Inflammation in the lungs of patients diagnosed with Cystic Fibrosis: Association with iron deficiency

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Cystic Fibrosis (CF) is characterized by a wide spectrum of phenotypic characteristics such as; deep coughing, increased mucous production, and weight loss. However, only recently was the role of inflammation on the etiology of the disease recognized. CF is characterized as a cyclic progression of infective exacerbations and stable periods initiated by the presence of Pseudomonas Aeruginosa (PA). An increase in inflammatory cytokines/mediators and a decrease in anti-inflammatory cytokines contribute to the net inflammation and overall tissue destruction of the lungs. PA is associated with the low iron status that is seen in 60-75% of the CF population, through the presence of iron sequestering siderophores which distract iron from the tissues. Iron deficiency (ID) initiates further symptoms such as; fatigue, tachycardia, weakness, brittle nails etc, in addition to those caused by CF. The colonization of PA may be the cause or a result of increased iron (ferritin) concentrations in the lungs, but independent of the original relationship, results in a decreased iron status. Iron is used by PA under hypoxic conditions such as in the fibrosis lung, as a source of energy. Studies on the relationship between CF and ID contribute a variety of possible causes although currently no direct connection has been discovered. At this stage, further studies in this area are needed. This review will primarily focus on the affects of CF on iron status in humans, and secondarily examine the effect of mediators of inflammation in respects to ID.

Cystic Fibrosis (CF) is a genetic disease (autosomal recessive) that occurs in childhood and results in premature death in the early adult years. It is the most common lethal recessive disease and currently no cure has been discovered. CF is a multi-system disease that presents as a spectrum of phenotypes that affect both the respiratory and gastrointestinal systems. It can be traced to over a thousand different mutations to the cystic fibrosis transmembrane conductance regulator (CFTR). A mutation of the CFTR gene alters the function of the epithelium, which results in the production of abnormally thick mucous. The blockage of airways as a result of the mucous may result in a decrease in mobilization of iron and therefore an increase in stores.

Even the slightest increase in iron is enough to initiate enough bacterial growth to overwhelm the immune system. Iron is seen increased in CF and initiates the growth and development of bacterial infections such as Pseudomonas Aeruginosa (PA). The colonization of bacteria (i.e. PA) results in recurrent episodes of chronic infection and inflammation throughout the lifespan, as these bacteria utilize iron sources in the lung through iron sequestering siderophores. An increase in the metabolism of iron in the epithelium results in an increased production of free hydroxyl-radicals, which damages airways, further stimulating the production of mucous, initiating a vicious cycle.

CF is an airway obstructive disorder that progresses towards the development of a hypoxic environment within the lungs. As a result the body initiates polycythaemia, which is an attempt to increase haemoglobin and hematocrit concentrations in the blood as a method of increasing bound $O_2$. The development of new blood cells is known as erythropoiesis and may result in ID, as it removes iron from other regions of the body.

ID or anaemia is defined as a transferrin saturation <16% in the blood and is commonly found in 60%-75% of the CF population, even in patients that meet the recommended dietary intake (RDI) of iron. ID remains a problem even in patients that meet the recommended dietary intake (RDI) of iron, those consuming supplementation, and in those consuming a diet low in iron. Symptoms such as anaemia, tachycardia, fatigue, pale skin, headaches, increased work of breathing (WOB) and weakness are all characteristic of ID. These symptoms in addition to symptoms of CF leave patients in a compromised state of well being.
Iron is an essential nutrient to the growth and development of CF patients. It is involved in various biological processes such as the synthesis of proteins, enzymes, and DNA, the storage and transport of iron, and the production of heme. In CF, the increased colonization of PA plays both a direct and indirect role in the progressive destruction of lung tissue. PA normally develops in CF children around the age of 6, and is responsible for periodic episodes of exacerbation throughout the lifespan. Under normal conditions, the lung maintains iron levels as close to zero as possible. If iron concentrations alter from this set point, there is an overwhelming response colon/vorization by pathogenic bacteria which use iron as an energy source for growth and respiration. PA has the ability to secrete iron-binding granules known as siderophores, which increase available iron in the lungs. Patients with CF have an unusually high concentration of free iron in their lungs compared to controls, and this may be a result of increased vascular permeability to the serum, increase in airflow TNF-α (decreases iron re-absorption), airway inflammation (accumulation of iron in airways), an increase in cell death (oxidative damage), or the pseudomonas siderophores.

Patients with both non-exacerbated and infective exacerbated symptoms of CF both have an increase in available iron concentration. Infective exacerbated symptoms include things such as: deep coughing, increased sputum concentrations, breathlessness, weight loss, and are experienced many times during the lifespan of patients with CF. During exacerbations patients are also found to have increased concentration of PA bacteria. As was stated above even the smallest increase in free iron is enough to initiate the colonization of PA. The increase in mucous found in CF leaves the lungs in a state of hypoxia, which means that bacteria must adapt their metabolism to anaerobic conditions. PA has adapted to hypoxia by developing branched aerobic respiratory side chains, termed 5-terminal-oxidases. For successful metabolism the PA is unique in that only one active chain is required. The side chains have a high affinity for O2, and allow for PA to flourish in hypoxic conditions. Secondarily, polycythemia is used in the lung as a response to a decrease in O2 concentrations in the blood. Polycythemia is an overproduction of red blood cells in an attempt to increase O2 delivery throughout the body. Patients with an infective bout of CF may have increased risk of ID due to increases in erythropoiesis. Erythropoiesis maxes out all other stores of iron available in an attempt to increase production of Hb. However, when tested this hypothesis proved incorrect. Patients with CF did have a decreased O2 saturation (SaO2%) but haemoglobin and hematocrit values did not differ significantly from control patients. Therefore erythropoiesis was not an improving factor in patients with CF; an over accumulation of red blood cells did not balance hypoxemia, and cannot be the reason for a decrease in iron status in CF.

Reid et al. collected sputum samples from patients with acute infective exacerbations, stable CF patients and control patients and analyzed total iron, ferritin, and tumour necrosis factor (TNF-α), both prior to and following a 14 day antibiotic treatment. Patients with the infective exacerbations showed significantly higher levels of total sputum cell count (TCC), iron, ferritin, and TNF-α than control patients. A significant association was also found between concentrations of TNF-α and iron homeostasis. Sputum iron and ferritin levels, 44.4 μmol/L and 3.6 mg/L, respectively exceeded normal ranges for iron (13-32 μmol/L) and ferritin (15-300 μmol/L). Reid also found a significant portion of the population was infected with PA, which most
likely contributed to the increase in iron measured. This data suggests that the increase in iron may act as a fuel for PA, increase disease pathogenesis, and lead to an increased severity of respiratory failure.

The increase in PA increases TNF-α released from macrophages and epithelial cells. Therefore research is inconclusive of whether the increase in iron seen in CF is an effect of increased PA, or whether the presence of PA increases available iron.

Levels of PA and iron content of the CF lung were recently found to be closely associated in CF lungs (Figure 1). Patients infected with PA had levels of sputum iron and ferritin of 59.73 µmol/l and 4002.20 mg/L, respectively. Patients who were diagnosed with CF but were not infected by PA also had increased concentrations of sputum iron and ferritin when compared to a control population, 29.1 µmol/l and 2972.30 mg/L. Therefore the increase in sputum iron occurs prior to infection by PA and therefore changes in iron must predate PA (Figure 1). The initial increase in iron may be a result of vascular leakage due to inflammation, although the direct connection remains unknown. This data suggests that the initial increase in iron may be responsible for the initial colonization of PA. The main proportion of extra-cellular iron found in CF lungs is in the form of ferritin. Iron in the lungs cannot be strictly supplied by damaged epithelial cells, as they contain very low concentrations of ferritin. It must therefore be derived from another source, which is supported by Stites et al.’s work, but contraindicated by research conducted by Reid et al., that suggested epithelial apoptosis was the main source of iron in the lungs. Iron bound to ferritin is more easily accessed then that attached to transferrin, which means that the increase in ferritin and the decrease in transferrin in the lungs influences the etiology of this disease. Reasons for the decrease in transferrin are unknown but there is reason to believe that it is cleaved by PA derived proteases.

Ferritin is a host binding molecule that readily releases ferric (Fe3+) iron in the presence of siderophores. PA is highly effective at obtaining ferric iron from its environment and increasing its solubility. Therefore an increase in ferritin stores will result in an increased free iron, as a result of siderophore digestion. Siderophores in the sputum of patients with CF bind to the iron repressible outer membrane of the cell, initiating ferric ion assimilation. When there were increased concentrations of PA, as seen during exacerbations, there was also an increase in siderophore presence.

Increased concentrations of two specific siderophores, pyoverdine and pyochelin, were found in patients with CF. The increase in siderophores produced by PA demonstrates the bacteria’s attempt to absorb iron for survival. In summary, siderophores are scavengers of iron that incorporate proteins into the outer membranes of the bacteria, allowing for the siderophores complex to bind and absorb iron more readily.

PA contributes both indirectly and directly to ID. The direct mechanisms involves removing iron from the airways as a source of energy through siderophore sequestering, whereas the indirect mechanism involves increasing cytokine production which, in turn increases iron storage and detracts iron from Haemoglobin (Hb) synthesis (Figure 1). Previous research felt removal of PA was impossible, whereas recent research has used chemical and/or physical therapy to reduce exacerbation caused by PA. Recent intervention strategies are used to treat the bacteria (the cause) instead of treating the inflammation (the effect). PA is detrimental to the body as it causes oxidative damage, inflammation, infections and uses up iron for metabolism that the body requires.

**Inflammation of the Lungs and Neutrophil Concentrations**

The mucous secretions of CF block the airways, resulting in not only hypoxic conditions as discussed above, but also a decreased re-absorption of iron, and endobronchial
inflammation and/or infection. The concentration of neutrophils in the sputum is a common marker of inflammation. Neutrophil concentrations are found to be significantly higher in patients with CF when compared with healthy subjects, 74% and 8%, respectively. Neutrophils are an essential aspect of the autoimmune response, as they defend the body against foreign particles attempting to infect the body, however in CF, excess concentrations of neutrophils are damaging. Neutrophils fight infection by inhibiting the spread of micro-organisms through phagocytosis and by increasing the concentration of pro-inflammatory cytokines and proteases.

When present in increased concentrations, neutrophils are associated with an increased production of proteases, such as elastase, which contributes to the development of fibrosis of the lungs. Proteases freely digest the elastin found in the airway walls, decreasing their elastic nature and increasing rigidity. The increase in elastase has secretagogue activity, which increases the production of mucous. The increase in PA associated with the increase in mucous, acts as a chemo-attractant to neutrophils (Figure 2). When neutrophils are recruited to a site of infection, the lifespan of inflammatory cytokines is increased (Figure 2). An increase in iron also contributes to the alterations in neutrophil behaviour.

The severity of inflammation in the lungs is also affected by errors in apoptosis and macrophage production. Macrophages are used in the process of removing dead neutrophils from the lungs. If neutrophils have an extended lifespan due to an error in apoptosis, the result is increased recruitment of inflammatory cytokines and a decrease in recruitment of anti-inflammatory cytokines. Pro-inflammatory cytokines concentrations are increased in CF. IL-8 is significantly higher in patients with PA (145.5ng/ml) than in those with no bacterial content (13.2ng/ml). An increase in IL-8 is related to an increase in neutrophil concentration and vice versa (Figure 3).

The increase in inflammation associated with CF is one of the main initiators of increased concentration of free and stored iron (ferritin) found in the lungs. Inflammation was found to be correlated with an increased PA population, which in turn affected iron concentrations. Inflammation independent of cause disrupts the homeostasis of iron in patients with CF.

Cytokine studies in general

Patients with CF in Nixon et al. study were compared to control patients, before and after antibiotic treatments and were found to have increased concentrations of IL-6, TNF-α, and CRP. Therefore patients diagnosed with anaemia or iron deficiency in CF have an association with increased concentrations of cytokines in their sputum or BAL. Cytokine presence is one of the largest contributors to anaemia in CF, as they cause red blood cell death and a decreased erythropoietin production. IL-6 and TNF-α also initiate trapping of iron by macrophages, and therefore aid in the promotion of ID by inhibiting the absorption of iron at the level of the duodenum. A decreased absorption inhibits the production of Hb, as it depends on iron as its central molecule.

Cytokines are produced from a variety of cells such as: airway, lung interstitial, air space or vascular components. They are produced in response to a variety of stimulants: hypoxia, neutrophils, increase in bacteria, and are activated by T and B lymphocytes. This occurs by lymphocytes binding foreign particles such as those that have been attacked by macrophages, and initiating further production of the T cells, B cells and cytokines. The following sections will further examine the role of individual cytokines and related inflammatory mediators in CF.

![Figure 2: Relationship between P. Aeruginosa, Neutrophil Concentration and Pro-Inflammatory Cytokine Production](image)

![Figure 3: Cell regulation of Neutrophil Concentration and Inflammatory Cytokines in the Presence of PA in CF lungs](image)
TNF-α and IL-8, Role in Respiratory Inflammation

TNF-α and IL-8 are the two major inflammatory cytokines produced by the airway epithelium and macrophages as a direct response to PA and indirectly to inflammation and infection. Both TNF-α and IL-8 play an essential role in the over recruitment of neutrophils to damaged areas of the body. The body produces TNF-α as a protective mechanism against PA by removing iron similar to IL-6 (as will be discussed below) however, the overwhelming concentrations result in an increase in damage to the lungs instead.

Concentrations of TNF-α were found increased in patients with CF. TNF-α was found at concentrations of (448pg/ml) whereas control samples concentrations were found on average to be much lower (9pg/ml). TNF-α reduces the availability of iron by diverting it away from Hb synthesis. It also recruits iron from other areas of the body by phagocytosis, increasing available free iron in the lungs and decreasing iron with in the rest of the body. Reid et al. found a positive correlation between a decrease in both TNF-α and ferritin. ID is directly related to the severity of lung disease, and was therefore found in increased concentrations in patients with exacerbated symptoms when compared to normal patients.

Reid et al found that an increase in TNF-α in the sputum of patients was closely correlated to a decrease in iron stores and an increase in sputum iron. TNF-α is responsible for up-regulation of cellular ferritin mRNA synthesis, which increases deposits of iron in the tissues of the lungs, providing a reservoir for PA. In support of these findings in a later study, Reid found a similar relationship, which was an increase in free iron in sputum of the lungs and an increase in TNF-α and IL-8. TNF-α was also found to correlate with IL-1β.

IL-6 a Pro & Anti Inflammatory Cytokine

IL-6 has both pro- and anti-inflammatory functions. As an anti-inflammatory it initiates negative feedback that decreases TNF-α and IL-1β production (potent inflammatory cytokines) and plays a pro-inflammatory role through stimulation of the liver to produce Acute Phase Proteins (APP) such as C-reactive protein (CRP). Increased concentrations of IL-6 are also utilized in the trapping of iron by macrophages, and play a role in the development of ID. Patients with CF had a higher concentration of IL-6 (8.7 +/- 0.4 pg/ml), in comparison to control patients who had significantly lower concentrations (2.6 +/- 0.1 pg/ml). As well, patients infected with PA had higher levels (9.3 +/- 0.3pg/ml) then CF patients infected with another stream of bacteria (6.9 +/-0.5pg/ml). However in the CF lung, IL-6 seems to play a pro-inflammatory role due to an increase in TNF-α.

Both IL-6 and LTB₄ concentrations significantly decreased after patients with PA were treated with antibiotics for two weeks. Similarly, Nixon et al. found that patients with CF had increased levels of IL-6 of (7.28pg/ml) compared to the control group which measured at (0.65pg/ml). After a 14 day treatment plan, patients with CF showed significant decreases in IL-6 (2.1 pg/ml) whereas the control group showed no significant changes. These studies both found IL-6 to play an inflammatory role. Carpagnano et al. compared a group of patients and analyzed BAL whereas Nixon et al. looked at less patients and measured concentrations in sputum. Cytokine concentrations present different characteristics depending on their origin in the body. More research is consequently needed in this area.

In contrast to the above studies Osika et al. found decreased IL-6 concentrations in patients with CF in comparison to healthy subjects and suggested that IL-6 had anti-inflammatory characteristics. This study measured the sputum concentration of IL-6 in a similar population to that in Carpagnano et al. and therefore is of similar quality of research. This study found that patients with CF had levels of IL-6 of (26 pg/ml) whereas the control group had levels of (225pg/ml). Although Osika et al. found that there was no significant difference between CF patients who were infected with PA, and patients with CF that were not, which conflicts with results found in most other studies. IL-6 as an anti-inflammatory cytokine aids in the disruption of homeostasis found between pro and anti-inflammatory cytokines in these patients. Therefore the increase in inflammation contributed to by changes in IL-6 activity increase the stability of PA and therefore further increase ID.

Anaemia is a common characteristic of inflammatory diseases such as CF. Inflammation in the body results in the recruitment of a large numbers of inflammatory cytokines that mediate the immune response. All the factors relating to ID such as: shortened red-cell survival, a decreased erythropoietin response to anaemia and decreased erythropoietin colony formation, can be related to the presence of these substrates. These cytokines attempt to remove any available iron from the PA, but as a result decrease concentration throughout the body instead. What initially is meant to be a protective mechanism during acute infection consequently harms the body during chronic disease.

Inflammatory Mediator; C - Reactive Protein

CRP is an APP that increases proportionally with IL-6. Increased concentrations of IL-6 stimulate the production of CRP in the liver. Inflammation initiates an increase in recruitment of IL-6, which results in increased CRP released in the serum and sputum. Mechanistically CRP binds to foreign or damaged particles and removes them from the body by actively binding and promoting the activity of macrophages. CRP aids in the development of ID by inducing the trapping of iron in macrophages, and decreasing its absorption as it stimulates macrophage activity. CRP also initiates cytokine induced hepatic expression of hepcidin,
which is produced in the gastrointestinal tract and inhibits the absorption of iron in the duodenum.\textsuperscript{12} The highest concentrations of CRP were recorded during exacerbation, but after 14 days of antibiotic treatment CRP levels were found to level off.\textsuperscript{27} Increased values of CRP result in a decrease in available iron.\textsuperscript{12}

Fischer et al. (2007) found that 40\% of CF patients presented with high levels of CRP, (0.5mg/dl) compared with normal values of (0.22mg/dl). After 3 and 6 months 60\% and 67\% of the group, respectively, had an increase in CRP. These patients had two times the concentration of CRP than the control patients. 88\% of the patients which had presented originally with non-exacerbated symptoms experienced an exacerbated infection as a result of PA by the end of the 12-month study,\textsuperscript{12} which also demonstrates the regular occurrence of exacerbations throughout the lifespan. As a result, a correlation is seen between CRP and infection. During periods of infection CRP levels were higher then in the non-exacerbated CF lung.\textsuperscript{12}

Nixon et al. found that CRP levels within each individual showed no increase over the 14 day study although consistent with Fischer et al. CRP values were greater in patients with CF than in the control group.\textsuperscript{23, 12} Both Nixon et al. and Fischer et al. found that once CRP levels reach their maximum, an increase in infection is no longer proportional to CRP. \textsuperscript{23, 12} Exacerbated CF patients began the trial with CRP levels of 21.32ug/ml but after treatment showed a significant decrease to 2.75ug/ml whereas before and after treatment control patients showed no significant change 0.33µg/ml to 0.34µg/ml.\textsuperscript{23}

**IL-10, an Anti-inflammatory Cytokine**

The airways of patients affected with CF are relatively deficient in the anti-inflammatory cytokine IL-10.\textsuperscript{3, 18, 25} IL-10 was recorded at (24pg/ml) in patients with CF whereas control subjects recorded a significantly lower average concentration of (45pg/ml).\textsuperscript{25} Sputum measurements in the control population were induced by treating subjects by induction of 200µg of inhaled salbutamol, as sputum is not produced under healthy conditions.\textsuperscript{25} IL-10 is an inhibitory cytokine that acts on IL-8 and TNF-α (pro-inflammatory). Unlike CRP which decreased when exacerbation was complete, IL-10 levels remain low in all stages of illness. Therefore low concentrations of IL-10 allow for the exaggerated effect of inflammatory cytokines IL-8 and TNF-α and indirectly contribute to ID and an increase inflammatory damage to the lung.\textsuperscript{3, 18}

**Nitric Oxide a Marker of Airway Inflammation**

Over the last ten years a number of studies have focused on nitric oxide (NO) and its role in pulmonary disease. NO is a marker of airway inflammation and is found in the EBC of individuals with chronic respiratory disease.\textsuperscript{13} NO is released from activated macrophages, neutrophils and infected lung epithelial cells, by stimulation of T and B immune cells.\textsuperscript{15} NOS, is the enzyme that is responsible for the proliferation of NO and is positively influenced by the presence of PA, and TNF-α, which are seen increased in CF.\textsuperscript{13, 16} Therefore, in individuals with increased NO, an increase in NOS can be assumed.\textsuperscript{13} Nitrite (NO\textsubscript{2}⁻) and Nitrate (NO\textsubscript{3}⁻) are the two stable end products produced by NO metabolism and are found in increased concentrations in CF.\textsuperscript{16} NO: NOS like other inflammatory factors contributes to ID through activating an increase in neutrophils, macrophages and damaged epithelial cells.\textsuperscript{13, 26} As a response to the infiltration of bacterial cultures, the body responds by increasing NO.\textsuperscript{26} Again, CF is a cyclic disease, and an increase in inflammation will further increase concentrations of PA resulting in increased utilization of iron, contributing to the overall ID in CF patients.

Patients with CF unexpectedly present with low levels of NO and high levels of NOS\textsuperscript{1, 13, 16, 26} Ho et al. hypothesized that due to the increase in mucous in the airways, NO is trapped close to the aqueous membrane, which allows for complete metabolism into NO\textsubscript{2} and NO\textsubscript{3}.\textsuperscript{16} When NO is trapped neighbouring the aqueous membrane of the epithelial cells, a reaction occurs with reactive oxygen species or with water and oxygen, stabilizing the molecule. Ho et al. found that NO\textsubscript{2} levels were significantly higher in patients with CF (2.15µm versus control 0.36µm) whereas NO showed no significant difference between the two populations (3.8 versus control 5.2 ppm).\textsuperscript{16} Nitrate levels which are much lower in CF, maybe due to its possible metabolic role in PA respiration.\textsuperscript{3} Under hypoxic conditions, nitrate may be used as a substitute of oxygen.\textsuperscript{1} It is therefore most likely that CF results in an increased production of NO but most of it is converted to bi-products.\textsuperscript{16, 22} The exact reasoning remains unknown as to whether it is an increase in NO metabolites which there is research to support or a decrease in the production of NO.\textsuperscript{13}

Bacteria which utilize iron as a source of energy may be responsible for the decrease seen in NO and hence the abnormal symptoms that are seen in CF. Consequently, the greater the inflammation, the more PA that can thrive on available iron and the larger the amount of NO that is metabolized by the PA to NO\textsubscript{2}. Possible research may be focused on developing a correlation between concentrations of NO\textsubscript{2} and a decrease in iron levels in the blood.

**Iron Deficiency in the Gastrointestinal (GI) tract**

Although this paper focuses on the decrease in iron at the respiratory level it will briefly touch on the effects of ID in the GI system. Davis & Biggs’ paper, which is original literature on the relationship between the pancreas and its role in iron absorption, found that a decrease in the function of the pancreas causes a decrease in iron absorption.\textsuperscript{8} Davis & Biggs’ study, however, was performed on rats, which must be considered when applying the findings to humans. When ferrous sulphate (FeSO\textsubscript{4}) was injected into the isolated and perfused region of the rat’s small intestine (pH of 1), the
mean value of iron absorption was 51%, whereas when the same dosage of iron was given with pancreatic extract, iron absorption decreased to 29%. It was concluded that pancreatic secretions play some role in the absorption of iron in the jejunum, and are associated with chronic CF, however, it is important to note that pancreatic secretions are only produced in CF once severe pancreatic damage has occurred. The natural pH of the jejunum in rats is roughly 6.4 whereas a human’s natural intestinal pH ranges between 6.63-7.4, which means that the slight differences between the data sets must be considered, and must give way to confirmation by further studies performed on human subjects (in vivo). In conjunction with these findings, research by Pond et al. also found that iron absorption is inhibited by an increase in volume of mucous on the pancreas. When pancreatic enzymes are inhibited, a decrease in iron absorption can be observed, eventually leading to ID, which can also be observed. For this reason, pancreatic extract, as a result of mucous production, resulting in a decreased absorption of iron, has been suggested to play an inhibitory role.

Recently, Fischer et al. tested patients for exocrine pancreatic insufficiency and found that all patients were limited in their ability to secrete pancreatic enzymes. Problems with this study included set backs such as looking at a small number of patients (at no fault to the researcher as CF patients are hard to find for testing), as well as using single measurements upon which to base comparisons, as these, in combination, may have been inaccurate when assessing overall health of subjects. This study also found that even patients who were taking iron supplementation prior to the study had iron levels below cut off. These findings suggested that ID is caused by an inability to absorb nutrients, and is also not the result of a dietary deficiency. Recent research, in contrast, has found evidence to support the hypothesis that that the degree of ID may be unrelated to pancreatic insufficiency, although further research is needed into this area. Reid et al. found that even when patients were given pancreatic supplements, iron levels showed no improvement. Absolute reasons for these observations have yet to be discovered and may also have some relation to other GI factors.

CONCLUSION AND FUTURE RESEARCH

Individuals with CF present a multitude of phenotypic mutations, along with significant sources of illness, which are together associated with an increase in mucous excretion by the lungs and pancreas. The increased mucous results primarily in a blockage of airways, producing a hypoxic environment, and results secondarily in an increase in inflammation. Research is unclear as to whether inflammation is the source of the colonization of PA, or whether PA is the reason behind the body’s inflammatory response. Independent of the cause, both inflammation and PA result in a decrease in iron and ferritin in both the blood and throughout the body. One of the areas of future research should be finding whether a direct connection between the role of NO, as an inflammatory marker, and ID is present. As well, further studies geared to adjusting the balance between the increase in pro-inflammatory, and decrease in anti-inflammatory cytokines and mediators, are required to ultimately improve health and wellness within patients. Research should emphasize the importance of treating PA and the sources of inflammation, as opposed to treating symptoms, not only to increase patients' lifespan, but more importantly, to enrich their quality of life.

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