DANVER —

Patients Over 40 Once Unheard-of Now a Growing Segment of the Population

Cystic fibrosis has been transformed in recent decades from a uniformly fatal childhood disease to a condition with middle-aged and older patients. A comprehensive analysis of cystic fibrosis patients over 40 by researchers at National Jewish Health highlights patients who have survived with the disease for decades and an emerging population of patients diagnosed as adults.

The results, published online May 20 in the American Journal of Respiratory and Critical Care, show that the majority of patients diagnosed as adults are females, which represents a striking reversal of the "gender gap" in cystic fibrosis. The researchers also discovered that these patients have delayed, but equally severe disease, and enjoy improved lung function once they begin receiving proper care.

"Our analysis offers insight into the status, disease progression and outcomes of middle-aged and older cystic fibrosis patients," said National Jewish Health pulmonologist and lead author Jerry Nick, MD. "It may help guide the treatment of cystic fibrosis as these patients become increasingly common."

Cystic fibrosis (CF), an inherited disease of the lungs and digestive system, affects about 30,000 people in the United States and is the most common genetic disease among Caucasians. A single defective gene causes a missing or non-functional channel for chloride to travel into and out of cells. Lack of this channel disrupts the water balance in the lungs of CF patients, causing the development of thick, dehydrated mucus, which serves as a fertile environment for bacterial growth. Most patients die of respiratory failure brought on by repeated severe bacterial infections in the lungs.

In 1962 the median predicted survival for children with cystic fibrosis was 10 years. Today, it is 37, and children diagnosed today can expect to live into their 50s. Improved care is helping patients live longer. In addition, many more patients with non-traditional symptoms are being diagnosed for the first time as adults.

These two groups - those diagnosed as children and those diagnosed for the first time as adults - comprise two distinct populations of CF patients over 40. Those diagnosed as adults usually have a genetic mutation that produces a partially functional gene and a slightly different set of symptoms from those diagnosed as children. Most commonly, they have functional pancreases and fewer digestive problems.

Dr. Nick and his colleagues analyzed epidemiological and health data on 156 CF patients over 40 year of age who receive care at National Jewish Health, the largest adult cystic fibrosis clinic in the nation. In addition, data were analyzed on nearly 3,000 patients from around the nation who were included in the Cystic Fibrosis Foundation Patient Registry from 1992-2007.

The researchers found that the fate of females changes considerably in the older CF population. It has long been recognized that a "gender gap" is present in CF, favoring males. Historically, females have been diagnosed later, had a poorer prognosis, and survived fewer years than males.
Accordingly, Dr. Nick’s analysis showed that fewer females diagnosed as children survived to age 40. However, among those diagnosed as adults, females represented a significant majority, accounting for 72 percent of patients in Colorado and 54 percent nationally. Among the adult diagnosed patients, females survived on average 9 to 14 years longer than males.

The complex factors that account for the differential fate of female CF patients is not understood, although Dr. Nick believes it could be a mixture of behavioral and biological factors.

Dr. Nick’s findings also indicate that patients diagnosed as adults do not really have milder diseases -- as is commonly believed -- just a delayed onset of an equally severe form of the disease. Although patients diagnosed as adults live longer than those diagnosed as children, the adult-diagnosed patients lose lung function as rapidly those diagnosed in childhood, and approximately 85% die of respiratory failure or post-transplant complications.

Dr. Nick believes there is a significant number of adults whose CF remains undiagnosed. His analysis indicates that once those patients are accurately diagnosed, proper care can significantly improve their health. Patients diagnosed as adults and subsequently followed at a CF center reversed progressive lung function decline and improved their lung function for at least four years.

Older patients commonly do not get specialized CF care. It is generally recognized that the team approach to treatment provided by the 112 CF Foundation-accredited Care Centers results in better clinical outcomes. However, less than half of long-term CF survivors continued to be seen at CF Centers as they pass 40 years, with the fewest among the adult-diagnosed patients.

"In the coming years, more and more cystic fibrosis patients will be living into their 40s, 50s and beyond," said Dr. Nick. "Our findings concerning the role of gender, in survival, progression of disease, and type of care in current long-term survivors provides important insights that will help us prepare for better treatment of the steadily aging CF population."

National Jewish Health is the leading respiratory hospital in the nation. Founded 125 years ago as a nonprofit hospital, National Jewish Health today is the only facility in the world dedicated exclusively to groundbreaking medical research and treatment of children and adults with respiratory, cardiac, immune and related disorders. Patients and families come to National Jewish Health from around the world to receive cutting-edge, comprehensive, coordinated care. To learn more, visit the media resources page.

Media Resources

We have many faculty members, from bench scientists to clinicians, who can speak on almost any aspect of respiratory, immune, cardiac and gastrointestinal disease as well as lung cancer and basic immunology.

- Accomplishments & Awards
- Annual Report
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