associated with pancreatic insufficiency, a fecal elastase value below 200 µg/g of stool, and/or signs of malabsorption. In addition to pancreatic insufficiency, there is increasing attention on other GI manifestations of CF including Gastroesophageal Reflux Disease (GERD) and CF liver disease.

Overall, a large proportion of individuals of all ages are prescribed PERT. The decrease in the proportion of older individuals with CF prescribed PERT is most likely due to individuals without pancreatic insufficiency (thus not on PERT) surviving longer.

For individuals age two years and older, the recommended upper limit for enzyme dosing is 2,500 lipase units/kg/meal and a total of 10,000 lipase units/kg/day. The mean dose of lipase units/kg/meal for individuals 2 to 19 years is 1,967, and for individuals 20 years and older, the mean dose is 1,796, suggesting that children and adults on average are not exceeding the maximum recommended dose of PERT.

Infants with evidence of pancreatic insufficiency are recommended to receive 2,000 to 5,000 lipase units per feeding (total lipase dose), with adjustments as the infant grows. Registry data show that the mean highest dose of lipase among children younger than two years is 1,670 units/kg/meal.

For infants with CF younger than two years, the infant clinical care guidelines recommend assessment of pancreatic functional status by measurement of fecal elastase. Data on fecal elastase test results have been collected in the Registry since 2010, with an increased number of individuals undergoing fecal elastase testing. About 70.9 percent of infants born in 2020 were tested and have a fecal elastase value. Among individuals with a clinic visit, aged younger than two years in 2020, 29.1 percent did not have a fecal elastase value reported. Of those not tested, 89.6 percent were prescribed PERT. Almost all individuals with a known fecal elastase value of less than 200 µg/g of stool were prescribed PERT. Approximately 24.0 percent of individuals with fecal elastase values greater than or equal to 200 µg/g of stool were also prescribed PERT.
In 2020, 88.4 percent of individuals age 2 to 19 and 79.8 percent of individuals age 20 and older were prescribed CF-specific vitamins, including vitamins A, D, E, and K. In terms of liver manifestations of CF, 12.6 percent of individuals were prescribed ursodeoxycholic acid, which is most commonly prescribed to those with abnormal liver function tests or suspected CF liver disease.

Acid blockers are commonly prescribed for people with CF to treat GERD and/or to decrease the acidity of the stomach to increase the effectiveness of PERT. Overall, proton pump inhibitors (PPIs) are prescribed more often (43.3 percent of individuals) than H2 blockers (15.3 percent of individuals). H2 blockers are used more frequently in younger individuals and their use declines among older individuals. Use of PPIs increases with age until age 20 and overall is prescribed to 48.9 percent of individuals age 20 and older.

In 2020, 88.4 percent of individuals age 2 to 19 and 79.8 percent of individuals age 20 and older were prescribed CF-specific vitamins, including vitamins A, D, E, and K. In terms of liver manifestations of CF, 12.6 percent of individuals were prescribed ursodeoxycholic acid, which is most commonly prescribed to those with abnormal liver function tests or suspected CF liver disease.
COMPLICATIONS

Recognition and management of the multiorgan system health issues associated with CF is important for maintaining an individual’s health and quality of life. Complications of CF can affect many different aspects of health; they can be the direct result of the malfunction or deficiency of the CFTR protein or a downstream effect of the disease or its treatment. The prevalence of some non-pulmonary CF complications is higher among individuals who received a lung transplant than among individuals who have never had a lung transplant. Since 2017, individuals who had a transplant have been removed from the analyses. As a result, some complications (e.g., CF-related diabetes (CFRD) and osteoporosis) are lower in 2020 than reported previously.

CFRD remains an important and highly prevalent complication that greatly impacts quality of life and is associated with increased morbidity and mortality. As the population ages, a larger proportion of individuals are reporting complications typically seen in older adults, including bone and joint disease and sinus disease. Furthermore, along with the recent publication and implementation of mental health screening guidelines, an increase in the reporting of anxiety and depression has been observed.

<table>
<thead>
<tr>
<th>Complications of CF, 2020</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Number of Individuals (n)</td>
</tr>
<tr>
<td>Percent with no complications</td>
</tr>
<tr>
<td>Percent with complications not reported&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Cystic Fibrosis-Related Diabetes</td>
</tr>
<tr>
<td>Cystic fibrosis-related diabetes (CFRD)&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Hepatobiliary</td>
</tr>
<tr>
<td>Gallstones&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Gallstones, requiring surgery/procedure&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Liver disease, cirrhosis&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Liver disease, noncirrhosis&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Acute hepatitis&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Hepatic steatosis</td>
</tr>
<tr>
<td>Liver disease, other&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Bone/Joints</td>
</tr>
<tr>
<td>Arthritis/arthropathy</td>
</tr>
<tr>
<td>Bone fracture&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Osteopenia</td>
</tr>
<tr>
<td>Osteoporosis</td>
</tr>
<tr>
<td>Pulmonary</td>
</tr>
<tr>
<td>Allergic bronchopulmonary aspergillosis (ABPA)</td>
</tr>
<tr>
<td>Asthma</td>
</tr>
<tr>
<td>Hemoptysis</td>
</tr>
<tr>
<td>Hemoptysis, massive&lt;sup&gt;e&lt;/sup&gt;</td>
</tr>
<tr>
<td>Pneumothorax requiring chest tube&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Table continues on the next page
Complications of CF, 2020 continued

<table>
<thead>
<tr>
<th>GI</th>
<th>Age &lt; 18 (%)</th>
<th>Age ≥ 18 (%)</th>
<th>All (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distal intestinal obstruction syndrome (DIOS)</td>
<td>1.7</td>
<td>1.7</td>
<td>1.7</td>
</tr>
<tr>
<td>Fibrosing colonopathy/colic stricture</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Gastroesophageal reflux disease (GERD)</td>
<td>30.6</td>
<td>41.8</td>
<td>36.7</td>
</tr>
<tr>
<td>GI bleed requiring hospitalization (non-variceal)</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>History of intestinal or colon surgery</td>
<td>4.8</td>
<td>2.3</td>
<td>3.4</td>
</tr>
<tr>
<td>Pancreatitis</td>
<td>0.4</td>
<td>1.4</td>
<td>1.0</td>
</tr>
<tr>
<td>Peptic ulcer disease</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Rectal prolapse</td>
<td>0.5</td>
<td>0.1</td>
<td>0.3</td>
</tr>
<tr>
<td>Mental Health</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anxiety disorder</td>
<td>5.4</td>
<td>25.5</td>
<td>16.4</td>
</tr>
<tr>
<td>Depression</td>
<td>3.6</td>
<td>28.2</td>
<td>17.0</td>
</tr>
<tr>
<td>Other Complications</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cancer confirmed by histology</td>
<td>0.0</td>
<td>0.2</td>
<td>0.1</td>
</tr>
<tr>
<td>Hearing loss</td>
<td>1.4</td>
<td>3.7</td>
<td>2.6</td>
</tr>
<tr>
<td>Hypertension</td>
<td>0.4</td>
<td>6.5</td>
<td>3.8</td>
</tr>
<tr>
<td>Kidney stones</td>
<td>0.1</td>
<td>1.3</td>
<td>0.7</td>
</tr>
<tr>
<td>Nasal polyps requiring surgery</td>
<td>1.1</td>
<td>1.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Renal failure requiring dialysis</td>
<td>&lt;0.1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>Sinus disease</td>
<td>19.3</td>
<td>53.5</td>
<td>38.0</td>
</tr>
</tbody>
</table>

A Individuals for whom the complications case report form was not completed were considered to not have any complications, as in previous years.

B See table on page 66 for secondary complications.

C At the end of 2015, the data entry for complications was revised such that acute complications would no longer carry forward from one encounter to the next. We hypothesize that this is the reason for the decrease in the observed number of reported acute complications, most notably DIOS.

D See table below for secondary complications.

E Cause other than CFRD.

The table below highlights the prevalence of clinical manifestations of portal hypertension among individuals with cirrhosis.

Complications of Cirrhosis, 2020 (n=903)

<table>
<thead>
<tr>
<th></th>
<th>Age &lt; 18 (%)</th>
<th>Age ≥18 (%)</th>
<th>All (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Individuals (n)</td>
<td>261</td>
<td>642</td>
<td>903</td>
</tr>
<tr>
<td>Esophageal varices</td>
<td>21.1</td>
<td>24.5</td>
<td>23.5</td>
</tr>
<tr>
<td>Gastric varices</td>
<td>7.3</td>
<td>5.1</td>
<td>5.8</td>
</tr>
<tr>
<td>GI bleed related to varices</td>
<td>3.1</td>
<td>2.3</td>
<td>2.5</td>
</tr>
<tr>
<td>Splenomegaly</td>
<td>39.5</td>
<td>33.2</td>
<td>35.0</td>
</tr>
<tr>
<td>Hypersplenism</td>
<td>12.3</td>
<td>11.1</td>
<td>11.4</td>
</tr>
<tr>
<td>Encephalopathy</td>
<td>0.0</td>
<td>2.2</td>
<td>1.6</td>
</tr>
<tr>
<td>Ascites</td>
<td>3.4</td>
<td>7.8</td>
<td>6.5</td>
</tr>
</tbody>
</table>
CF Complications by Age, 2020

Reported complications differ in their distribution by age. Some are relatively stable over ages while others increase with age. The prevalence of bone disease and GERD is higher among older age groups. The prevalence of asthma peaks during adolescence and then decreases among adults while the prevalence of sinus disease increases in adolescence and in young adults and then remains high through the older ages. The prevalence of CFRD is higher in adolescence and adulthood. ABPA and DIOS are less common overall, but occur in all age groups. Liver disease is more prevalent in children. The prevalence of anxiety and depression rises during adolescence and young adulthood, and then appears to stabilize.
Cystic Fibrosis Related-Diabetes (CFRD)

Distal Intestinal Obstruction Syndrome (DIOS)

Liver Disease (All Types)

Allergic Bronchopulmonary Aspergillosis (ABPA)

Depression or Anxiety
Cystic Fibrosis-Related Diabetes (CFRD)

CFRD is an important complication of CF and is associated with weight loss, lung function decline, and increased mortality. Early diagnosis and treatment may minimize the impact of CFRD. The CF Foundation/American Diabetes Association clinical practice guidelines for CFRD recommend screening all individuals annually, starting at age 10, with an oral glucose tolerance test (OGTT).

Blood glucose testing is routinely performed at most CF Care Centers. The recommended OGTT is used less frequently, and substantial variation exists across CF Care Centers. It is encouraging to note that rates of screening adolescents for CFRD using the OGTT continues to increase (with the exception of 2020) since the CF Foundation clinical care guidelines for CFRD were published in 2010, while use in adults lags behind.

### Percentage of Non-Diabetic Individuals Receiving Glucose Testing, by Center

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>50</th>
<th>100</th>
<th>Median</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any Glucose Measurements in Non-Diabetic Individuals 10 to 17 Years</td>
<td></td>
<td></td>
<td></td>
<td>89.2</td>
<td>66.7</td>
<td>100.0</td>
</tr>
<tr>
<td>Any Glucose Measurements in Non-Diabetic Individuals 18 Years and Older</td>
<td></td>
<td></td>
<td></td>
<td>81.8</td>
<td>47.8</td>
<td>100.0</td>
</tr>
<tr>
<td>OGTT Screening in Non-Diabetic Individuals 10 to 17 Years</td>
<td></td>
<td></td>
<td></td>
<td>53.1</td>
<td>0.0</td>
<td>97.4</td>
</tr>
<tr>
<td>OGTT Screening in Non-Diabetic Individuals 18 Years and Older</td>
<td></td>
<td></td>
<td></td>
<td>24.0</td>
<td>0.0</td>
<td>76.2</td>
</tr>
</tbody>
</table>

### Blood Glucose and OGTT Screening for Individuals without CFRD, 2003–2020

[Graph showing blood glucose and OGTT screening percentages over years]
Prevalence of CFRD is higher among adults as compared to children with CF. Impaired glucose tolerance is most prevalent in adolescence; these individuals are at increased risk for developing CFRD and may benefit from increased monitoring.

The vast majority of individuals who were diagnosed with CFRD are noted in the Registry as being treated with insulin, as recommended in the CF Foundation CFRD clinical care guidelines.³⁹

<table>
<thead>
<tr>
<th>CFRD Treatment in 2020</th>
<th>Percentage of People with CFRD on Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dietary change</td>
<td>21.2</td>
</tr>
<tr>
<td>Oral hypoglycemic agents</td>
<td>4.0</td>
</tr>
<tr>
<td>Intermittent insulin (with illness, steroids, etc.)</td>
<td>4.6</td>
</tr>
<tr>
<td>Chronic insulin</td>
<td>71.4</td>
</tr>
<tr>
<td>No treatment noted in reporting year</td>
<td>13.3</td>
</tr>
</tbody>
</table>

*The data are not mutually exclusive and represent CFRD treatment at any point during the year.*
The clinical practice guidelines recommend regular hemoglobin A1c (HbA1c) measurements for individuals with CFRD. Although there is variation by CF Care Centers in the percentage of individuals with CFRD with one or more HbA1c measurements during the year, the majority of centers test most of their patients at least annually.

<table>
<thead>
<tr>
<th>Complications of CFRD in 2020</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Number of Individuals (n)</strong></td>
</tr>
<tr>
<td>--------------------------------</td>
</tr>
<tr>
<td>Number of Individuals (n)</td>
</tr>
<tr>
<td>Retinopathy</td>
</tr>
<tr>
<td>Microalbuminuria</td>
</tr>
<tr>
<td>Chronic renal insufficiency</td>
</tr>
<tr>
<td>Chronic renal failure requiring dialysis</td>
</tr>
<tr>
<td>Peripheral neuropathy</td>
</tr>
<tr>
<td>Any episodes of severe hypoglycemia</td>
</tr>
</tbody>
</table>

The goal established by the CF Foundation guidelines for CFRD is an HbA1c less than 7.0 percent for individuals with CFRD. More than half of individuals with CFRD are meeting this guideline.

Rates of secondary complications of CFRD, including retinopathy, microalbuminuria, kidney disease, and neuropathy, remain low. Episodes of severe hypoglycemia can be dangerous in individuals with CFRD. In 2020, they occurred slightly more in adults compared to children (4.7 percent vs. 3.0 percent). As the CF population continues to age, adult CF care providers should continue to screen individuals for these complications, as recommended by the CF Foundation clinical care guidelines for CFRD.
Depression and Anxiety

Addressing the mental health of all individuals with CF is critical to maintaining their overall health and quality of life. In 2015, the CF Foundation and the European CF Society published guidelines on screening and treatment for depression and anxiety among individuals with CF and caregivers of children with CF. These guidelines recommend annual screenings for all individuals with CF who are age 12 and older, as well as caregivers of children with CF. Uptake of mental health screening was rapid across CF Care Centers, and the majority of individuals were able to be screened in 2020 despite the pandemic and use of telehealth.

No differences are observed between screening rates for adolescents and adults. Of note, wide variation in screening across the CF Care Center network remains. Data on caregiver screening are not included in the Registry at this time.
Prevalence of both anxiety and depression increases through adolescence and early adulthood, then remains high at older ages. There is substantial overlap in individuals who experience both anxiety and depression. Among individuals who report anxiety or depression, 44.7 percent report both conditions.
TRANSPLANTATION

The majority of transplant procedures in individuals with CF are lung transplants, but other types of transplants are necessary as well. Lung transplantation remains an option for some individuals with severe lung disease. The number of lung transplant procedures for individuals with CF fluctuates yearly, and there was an overall upward trend until 2020 when a marked decrease in transplants occurred.

In 2020, there were 1,916 individuals in the Registry who were reported to have ever received a lung, kidney, heart, or liver transplant.

<table>
<thead>
<tr>
<th>Transplant Status of People with CF in 2020 (All Organs)</th>
<th>Number of Individuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accepted, on waiting list</td>
<td>73</td>
</tr>
<tr>
<td>Evaluated, rejected</td>
<td>154</td>
</tr>
<tr>
<td>Received transplant this year</td>
<td>116</td>
</tr>
<tr>
<td>Received transplant in a prior year</td>
<td>1,800</td>
</tr>
</tbody>
</table>

Lung Transplantation

There were 1,735 lung transplant recipients in the Registry in 2020, including 91 individuals who were reported to have received a lung transplant in 2020. The number of lung transplants among individuals with CF reported to the Registry in 2020 is far lower than reported in 2019 (62.7 percent) as compared to only a modest decrease (6.4 percent) in total lung transplants in the United States in 2020 (2,539) compared to 2019 (2,714). Of CF transplants performed in 2020, 5.5 percent were among individuals younger than age 18.
From 1992 to 2019, there has been a steady increase in the number of lung transplants observed among individuals with CF. In 2020, only 91 transplants were performed (37.3 percent of the number of lung transplants reported in 2019). This appears to be more than just the result of the pandemic and decreased utilization of health care, and is likely a result of the positive impact of elexacaftor/tezacaftor/ivacaftor on people with advanced CF lung disease.

In 2020, 32.0 percent of lung transplants occurred among individuals aged 18 to 29 years. With increasing eligibility of elexacaftor/tezacaftor/ivacaftor to younger children over time, hopefully the number of children needing lung transplant will sharply decline.

Overall, lung transplant recipients are a relatively small proportion of individuals included in the Registry; the majority are age 30 years and older.
While most CF care occurs within CF Foundation-accredited Care Centers, transplant and post transplant care typically occurs at transplant centers. Therefore, some of these individuals are lost to follow-up in the Registry. Optimal care for CF transplant recipients should include periodic follow-up at a CF Foundation-accredited Care Center.\textsuperscript{41}

### Status of Lung Transplant Recipients by Year of Transplant, 2001-2020

The last documented FEV\textsubscript{1} percent predicted prior to lung transplant demonstrates that the vast majority of individuals have FEV\textsubscript{1} percent predicted values less than 30 percent with 35.5 percent having FEV\textsubscript{1} percent predicted values less than 20 percent. The data underscore the severity of disease at the time of transplant and the potential benefit of earlier conversations about lung transplantation.\textsuperscript{42}

### Last FEV\textsubscript{1} Percent Predicted Prior to Lung Transplant, 2018-2020

Last FEV\textsubscript{1} Percent Predicted

- \textless 20: 14.7%
- 21-30: 46.5%
- 31-40: 35.5%
- \textgreater 41: 3.2%
SURVIVAL

Over time, there have been substantial improvements in the survival of people with CF. There are a number of different metrics to describe the survival of people living with CF in the United States. Definitions for these metrics are provided in the Technical Supplement that can be found on cff.org.

Median Predicted Survival

In 2020, the median predicted survival age in 2020 was 59.0 years (95 percent confidence interval: 56.4 - 65.1 years). Given the instability of annual survival estimates due to the relatively low number of deaths in any given year, the data is grouped into five-year increments. The following graph shows gains in median predicted survival from 1988 to 2020 in five-year increments.* Between 2016 and 2020, the median predicted survival age was 50.0 years (95 percent confidence interval: 48.5 - 51.3 years). This means that half of individuals born from 2016 to 2020 are predicted to live beyond 50.0 years of age. This prediction assumes no further improvement in mortality rate and thus does not take into account the potential impact of CFTR modulators on younger CF patients and other improvements in clinical care.

*Using the currently recommended method for calculating median predicted survival. For more information about the methodology, please see the Technical Supplement available at cff.org.
Median Conditional Predicted Survival

Median conditional predicted survival at specific ages is another metric to help understand changes in survival over time that considers an individual’s current age. This metric reflects the population-based median for all individuals of a specific attained age and does not take individual characteristics into consideration. The figure ends at age 40 because the numbers at older ages are currently too small to accurately predict survival. This metric does not take into account the potential impact of CFTR modulators and other improvements in clinical care.

Mortality Rate

The mortality rate in 2020 was 0.8 deaths per 100 individuals with CF in the Registry. There has been a steady decrease in the mortality rate over the last 30 years. This is encouraging, given that the median age of individuals in the Registry has increased from 12.3 years in 1990 to 20.8 years in 2020.
Median Age at Death

The median age at death was 34.1 years for the 252 people with CF who were reported to have died in 2020. About 11.5 percent of deaths occurred before 20 years of age. A comparison of the cumulative percentage for age at death between 1990 and 2020 shows a substantial shift of the curve toward the right with a larger proportion of deaths occurring at older ages. The median age at death reflects the age distribution of deaths in 2020 and cannot be used to predict survival of the entire population.

Causes of Death

Among the 252 deaths in 2020, the primary causes were respiratory/cardiorespiratory and transplant-related, similar to previous years. Of these, 40.5 percent of deaths occurred in people who were F508del homozygotes, reflecting their distribution in the Registry. Although 5.5 percent of people in the Registry are post transplant, 100 deaths (39.7 percent) occurred in transplant recipients. Note that the primary cause of death for these individuals was not always categorized as transplant-related. COVID-19 was a contributing factor in 11 deaths, including seven deaths in people who had received a lung transplant.

### Primary Cause of Death in 2020

<table>
<thead>
<tr>
<th>Cause</th>
<th>Number of Individuals</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory/cardiorespiratory</td>
<td>127</td>
<td>50.4</td>
</tr>
<tr>
<td>Transplant-related</td>
<td>45</td>
<td>17.9</td>
</tr>
<tr>
<td>Other</td>
<td>44</td>
<td>17.5</td>
</tr>
<tr>
<td>Unknown</td>
<td>17</td>
<td>6.7</td>
</tr>
<tr>
<td>Liver Disease/Liver Failure</td>
<td>8</td>
<td>3.2</td>
</tr>
<tr>
<td>Suicide or Drug Overdose</td>
<td>11</td>
<td>4.4</td>
</tr>
</tbody>
</table>
## CF DIAGNOSIS

**History of patient diagnosis***

**Date of Diagnosis:** (MM/DD/YYYY)  
Date is an approximation: □

**Diagnosis:**
- □ Cystic Fibrosis  
- □ CFTR-related metabolic syndrome  
- □ CFTR-related disorder  
- □ CF, CRMS and CFTR-related disorder all ruled out  

Patient was diagnosed with CF after false negative result by newborn screening:
- □ Yes  
- □ No  
- □ Unknown

**Diagnosis Suggested by the following:**
- □ Acute or persistent respiratory abnormalities  
- □ CBAVD (absent vas deferens) or related abnormalities  
- □ Classic clubbing  
- □ DNA Analysis  
- □ Electrolyte imbalance  
- □ Elevated immunoreactive trypsinogen (IRT) at CF newborn screening  
- □ Failure to thrive/malnutrition  
- □ Family history  
- □ Infertility/GU abnormalities  
- □ Less than 2 identified disease causing mutations  
- □ Liver problems  
- □ Meconium ileus/other intestinal obstruction (provide details below)
  - □ Meconium ileus with perforation  
  - □ Meconium ileus without perforation  
- □ Other neonatal bowel obstruction: ________________________________  
- □ Nasal polyps/sinus disease  
- □ Newborn (neonatal) screening  
- □ Non-diagnostic sweat chloride value (≥60 mmol/L)  
- □ Pancreatitis (not explained by other etiologies)  
- □ Persistent respiratory colonization/infection with a typical CF pathogen(s) (e.g., *Pseudomonas aeruginosa*)  
- □ Prenatal screening (CVS, amnio)  
- □ Pulmonary mycobacterial infection  
- □ Rectal prolapsed  
- □ Repeat Normal Sweat Testing  
- □ Steatorrhea/abnormal stools/malabsorption  
- □ Transepithelial potential differences  
- □ Other, specify: ________________________________  
- □ Unknown

**Date & value of documented positive quantitative pilocarpine iontophoresis sweat test (Chloride)**

**Date of Test:** MM/DD/YYYY
**Value (mmol/L): ________________________________
**Quantity Not Sufficient:** □

If sweat test value ≤60, CF diagnosis was suggested by:
- □ DNA Analysis/genotyping  
- □ Transepithelial potential differences  
- □ Clinical presentation (pancreatic fxn tests, Microbiology, etc.)  
- □ Unknown

### DEMOGRAPHIC DATA

**Demographics**
- CFF Patient Number: ______________________  
- Last Name: ______________________________  
- Last Name at Birth (if different): ______________  
- First Name: _____________________________  
- Middle Name: ____________________________  
- Last 4 digits of SSN: _______________________
- Date of Birth: (MM/DD/YYYY)  
- State of Birth: ____________________________  
- Gender:
  - □ Male  
  - □ Female  
- Current Zip: ______________________________  
- Is patient residing in the US permanently?
  - □ Yes  
  - □ No  
  - □ Unknown  
- Emergency Phone: ________________________  
- Email: __________________________________

**Race/Ethnicity Information**

**Race:**
- □ White  
- □ Black or African American  
- □ American Indian or Alaska Native  
- □ Asian  
- □ Native Hawaiian or Other Pacific Islander  
- □ Some other race  
- □ Two or more races  
  - If two or more races, specify Mixed Race components:
    - □ White  
    - □ Black or African American  
    - □ American Indian or Alaska Native  
    - □ Asian  
    - □ Native Hawaiian or Other Pacific Islander  
- Is the Patient of Hispanic Origin?
  - □ Yes  
  - □ No  
  - □ Unknown

**Death Information**

**Date of Death:** (MM/DD/YYYY)  
Check if date of death is approximate: □

**Primary Cause of death:**
- □ Respiratory/cardiorespiratory  
- □ Liver Disease/Liver Failure  
- □ Trauma  
- □ Suicide  
- □ Transplant related: Bronchiolitis obliterans  
- □ Transplant related: Other  
- □ Drug Overdose  
- □ Other  
- □ Unknown

**Additional Information**

**Additional Information:** ________________________________
## 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

### Parents' Information
*Information not required for patients 21 years of age and older*
- Not available: □
- Mother height: _____ cm _____ inches
- Father height: _____ cm _____ inches

### Birth Measurements
- Baby delivered:
  - Full term (>= 37 weeks gestational age)
  - Premature (< 37 weeks gestational age)
  - Unknown
- Specify gestational age (only if premature): _______
- Birth length: _____ cm _____ inches
- Birth weight: _____ lb _____ kg

### Genotype Information
For a list of mutation options, please contact reghelp@cff.org
- Has this patient been genotyped?  ___ Yes     ___ No
- Select Mutation 1: ________  Other genotype: ___________
- Poly T tract:
- Select Mutation 2: ________  Other genotype: ___________
- Poly T tract:
- Select Mutation 3: ________  Other genotype: ___________
- Poly TG repeats:
- Other genotype:
- Additional information about genotype not captured above: ____________________________

### ENCOUNTER DATA

#### Vital Signs/Encounter Start
- Encounter date: (MM/DD/YYYY)
- Location:  Clinic  Hospital  Home IV  Other
- Height: _____ cm _____ inches
  - [Height Percentile _______]  kg
  - [Weight Percentile _______]  lb
  - [BMI value: _______]  
  - [BMI Percentile: _______]  
  - [Weight for Length percentile: ___________]

#### Exacerbation Assessment
- Were there crackles (rales) on physical exam at this visit?
  - Yes  No  Physical exam data not available
- What was your assessment regarding pulmonary exacerbation at this visit?
  - Absent
  - Mild exacerbation
  - Moderate exacerbation
  - Severe exacerbation
  - Don't know/unable to answer
- Was a follow up visit scheduled?
  - Yes  No  Unknown
  - If Yes, indicate when:*
  - In less than 2 weeks
  - 2-4 weeks
  - 5-6 weeks
  - 7 weeks or later

If you determined that an exacerbation was present, please select the treatment course prescribed to treat the exacerbation:
- Increased airway clearance, exercise, and/or bronchodilators
- Oral NON-quinolone antibiotic (e.g. azithromycin, Bactrim, Augmentin, etc.)
- Oral quinolone antibiotic (e.g. ciprofloxacin (Cipro), levofloxacin)
- Inhaled antibiotic
- Inhaled antibiotic PLUS Oral NON-quinolone antibiotic
- Inhaled antibiotic PLUS an oral quinolone antibiotic
- None of the above
- (Note: if you elected to treat with hospital or home IV antibiotics, please start a care episode and enter the requested data.)

#### Consultations with Care Team Members
- Patient consulted with a Social Worker at this visit
- Patient consulted with a Dietitian/Nutritionist at this visit
- Patient consulted with a Physical therapist at this visit
- Patient consulted with a Respiratory therapist at this visit
- Patient consulted with a Pharmacist at this visit
- Patient was seen by a Mental Health Coordinator at this visit

#### Other
- Record any additional information about this encounter:
  - Custom field 1: ____________________________
  - Custom field 2: ____________________________
  - Custom field 3: ____________________________

#### Microbiology

##### Bacterial Culture
- Bacterial culture done?  □
- Date of Culture: (MM/DD/YYYY)
- Type of Specimen:
  - sputum
  - induced sputum
  - throat/nasal
  - bronchoscopy
- Culture Results:
  - Microorganisms
  - Normal flora
  - No growth/sterile culture
- Staphylococcus aureus:
  - MRSA (methicillin resistant Staph aureus)
  - MSSA (methicillin sensitive Staph aureus)
- *repeated entries can be recorded
- [ ] indicates values calculated by the registry
### 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

**Haemophilus influenzae** (any species): ☐

**Pseudomonas aeruginosa**: ☐
- mucoid ☐
- non mucoid ☐
- mucoid status unknown ☐

**Susceptibility Testing** (Please use the most resistant PA strain. If multiple PA strains are resistant to the same number of classes of antibiotics then use the following schema: Beta lactams> Quinolones>Aminoglycosides).

Resistant to All Aminoglycosides Tested (e.g., tobramycin, gentamicin, amikacin):
- Yes ☐
- No ☐
- Testing not done ☐

Resistant to All Quinolones Tested (e.g., ciprofloxacin, levofloxacin, moxifloxacin):
- Yes ☐
- No ☐
- Testing not done ☐

Resistant to All Beta Lactams Tested (e.g., ceftriaxone, imipenem, meropenem, piperacillin/tazobactam (Zosyn), ticarcillin/clavulanic acid (Timentin), aztreonam):
- Yes ☐
- No ☐
- Testing not done ☐

**Burkholderia species**: ☐
- B. gladioli ☐
- B. cenocepacia ☐
- B. multivorans ☐
- B. contaminans ☐
- B. stabilis ☐
- B. viennensis ☐
- B. dolosa ☐
- B. anhina ☐
- B. ambifaria ☐
- B. pyrrocina ☐
- B. ubonensis ☐
- B. arboris ☐
- B. latens ☐
- B. lata ☐
- B. metallica ☐
- B. seminatis ☐
- B. contaminans ☐
- B. diffusa ☐
- B. pseudomallei ☐

**Mycobacterial culture**
- Was Mycobacterial culture done? ☐
- Date of Culture: (MM/DD/YYYY)

**Type of Specimen**:
- sputum ☐
- induced sputum ☐
- bronchoscopy ☐

**AFB Smear**:
- Positive ☐
- Negative ☐
- Not done ☐

**Culture Results**:
- Microorganisms ☐
- Normal flora ☐
- No growth/sterile culture ☐

**Mycobacterial Species**:
- Mycobacterial tuberculosis ☐
- Mycobacterium abscessus/chelonae ☐
- Mycobacterium avium complex (MAC) ☐
- Mycobacterium fortuitum group ☐
- Mycobacterium gordonae ☐
- Mycobacterium kansasi ☐
- Mycobacterium marinum ☐
- Mycobacterium terrae ☐
- Other ☐

**Mycobacterium gordonae** ☐
**Mycobacterium kansasi** ☐
**Mycobacterium avium complex (MAC)** ☐
**Mycobacterium avium complex** ☐
**Mycobacterium avium** ☐
**Mycobacterium avium subsp. hominis** ☐
**Mycobacterium avium subsp. paratuberculosis** ☐
**Mycobacterium avium subsp. avium** ☐
**Mycobacterium kansasii** ☐
**Mycobacterium bovis** ☐
**Mycobacterium fortuitum** ☐
**Mycobacterium leprae** ☐
**Mycobacterium gastri** ☐
**Mycobacterium xenopi** ☐
**Mycobacterium smegmatis** ☐
**Mycobacterium tuberculosis** ☐

**Other**: ☐

**Key:**
- ☐ radio buttons (select one option only)
- ☐ check box (multiple selections allowed)

---

### Pulmonary Medications

**Antibiotics – inhaled and/or oral**

**Tobramycin Based Medications**

**Tobramycin solution for inhalation** (i.e. TOBI):
- Frequency: ☐
  - 300 mg BID alternate month schedule ☐
  - Other regimen (different dose or freq) ☐
  - Eradication ☐

**Tobi Podhaler** (Tobramycin Inhalation Powder):
- Frequency: ☐
  - 4.2mg capsules BID alternate month ☐
  - Other regimen (different dose or freq) ☐
  - Eradication ☐

**Bethkis**:
- Frequency: ☐
  - 300 mg BID alternate month ☐
  - Other regimen (different dose or freq) ☐
  - Eradication ☐

*repeated entries can be recorded
[ ] indicates values calculated by the registry
2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

Other inhaled aminoglycoside (e.g. gentamicin, amikacin, or tobramycin preparation): ☐
Frequency:
☐ Alternate Month
☐ Continuous
☐ Other regimen (different dose or freq)
☐ Eradication

Colistin: ☐
Frequency:
☐ Alternate Month
☐ Continuous
☐ Other regimen (different dose or freq)
☐ Eradication

Aztreonam – Inhaled: ☐
Frequency:
☐ 75 mg TID Alternate Month Schedule
☐ 75 mg TID Continuous
☐ Other Regimen
☐ Eradication

Other inhaled antibiotics: ☐
Comments:___________________________

Oral macrolide antibiotic: ☐
☐ azithromycin (Zithromax)
☐ clarithromycin (Biaxin)

Other oral antibiotic: ☐
☐ Quinolone (Cipro, Levaquin, gatifloxacin, etc.)
☐ Cephalosporin (cephalexin, Keflex, cefixime, etc.)
☐ Sulfam (Bactrim, Septra, etc.)
☐ Amoxicillin (Augmentin, etc.)
☐ Tetracycline (doxycycline, Vibramycin, minocycline, etc.)
☐ Other

CFTR Modulators
Ivacaftor Monotherapy (i.e. Kalydeco): ☐
Frequency:
☐ 50 mg BID
☐ 75 mg BID
☐ 150 mg BID
☐ Other Regimen (different dose or freq)

Ivacaftor/Lumacaftor Combination Therapy (i.e. Orkambi): ☐
Frequency:
☐ Full dose BID
☐ Half dose BID
☐ Other Regimen (different dose or freq)

Tezacaftor/Ivacaftor Combination Therapy: ☐
Frequency:
☐ Full dose BID
☐ Half dose BID
☐ Other Regimen (different dose or freq)

Elexacaftor/Tezacaftor/Ivacaftor Triple Combination Therapy: ☐
Frequency:
☐ Full dose BID
☐ Half dose BID
☐ Other Regimen (different dose or freq)

Other Medications
Dornase alfa (i.e. Pulmozyme): ☐
Frequency:
☐ 2.5 mg QD
Key:
☐ 2.5 mg BID
☐ Other regimen (different dose or frequency)

Bronchitol (Mannitol): ☐
Frequency:
☐ 400 mg BID
☐ Other regimen (different dose or frequency)

Acetylcysteine or Mucomist: ☐
High-dose ibuprofen (e.g. 25-30 mg/kg): ☐
Total (mg/dose): ______
Hypertonic saline: ☐
Concentration (%): 3 4 5 6 7 8 9 10
Frequency: ☐ QD ☐ BID ☐ Other

Bronchodilators (oral): ☐
☐ Beta agonist (e.g. Proventil Repetabs, Volmax, etc.)
☐ Theophylline product (e.g. Theodur, Slo-bid, Uniphyl)

Bronchodilators (inhaled):
☐ Short acting beta agonist (e.g. albuterol, Proventil, Ventolin, Xopenex, etc.)
☐ Long acting beta agonist (e.g. salmeterol, Serevent, Foradil, Brovana, etc.)
☐ Short acting anticholinergic (e.g. ipratropium, Atrovent)
☐ Long acting anticholinergic (e.g. tiotropium, Spiriva, etc.)
☐ Combination beta agonist and anticholinergic (e.g. Combivent, DuoNeb, etc.)

Corticosteroids:
☐ Oral (e.g. prednisone)
☐ Inhaled (e.g. fluticasone, Flovent, budesonide, Pulmicort, etc.)
☐ Inhaled in combination with a bronchodilator (e.g. Advair, Symbicort)

Other:
☐ Leukotriene modifiers (e.g. montelukast, Singular, zafirlukast, Accolate, zileuton, Zyflo, etc.)
☐ Mast cell stabilizers (e.g. cromolyn, Intal, nedocromil, Tilde, etc.)
☐ Antifungals (e.g. itraconazole, Sporanox) Note: exclude topical agents for skin conditions and agents used for oral thrush)

Drug Intolerance/Allergies:
☐ Dornase alfa (i.e. Pulmozyme)
☐ TOBI or other aminoglycoside
☐ Aztreonam
☐ Colistin
☐ Macrolide antibiotics
☐ High-dose ibuprofen
☐ Hypertonic saline
☐ Ivacaftor (i.e. Kalydeco)
☐ Ivacaftor Lumacaftor (i.e. Orkambi)
☐ Tezacaftor/Ivacaftor (i.e. Symdeko)

GI/Nutrition/Endrocrine Medications
This Patient is on enzyme medications: ☐ Yes ☐ No
For all enzymes, “capsules per largest meal” options are: .5 .1 .2 .3 .4 .5 .6 .7 .8 .9
*repeated entries can be recorded
[ ] indicates values calculated by the registry
2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

**Enzymes**

**Creon**

Creon 1203: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Creon 1206: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Creon 1212: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Creon 1224: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Creon 1236: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

**Pancreaze**

Pancreaze MT4: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Pancreaze MT10: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Pancreaze MT16: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Pancreaze MT20: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

**Ultresa**

Ultresa 14: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Ultresa 20: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Ultresa 23: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

**Pertzye (Pancrecarb)**

Pertzye 4000: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Pertzye 8000: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Pertzye 16000: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Pertzye 24000: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

**Zenpep**

Zenpep 3: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Zenpep 5: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Zenpep 10: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Zenpep 15: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Zenpep 20: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Zenpep 25: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Zenpep 40: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

**Viokace**

Viokace 10: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

Viokace 20: ☐
- Number of capsules per largest meal of the day: ____
  - Total capsules per day: ____

**Other Enzymes**

Please specify if other enzymes: ______________________

**Acid Blocker**

Acid Blocker (Daily use. Check all that apply since last visit):
- ☐ H2 Blocker (e.g. Zantac, Pepcid, etc.)
- ☐ Proton Pump Inhibitor (e.g. Prilosec, Nexium, etc.)
- ☐ Unknown

**GI other**

Ursodeoxycholic acid: ☐

**Pulmonary**

**Pulmonary Function Tests (PFTs)**

Unable to Perform test: ☐
- Reason why PFTs have not been done: ___________

FVC measure (L): ___________
  - [Predicted value: _______ ]
  - [Reference equation: _______ ]
  - [% Predicted: _______ ]
  - [Relative change since previous measurement: _______ ]
  - [Days since last measured: _______ ]

*repeated entries can be recorded*  
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## 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

### FEV1 measure (L): 
- Predicted value:_____  
- Reference equation:_______  
- [% Predicted:_______ ]  
- [Relative change since previous measurement:_____ ]  
- [Days since last measured:____ ]

### FEF25-75 measure (L/sec): __________
- Predicted value:_______  
- Reference equation:_______ 
- [% Predicted:_________ ]

### CF Specific FEV1 percentile (ages 6-21): 

### GI/Nutrition

#### Assessment of Oral Intake:
- ○ Done  ○ Not done

Is patient currently receiving supplemental feeding?
- ○ Yes  ○ No  ○ Unknown

**Feeding:**
- ☐ oral supplementation (Scandishakes, Pediasure, Instant Breakfast, etc.)
- ☐ nasogastric tube (NG)
- ☐ gastrostomy tube/button (G-Tube)
- ☐ jejunal tube (J-tube)
- ☐ total parenteral nutrition (TPN)

If using a tube (NG, G-tube or J-tube), was the patient recommended to use pancreatic enzymes with supplemental feedings?
- ☐ No enzymes recommended with tube feeding
- ☐ Yes enzymes mixed into the formula
- ☐ Yes, enzymes administered directly through the tube (i.e. not into the formula)
- ☐ Yes, enzymes taken by mouth prior to, during and/or after the feeding
- ☐ Yes, formula infused through Relizorb (enzyme cartridge)
- ☐ Other

**CF specific vitamins (i.e. with additional vitamins A, D, E, and K):**
- ○ Yes  ○ No

#### Infants under 2 years of age

**Salt supplementation:**
- ○ Yes  ○ No

Select type of feeding:
- ○ Breast milk
- ○ Formula exclusively
- ○ Other food
- ○ Unknown

If receiving any formula feeding, select type of formula and caloric density:
- ○ Cow's milk
- ○ Soy milk
- ○ Predigested
- ○ Other

**Caloric Density:**
- ○ 20 cal/oz
- ○ 22 cal/oz
- ○ 24 cal/oz
- ○ 27 cal/oz
- ○ 30 cal/oz
- ○ Other, specify:_____________

### Complications

**Patient does not have any complications:**
- ☐

### Diabetes Status
- ○ Impaired Glucose Tolerance (FBG < 126, 2-h PG 140-199)
- ○ CFRD with or without fasting hyperglycemia
- ○ Type 1 Diabetes
- ○ Type 2 Diabetes

**CFRD secondary complications:**
- ☐ Retinopathy
- ☐ Microalbuminuria
- ☐ Chronic renal insufficiency
- ☐ Chronic renal failure requiring dialysis
- ☐ Peripheral neuropathy

### Hepatobiliary
- ☐ Gall stones
- ☐ Gall stones, requiring surgery/procedure
- ☐ Liver disease, cirrhosis

**Please specify complications related to cirrhosis:**
- ☐ Esophageal varices
- ☐ Gastric varices
- ☐ GI bleed related to varices
- ☐ Splenomegaly
- ☐ Hypersplenism (i.e., WBC <3.0 or platelets <100,000)
- ☐ Ascites
- ☐ Encephalopathy
- ☐ Liver disease, non-cirrhosis
- ☐ Acute Liver Failure (No underlying liver disease, ALT>3X ULN, INR>2, not responsive to vitamin K)
- ☐ Hepatic Steatosis
- ☐ Liver disease, other: _____________

**Acute Hepatitis (ALT > 5X ULN and duration of illness < 6 months):**
- ☐ Infectious (Hepatitis A,B,C,EBV,CMV or other known infectious cause)
- ☐ Non-infectious (Autoimmune, Drug Induced, Alcohol or other known cause)
- ☐ Unknown

### Bone/Joints
- ☐ Arthritis/Arthropathy
- ☐ Bone fracture
- ☐ Osteopenia
- ☐ Osteoporosis

### Pulmonary
- ☐ Allergic Bronchial Pulmonary Aspergillosis (ABPA)
- ☐ Asthma
- ☐ Hemoptysis

**Please specify selection of hemoptysis:**
- ☐ Hemoptysis, massive
- ☐ Hemoptysis, other
- ☐ Pneumothorax requiring chest tube

### GI
- ☐ Distal intestinal obstruction syndrome (DIOS, Meconium

---

*Key:*
- ○ radio buttons (select one option only)
- ☐ check box (multiple selections allowed)

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2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

FORM NAME

Radio buttons (select one option only)

Check box (multiple selections allowed)

Key:

- ileus equiv.
- Fibrosing colonopathy/colonic stricture (report incidence only)
- GERD (Gastro-Esophageal Reflux Disease)
- GI Bleed req hosp non variceal
- History of intestinal or colon surgery
- Pancreatitis
- Peptic ulcer disease
- Rectal prolapse

Other Complications
- Absence of Vas Deferens
- Anxiety Disorder
- Cancer confirmed by histology
- Depression
- Hearing loss
- Hypertension
- Kidney Stones
- Nasal polyps requiring surgery
- Renal failure requiring dialysis (cause other than CFRD)
- Sinus Disease (symptomatic)

Complications not listed above
Enter additional complications:

Lab

Blood counts
- WBC count x1,000/microL (typical clinical value: 3.0 to 30.0):_____
- Platelet Count x1,000/microL (typical clinical value: 100 to 500):_____
- Hemoglobin (grams per deciliter):_____

Serum Creatinine
- Serum Creatinine Level (mg/dL):_____

Liver Function Tests (LFTs)
- Alanine Aminotransferase (ALT or SGPT), IU/L:_____
- GGTP (gamma glutamyl transpeptidase), IU/L:_____
- Aspartate Aminotransferase (AST), IU/L:_____
- Alkaline phosphatase (ALP), IU/L:_____
- Total Bilirubin, mg/dL:_____

Glucose Test
- Random blood glucose (mg/dL):_____
- Fasting blood glucose (mg/dL):_____

If OGTT performed:
- OGTT Fasting glucose level (mg/dL):_____
- 1 hour (mg/dL)(not required):_____
- 2 hour (mg/dL):_____

Hemoglobin A1C (Hgb A1C)
- Hgb A1C value, %:_____

Fecal Elastase
- Fecal Elastase Value (microg/g of stool):_____

Act/Exercise

Primary Airway Clearance Technique (ACT)
- Positive Expiratory Pressure (PEP)
- Postural drainage with clapping (CPT)
- Forced expiratory techniques (e.g. autogenic drainage, huff cough, active cycle breathing)
- Oscillating PEP (e.g. Flutter, acapella, IPV)
- High frequency chest wall oscillation (e.g. Vest)
- Exercise
- None
- Other
- Specify if other technique:_________

Secondary Airway Clearance Technique (ACT)
- Positive Expiratory Pressure (PEP)
- Postural drainage with clapping (CPT)
- Forced expiratory techniques (e.g. autogenic drainage, huff cough, active cycle breathing)
- Oscillating PEP (e.g. Flutter, acapella, IPV)
- High frequency chest wall oscillation (e.g. Vest)
- Exercise

CARE EPISODE

Care Episode Segment*

Start date: (MM/DD/YYYY)
End date: (MM/DD/YYYY)
Location: ○ Hospital ○ Home IV
Reasons:
- Pulmonary Exacerbation
- Pulmonary Complication Other than exacerbation
- GI Complications
- Transplant related
- Sinus infection
- Non-transplant surgery
- NTM Pulmonary Infection
- Other
- Please specify reason:_________

Care Episode Measurements

At the beginning of Care Episode:
- FVC (L):_____
- FEV1 (L):_____
- FEF25-75 (L):_____

Height: ____ cm ____ inches
Weight: ____ kg ____ lb
Date recorded: (MM/DD/YYYY)
Check if data were impossible to measure:□

At the end of Care Episode:
- FVC (L):_____
- FEV1 (L):_____
- FEF25-75 (L):_____

Height: ____ cm ____ inches
Weight: ____ kg ____ lb
Date recorded: (MM/DD/YYYY)
Check if data were impossible to measure:□

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### 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

Comments:___________________________

<table>
<thead>
<tr>
<th>ANNUAL REVIEW</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual Review Year: (YYYY)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Patient Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Encounters recorded by Center: [ ]</td>
</tr>
<tr>
<td>Number of Encounters recorded by other Care Centers: [ ]</td>
</tr>
<tr>
<td>Number of Care Episodes recorded by Care Center: [ ]</td>
</tr>
<tr>
<td>Number of Care Episodes recorded by Other Care Centers: [ ]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Demographics Update</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current Zip:_________</td>
</tr>
<tr>
<td>Patient is: [alive or dead]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pulmonary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Did this patient use oxygen therapy during the reporting year?</td>
</tr>
<tr>
<td>o Yes, Continuously</td>
</tr>
<tr>
<td>o Yes, Nocturnal and/or with exertion</td>
</tr>
<tr>
<td>o Yes, During exacerbation</td>
</tr>
<tr>
<td>o Yes, pm</td>
</tr>
<tr>
<td>o No</td>
</tr>
<tr>
<td>o Unknown</td>
</tr>
</tbody>
</table>

| Did this patient use non-invasive ventilation during the reporting year (i.e., assisted breathing, BiPap, CPAP, etc) |
| o Yes |
| o No |
| o Unknown |

| Was a Chest X Ray performed during the reporting year? |
| o Yes |
| o No |
| o Unknown |

| Did the patient receive an influenza vaccination this season (Sept through Jan)? |
| o Yes |
| o No |
| o Unknown |

<table>
<thead>
<tr>
<th>Covid-19 Vaccination</th>
</tr>
</thead>
<tbody>
<tr>
<td>Did the patient receive a Covid-19 vaccination this year?</td>
</tr>
<tr>
<td>o Yes (Please complete Covid-19 Vaccination form)</td>
</tr>
<tr>
<td>o No</td>
</tr>
<tr>
<td>o Unknown</td>
</tr>
</tbody>
</table>

| Specify why vaccination wasn't done: |
| o Ineligible by Age |
| o Allergic |
| o Patient refused vaccination |
| o Other |

| Specify other reasons why vaccination wasn't done: ________ |

<table>
<thead>
<tr>
<th>Mycobacterial Culture</th>
</tr>
</thead>
<tbody>
<tr>
<td>[According to the encounters a Mycobacterial culture has been performed during this reporting year: o Yes o No ]</td>
</tr>
</tbody>
</table>

| Please check to confirm the above is correct: □ |

| Was treatment INITIATED for a pulmonary mycobacterial infection during this reporting year? |
| o Yes |
| o No |
| o Unknown |

| Was an IgE screening for ABPA performed in this reporting year? |
| o Yes |
| o No |
| o Unknown |

<table>
<thead>
<tr>
<th>Key: FORM NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>o radio buttons (select one option only)</td>
</tr>
<tr>
<td>□ check box (multiple selections allowed)</td>
</tr>
</tbody>
</table>

| Did this patientsmoke cigarettes during the reporting year? |
| o No |
| o Occasionally |
| o Yes, Regularly, less than 1 ppd |
| o Yes, Regularly, 1 ppd or more |
| o Declined to answer |
| o Not Known |
| o Not Applicable |

| Does anyone in the patient’s household smoke cigarettes? |
| o Yes o No o Unknown |

| During the reporting year, how often was this patient exposed to secondhand smoke? |
| o Daily |
| o Several Times Per Week |
| o Several Times Per Month or less |
| o Never |
| o Declined to answer |
| o Not Known |

<table>
<thead>
<tr>
<th>Liver</th>
</tr>
</thead>
<tbody>
<tr>
<td>[According to the encounters data liver function tests were done in this reporting year o Yes o No ]</td>
</tr>
</tbody>
</table>

| Please check to confirm that information about liver function tests above is correct. If it is incorrect, please return to the encounter forms and enter correct information into the lab section of the encounter form: □ |

<table>
<thead>
<tr>
<th>Eye</th>
</tr>
</thead>
<tbody>
<tr>
<td>Was any eye exam performed to check for cataracts in this reporting year?</td>
</tr>
<tr>
<td>o Yes</td>
</tr>
<tr>
<td>o No</td>
</tr>
<tr>
<td>o Unknown</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Growth and Nutrition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fat soluble vitamin levels measured?</td>
</tr>
<tr>
<td>o Yes</td>
</tr>
<tr>
<td>o No</td>
</tr>
<tr>
<td>o Unknown</td>
</tr>
</tbody>
</table>

| Has this patient been on growth hormone in the reporting year? |
| o Yes |
| o No |
| o Unknown |

| Was a DEXA scan for bone density performed in the reporting year? Please enter findings of osteoporosis or osteopenia into the complications section of last patient encounter. |
| o Yes |
| o No |
| o Unknown |

| Results of DEXA Scan: |
| o Normal |
| o Osteopenia |
| o Osteoporosis |
| o Other |
| o Unknown |

<table>
<thead>
<tr>
<th>Diabetes Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Status from recent encounter [does or does not] indicate CFRD.</td>
</tr>
<tr>
<td>o Normal Glucose Metabolism (includes normal, random, fasting, or OGTT)</td>
</tr>
<tr>
<td>o Impaired Glucose Tolerance (FBG &lt; 126, 2-h PG 140-199)</td>
</tr>
<tr>
<td>o CFRD with or without fasting hyperglycemia (2-h PG &gt;= 200)</td>
</tr>
<tr>
<td>o Type 1 Diabetes</td>
</tr>
</tbody>
</table>

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2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

○ Type 2 Diabetes

Was a retinal eye exam performed by an ophthalmologist in this reporting year? ○ Yes ○ No ○ Unknown
Was a spot urine sent for albumin/creatinine ratio in this reporting year? ○ Yes ○ No ○ Unknown
If positive findings, please update CFRD secondary complications in the most recent encounter of the patient.

Was the patient prescribed treatment for CFRD? ○ Yes ○ No
Select all that apply:
□ Dietary change
□ Oral hypoglycemic agents
□ Intermittent insulin (with illness, steroids, etc.)
□ Chronic insulin

Did the patient experience any episodes of severe hypoglycemia (became unconscious or required help to resolve) during the reporting year? ○ Yes ○ No ○ Unknown

Transplantation
What is the transplantation status of the patient currently? If the patient had transplantation in previous years please select or keep "Had transplantation" option.
○ Not pertinent
○ Accepted, on waiting list
○ Evaluated, final decision pending
○ Evaluated, rejected
○ Had transplantation

Transplant
□ Lung: Bilateral
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
□ Heart/Lung
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
□ Lung: Lobar/Cadaveric
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
□ Lung: Lobar/living donor
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
□ Liver
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
□ Kidney
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
□ Other
   Number this year: ___ Date of last transplant: (MM/DD/YYYY)
Specify transplant type: ________

Were there post transplant complications? □
Select those that apply:
□ Bronchiolitis obliterans syndrome
□ Lympho-proliferative disorder
□ Other
   Specify other complication: ________________________

Colorectal Cancer Screening/Surveillance
Did the patient undergo a colonoscopy (screening or surveillance during the reporting year)?
Key: ○ Yes ○ No ○ Unknown
What were the results of the colonoscopy?
○ Normal
○ Colorectal Cancer
○ Adenomatous polyps
○ Indeterminate results (e.g. inadequate preparation)

Clinical Trials
Has this patient participated in any interventional (drug) studies? ○ Yes ○ No ○ Unknown
Has this patient participated in any observational studies? ○ Yes ○ No ○ Unknown

Health Insurance Coverage
It is important for us to have accurate numbers of patients who have specific types of coverage:
□ Health Insurance Policy (e.g. Private Insurance)
□ Medicare
□ Medicaid
□ State special needs program, e.g., BCN, CCS, CRS, GHPP, etc.
□ TriCare or other military health plan
□ Indian Health Service
□ Other
   Specify if other insurance: ________________________

Patient has no health insurance: □

Was patient covered under parent’s health insurance plan? ○ Yes ○ No ○ Unknown
Did patient receive free medicine or co-pay/deductible assistance from a Patient Assistance Program? ○ Yes ○ No ○ Unknown

Socio-economic Status
Education of Patient:
○ Less than High School
○ High School diploma or equivalent
○ Some College
○ College Graduate
○ Masters/Doctoral level degree
○ Unknown/Not applicable

Education of father of patient:
○ Less than High School
○ High School diploma or equivalent
○ Some College
○ College Graduate
○ Masters/Doctoral level degree
○ Unknown/Not applicable

Education of mother of patient:
○ Less than High School
○ High School diploma or equivalent
○ Some College
○ College Graduate
○ Masters/Doctoral level degree
○ Unknown/Not applicable

Education of spouse of patient:
○ Less than High School
○ High School diploma or equivalent
○ Some College
○ College Graduate
○ Masters/Doctoral level degree
○ Unknown/Not applicable

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## 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

### Key:
- ○ radio buttons (select one option only)
- □ check box (multiple selections allowed)

#### What was the total combined income of the household before taxes where the patient resided for the majority of the reporting year?
- ○ $<10,000
- ○ $10,000 to $19,999
- ○ $20,000 to $29,999
- ○ $30,000 to $39,999
- ○ $40,000 to $49,999
- ○ $50,000 to $59,999
- ○ $60,000 to $69,999
- ○ $70,000 to $79,999
- ○ $80,000 to $89,999
- ○ $90,000 or more

#### How many people currently live in the patient’s household (including the patient)?
- ○ 1
- ○ 2
- ○ 3
- ○ 4
- ○ 5
- ○ 6
- ○ 7
- ○ 8
- ○ 9
- ○ 10
- ○ 11
- ○ 12 or more
- ○ Unknown

#### Mental Health

- Was the patient screened for symptoms of classic depression using Patient Health Questionnaire (PHQ-9) or other valid depression screening tools?
  - ○ Yes
  - ○ No
  - ○ Unknown

- Was the patient screened for the anxiety disorder using Generalized Anxiety Disorder Tool (GAD-7 or similar)?
  - ○ Yes
  - ○ No
  - ○ Unknown

#### Age 18 and Older

- Marital Status:
  - ○ Single (never married)
  - ○ Living Together
  - ○ Married
  - ○ Separated
  - ○ Divorced
  - ○ Widowed
  - ○ Unknown

- Employment:
  - ○ Part Time
  - ○ Full time homemaker
  - ○ Full time employment
  - ○ Unemployed
  - ○ Student
  - ○ Disabled
  - ○ Retired
  - ○ Unknown

- Pregnancy
  - Was patient pregnant during the reporting year?
    - ○ Yes
    - ○ No
    - ○ Unknown
  
  - If Yes, indicate outcome:
    - ○ Live Birth

#### Other

- Please use this field to record any additional information about this patient: _______________

### ALD INITIATION

**Date patient flagged for ALD: (MM/DD/YYYY)**

Select all reasons for flagging ALD:
- ○ Stable FEV1pp<40%
- ○ Hemoptysis
- ○ Pneumothorax
- ○ Rapid FEV1 decline
- ○ Respiratory failure
- ○ Supplemental Oxygen at rest
- ○ Pulmonary Hypertension
- ○ 6 mins walk distance <400m
- ○ Hypercapnia
- ○ Other

Specify other reasons for including patient: _______________

### ALD ANNUAL

**Interventions**

Review Year:

**Thoracic Procedure(s)**

- Did patient have a thoracic procedure during the reporting year?
  - ○ Yes
  - ○ No
  - ○ Unknown

- Thoracic procedure type(s):
  - ○ Lung resection
  - ○ Thoracotomy
  - ○ Video thoracotomy
  - ○ Tube thoracotomy
  - ○ Pleurodesis

- Pleurodesis side:
  - ○ Unilateral
  - ○ Bilateral
  - ○ Unknown

### Other Interventions

*repeated entries can be recorded

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2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

Was patient enrolled in pulmonary rehabilitation during the reporting year?
- Yes
- No
- Unknown

Was the patient seen by a palliative care specialist during the reporting year?
- Yes
- No
- Unknown

Specify type(s) of palliative care services:
- Documented advance care planning conversation
- Documentation of an advance directive
- Palliative care consultation
- Referral to hospice
- Referral to outpatient/community palliative care

Embolization For Hemoptysis
Embolization Date (if performed): (MM/DD/YYYY)

ICU Admissions
Date of admission to ICU: (MM/DD/YYYY)
Date of discharge from ICU: (MM/DD/YYYY)
Primary reason for admission:
- Post-surgical procedure with observation
- Respiratory failure
- Hemoptysis
- Pneumothorax
- Shock
- Densitization for antibiotics
- Other
- Unknown

Intubation
Was patient intubated?
- Yes
- No
- Unknown

Date of intubation: (MM/DD/YYYY)
Date of extubation: (MM/DD/YYYY)

ECMO/Novalung
Was patient placed on ECMO/Novalung?
- Yes
- No
- Unknown
Date ECMO/Novalung started: (MM/DD/YYYY)
Date ECMO/Novalung ended: (MM/DD/YYYY)
Cannulation strategy used:
- VV-Single Cannula
- Code 2VV-Double Cannula
- Veno-Arterial
- Veno-Arterial-Venous
- Unknown

Was ECMO/Novalung used in combination with ventilator?
- Yes
- No
- Unknown

Labs/Tests
Echocardiogram
Was ECHO performed during the reporting year?
- Yes
- No
- Unknown

Date of last echocardiogram: (MM/DD/YYYY)
Could PA pressure be estimated?
- Yes
- No
- Unknown

Estimated RVSP or PASP, mmHg: ___________________
Tricuspid annular plane systolic excursion, mm: ____________

Right Heart Catheterization (RHC)
Was Right Heart Catheterization (RHC) done during the reporting year?
- Yes
- No
- Unknown

Date of last RHC: (MM/DD/YYYY)
Mean PA pressure from RHC, mmHg: ________________
Systemic systolic pressure, mmHg: ________________
Systemic diastolic pressure, mmHg: ________________
Heart Rate: _______________________

PA systolic pressure, mmHg: __________________________
PA diastolic pressure, mmHg: __________________________
Pulmonary capillary wedge pressure (PCW), mmHg: ______
Cardiac Index (L/min/m²): __________________________

Six Minute Walk
Date of six-minute walk test: (MM/DD/YYYY)
Six minute walk distance, m: _______________________
Was supplemental oxygen titrated?
- Yes
- No
- Unknown

Blood Gas Analysis
Date of blood gas analysis: (MM/DD/YYYY)
Blood source:
- Venous
- Arterial
Type of blood gas analysis:
- Stable blood gas
- Worst blood gas in a year
- Last blood gas prior to discharge
Blood pH: ________________________
PaO2, mm Hg: ______________________
PaCO2, mm Hg: ______________________

Transplant Consideration and Referral
Transplant Consideration
Was transplant discussed with patient at the care program?
- Yes
- No
- Unknown

Transplant Referral And Evaluation
Was patient referred to a transplant program?
- Yes
- No
- Unknown

If patient was not referred for transplant, select reasons:
- Patient stable
- Patient declined to pursue
- Substance misuse (alcohol, marijuana, illicit drugs etc.)
- Chronic opioid use
- Smoking tobacco
- Inadequate medical insurance
- Limited social support
- Adherence issues
- Poor nutritional status
- Microbiological reasons
- Mental health
- Uncontrolled diabetes
- Physical inactivity/deconditioning

*repeated entries can be recorded
[ ] indicates values calculated by the registry
### 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

#### Primary referral program:
- ☐ Other
  Specify other reasons for not referring: __________________

#### Date patient referred for transplant: (MM/DD/YYYY)

#### State of primary referral program:
- ☐ AL
  - University of Alabama Hospital
- ☐ AZ
  - Banner University Medical Center – Tucson
  - Mayo Clinic
  - St Joseph’s Hospital and Medical Center
- ☐ CA
  - Cedars-Sinai Medical Center
  - Children's Hospital Los Angeles
  - Keck Hospital of USC
  - Lucile Salter Packard Children’s Hospital at Stanford
  - Stanford Health Care
  - University of California San Diego Medical Center
  - University of California San Francisco Medical Center
  - University of California at Los Angeles Medical Center
- ☐ CO
  - University of Colorado Hospital/Health Science Center
- ☐ FL
  - Florida Hospital Medical Center
  - Jackson Memorial Hospital University of Miami School of Medicine
  - Mayo Clinic Florida
  - Tampa General Hospital
  - UF Health Shands Hospital
- ☐ GA
  - Emory University Hospital
- ☐ IA
  - University of Iowa Hospitals and Clinics Transplant Programs
- ☐ IL
  - Advocate Christ Medical Center
  - Loyola University Medical Center
  - Northwestern Memorial Hospital
  - University of Chicago Medical Center
- ☐ IN
  - Indiana University Health
- ☐ KY
  - Jewish Hospital
  - University of Kentucky Medical Center
- ☐ LA
  - Ochsner Foundation Hospital
- ☐ MA
  - Boston Children’s Hospital
  - Brigham and Women’s Hospital
  - Massachusetts General Hospital
- ☐ MD
  - Johns Hopkins Hospital
  - University of Maryland Medical System
- ☐ MI
  - Henry Ford Hospital
  - Spectrum Health
  - University of Michigan Medical Center
- ☐ MN
  - Saint Marys Hospital (Mayo Clinic)
  - University of Minnesota Medical Center, Fairview

#### Key:
- ☐ radio buttons (select one option only)
- ☐ check box (multiple selections allowed)

#### Checklist:
- ☐ MO
  - Barnes-Jewish Hospital
  - St Louis Children’s Hospital at Washington University Medical Center
- ☐ NC
  - Duke University Hospital
  - University of North Carolina Hospitals
- ☐ NE
  - The Nebraska Medical Center
- ☐ NJ
  - Newark Beth Israel Medical Center
- ☐ NY
  - Mount Sinai Medical Center
  - NY Presbyterian Hospital/Columbia Univ Medical Center
- ☐ OH
  - Children’s Hospital Medical Center
  - Nationwide Children’s Hospital
  - Ohio State University Medical Center
  - The Cleveland Clinic Foundation
  - University Hospitals of Cleveland
- ☐ OK
  - Integris Baptist Medical Center
- ☐ PA
  - Allegheny General Hospital
  - Children’s Hospital of Pennsylvania
  - Children’s Hospital of Pittsburgh of UPMC
  - Hospital of the University of Pennsylvania
  - Temple University Hospital
  - University of Pittsburgh Medical Center
- ☐ SC
  - Medical University of South Carolina
- ☐ TN
  - Baptist Memorial Hospital
  - Vanderbilt University Medical Center and Nashville VA Medical Center
- ☐ TX
  - Baylor University Medical Center
  - CHI St. Luke’s Health Baylor College of Medicine Medical Center
  - Houston Methodist Hospital
  - Memorial Hermann Hospital, University of Texas at Houston
  - Scott and White Memorial Hospital
  - Texas Children’s Hospital
  - UT Southwestern Medical Center/William P Clements Jr University Hospital
  - University Hospital, University of Texas Health Science Center
  - University of Texas Medical Branch at Galveston
- ☐ UT
  - University of Utah Medical Center
- ☐ VA
  - Inova Fairfax Hospital
  - University of Virginia Health Sciences Center
- ☐ WA
  - University of Washington Medical Center
- ☐ WI
  - Aurora St. Luke’s Medical Center
  - Froedtert Memoria Lutheran Hospital
  - University of Wisconsin Hospital and Clinics

#### Was patient evaluated for transplant at primary referral program?
- ☐ Yes
- ☐ No
- ☐ Unknown

*repeated entries can be recorded*

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### 2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

If patient evaluated for transplant at primary referral program, specify outcome:
- [ ] Listed
- [ ] Delisted
- [ ] Denied
- [ ] Pt declined to pursue
- [ ] Unknown

Date listed for transplant: **(MM/DD/YYYY)**

If transplant deferred, specify reason:
- [ ] Patient stable
- [ ] Additional testing necessary
- [ ] Poor nutritional status
- [ ] Limited social support
- [ ] Financial hardship
- [ ] Substance misuse
- [ ] Other

Other reason(s) for transplant deferral: ______________________

If patient was denied for transplant, specify reason:
- [ ] Too sick
- [ ] Microbiological reasons
- [ ] Limited social support
- [ ] Substance misuse
- [ ] Malignancy
- [ ] Poor nutritional status
- [ ] Renal disease
- [ ] Liver disease
- [ ] Financial hardship
- [ ] Other, specify

Other reasons patient denied for transplant: ______________________

Was pt referred to another program for second opinion?
- [ ] Yes
- [ ] No
- [ ] Unknown

State of second referral program:
- [ ] Second referral program:
  - [ ] AL
  - [ ] AZ
  - [ ] CA
  - [ ] CO
  - [ ] FL
  - [ ] GA
  - [ ] IA
  - [ ] IL
  - [ ] IN
  - [ ] KY
  - [ ] LA
  - [ ] MA
  - [ ] MD
  - [ ] MI
  - [ ] MN
  - [ ] MO
  - [ ] NC
  - [ ] NE
  - [ ] NJ
  - [ ] NY
  - [ ] OK
  - [ ] PA
  - [ ] SC
  - [ ] TN
  - [ ] TX
  - [ ] VA
  - [ ] WV
  - [ ] WI
  - [ ] WY

Key:
- [ ] radio buttons (select one option only)
- [ ] check box (multiple selections allowed)

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2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

- Vanderbilt University Medical Center and Nashville VA Medical Center
- Baylor University Medical Center
- CHI St. Luke’s Health Baylor College of Medicine Medical Center
- Houston Methodist Hospital
- Memorial Hermann Hospital, University of Texas at Houston
- Scott and White Memorial Hospital
- Texas Children’s Hospital
- UT Southwestern Medical Center/William P Clements Jr University Hospital
- University Hospital, University of Texas Health Science Center
- University of Texas Medical Branch at Galveston
- Vanderbilt University Medical Center
- UT Medical Center
- VA Inova Fairfax Hospital
- University of Utah Medical Center
- Inova Fairfax Hospital
- University of Virginia Health Sciences Center
- University of Washington Medical Center
- Aurora St. Luke’s Medical Center
- Froedtert Memoria Lutheran Hospital
- University of Wisconsin Hospital and Clinics

If patient evaluated for transplant at second program, specify outcome:
- Listed
- Delisted
- Denied
- Pt declined to pursue
- Unknown

COVID-19

- Nasopharyngeal swab (antigen test)
- Testing Result:
  - Positive
  - Negative
  - Inconclusive
- Test Comments: ___________________________________

Please describe confirmed or suspected event:

Did patient have any symptoms related to COVID-19 at presentation?
- Yes, symptomatic
- No, patient was asymptomatic

Date of first symptoms: (MM/DD/YYYY)

Signs and Symptoms at manifestation:
- Cough, different from CF
- New shortness of breath or difficulty breathing
- Chills
- Fever (>=38C)
- Headache
- Muscle pain
- New loss of taste or smell
- Repeated shaking with chills
- Other

Other symptoms, not included above: __________________

Treatment

Where was the patient treated for Covid-19 symptoms?
- Self-quarantine at home
- Admitted to hospital
- Other

Specify other treatment location: ______________________

Date admitted to hospital: (MM/DD/YYYY)

Make sure to create a care episode if a patient was hospitalized.

Was patient admitted to ICU?
- Yes
- No

Was patient on supplemental oxygen?
- Yes
- No

Did the patient require non-invasive respiratory support (CPAP, biPAP; High-flow)?
- Yes
- No

Did the patient require mechanical ventilation?
- Yes
- No

Did the patient require ECMO support?
- Yes
- No

Did the patient require treatment therapies?
- Yes
- No
- Unknown

Specify if the following therapies were used to treat the patient:
- Antiviral therapy
- Antibiotics
- Systemic steroids
- Immune modulators
- Other

Other Therapies: __________________________________

Complications/Co-morbidities

COVID-19 associated complications/comorbidities (e.g. stroke, secondary infection): _______________

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2020 Cystic Fibrosis Foundation Patient Registry Questionnaire

Please be sure to enter or update patient data in the appropriate tabs of an encounter form (e.g. complications, labs). For any data entry questions, contact reghelp@cff.org.

Follow-up
Has patient recovered?
○ Yes
○ Currently Hospitalized
○ Currently in ICU
○ Currently in domestic quarantine
○ Deceased
○ Unknown

Date of recovery (i.e. end of symptoms OR negative COVID-19 test): (MM/DD/YYYY)

COVID-19 Vaccination

Covid-19 Vaccination
Please enter the date of first vaccination shot, even if it is approx.: (MM/DD/YYYY)
Please enter the date of second vaccination shot, if known and required: (MM/DD/YYYY)
The patient received a vaccine developed by:
○ Pfizer
○ Moderna
○ AstraZeneca
○ Novavax
○ Johnson&Johnson
○ Other or Unknown
Specify other vaccine manufacturer, if known: ____________

Key:

- FORM NAME
  ○ radio buttons (select one option only)
  [ ] check box (multiple selections allowed)
  *repeated entries can be recorded
  [ ] indicates values calculated by the registry
REFERENCES


