

## Cystic Fibrosis

### I. Background and History of Cystic Fibrosis Research

Cystic Fibrosis is a disease caused by genetic mutations, affecting more than 300 million people worldwide, that causes a buildup of mucus in the alveoli of the lungs (CF Foundation). This disease is so detrimental because viscous mucus coats the alveoli, severely obstructing breathing and preventing the exchange of oxygen and carbon dioxide in the lungs at times. Figure 1, obtained from the Mayo Clinic, shows the difference between the alveoli of an uninfected person versus someone who suffers from Cystic Fibrosis (Figure 1). Cystic Fibrosis, abbreviated CF, is usually diagnosed in patients under two years of age, because the lifespan of someone with CF is only about forty years (CF Foundation). The symptoms for CF include persistent coughing episodes, shortness of breath, buildup of phlegm, salty tasting skin, poor weight gain, male infertility, and frequent *Staphylococcus aureus*, *Haemophilus influenzae*, and *Pseudomonas aeruginosa* infections, (CF Foundation). These infections occur so frequently because the warm, viscous mucus forms a sustainable environment for bacterial colonies to grow. Amoxicillin or Erythromycin can cure the bacterial infections, but there is no known genetic cure for CF.

Cystic Fibrosis was originally thought to be caused by bacterial infections alone, until the CF gene was discovered on human chromosome seven in 1989 (CF Foundation). Early research led to the

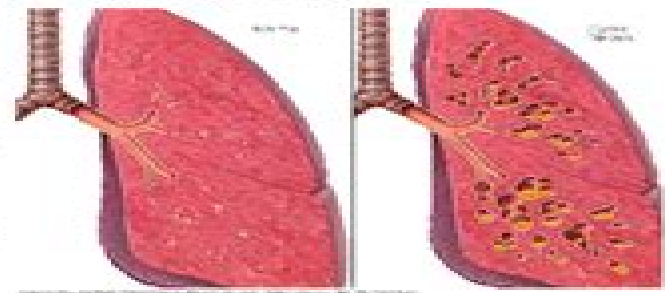


Figure 1 - Diagram of normal lungs versus lungs that are filled with mucus due to Cystic Fibrosis (CF Foundation).

# Cystic Fibrosis Research Paper Thesis

**Christian G. Meyer**



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