Gastro-Esophageal Reflux and Pulmonary Medicine; Where Are We and What Should We Do?

Mostafa Ghanei\textsuperscript{a} and Amin Saburi

Chemical Injury Research Center, Baqiyatallah University of Medical Sciences, Tehran, I.R. Iran

There is growing concern about relation between Gastro-Esophageal Reflux Disease (GERD) and pulmonary diseases. It seems that researches which are focused on pulmonary effects of GERD are now being enhanced during the past decades. GERD is one of the commonest gastrointestinal disorders which is affecting more than one/fourth of adults in western countries [1]. Recently, the prevalence of GERD is rising especially in developing countries. The same entities of these evidences support this theory that the non-acidic components have the same embryonic origin [2]. Accordingly, the same originality of airways and esophagus can explain the pathogenesis of both GERD and pulmonary related disorders. Also, there are evidences about the relation between airway narrowing and esophageal distention due to intra-esophageal hydrochloric acid provocation via vague nerve [2,3]. Accompanying between these two diseases especially in incidences and responses to the treatment enhance this hypothesis.

The role of gastric especially acidic contents in pulmonary complication of GERD has been identified well. Increase in exhaled oxidative stress biomarkers (e.g., 8-isoprostane), neuroinflammation increases in vagal tone and PH metery findings in especially symptomatic patients are considered as some evidence in the pathogenic role of acid reflux in GERD [4].

Aspiration of non acidic gastric contents such as pepsin, pepsinogen, pancreatic enzymes and bile derivates have been also reported to play a role in pulmonary symptoms of asthma [5,6]. Increasing in some pro-inflammatory cytokine such as IL-8 was detected in bronchial epithelial cells of patients who are under acid suppressive treatment. Treatment with non-acidic components such as PPI in patients who are under acid suppressive treatment may induce inflammatory responses of lung to GERD. Although reaction to the microbial agents or endotoxins in the gastric content may even induce the pulmonary inflammatory responses, it has not been yet identified which part of the gastric content has more significant influence on the respiratory epithelium [7]. Asthma, chronic cough, bronchitis, bacterial pneumonia and interstitial lung fibrosis are well determined as pulmonary consequences of GERD [1].

However still some questions have been remained to be answered in this filed before designing an effective therapeutic protocol. Traditional treatments of GERD reduce acid related symptoms in these patients although the injury of non-acidic reflux can be continued and consequently the symptoms persist. Regarding to the failure of traditional anti-reflux medication in many cases, new therapeutic agents should be considered in future studies. New generation of anti-reflux medications such as MGlur5 antagonists and γ-Amino butyric acid agonists have been recommended which is affecting the transient lower esophageal sphincter relaxation that is the basic underlying cause of reflux [8]. On the other side, GERD is regularly diagnosed and treated only in symptomatic stage. However the "silent" form of disease which can be associated with microaspiration usually is neglected.

The low effectiveness of current remedies to suppress gastric acidity even with powerful proton pump inhibitors on extra-esophageal symptoms of GERD may suggest that reflux can exert its effects through mechanisms beyond excessive acid production. Non acidic component reflux was presented as a justifying mechanism for extra-esophageal symptoms of patients who were remained unresponsive to acid suppressive treatment [9]. Symptomatic relief in response to surgical repair of lower esophageal sphincter that targets the basic mechanism of reflux as a motility disorder can support this idea [10]. The symptoms of high acid reflux can be controlled by acid suppressive treatment but the Pathophysiology remains intact. Accordingly, we need to an effective marker or device to evaluate the efficacy of treatment especially in non-acidic reflux although bile salts or pepsin of sputum was faintly introduce as diagnostic biomarkers for GERD. Also, the neural reflex between the esophagi sensitizes and cough reflex in patients with GERD and chronic cough was previously described [11]. Therefore, Even if the aspiration (micro or macro) is controlled in patients with reflux, the symptoms may not relief although microaspiration can be difficult to control and monitor.

Bronchiolitis obliterans is the main cause of lung transplant rejection and the microaspiration has been implicated as the latent causative factor leading to Bronchiolitis obliterans in this patients [12].

Also, we need studies which evaluate the efficacy of microaspiration treatment on the extra-pulmonary manifestations of GERD to confirm the role of any degree of reflux in pathogenesis of GERD in lung tissue. In addition, there are no methods for following up of treatment’s efficacy and it can be proposed for future studies. There are some biomarkers increased in respiratory system of patients with GERD which it can be useful for this issue. For example substance P and neurokinin A are increasing in response of inflammatory process at the respiratory epithelium [13]. Salivary or sputum pepsin and pepsinogen level can be measured for this issue as well. It seems that replacing PH metery with another diagnostic and confirming method is helpful to achieve to more accurate diagnosis in non-acidic gastro esophageal reflux.

References


*Corresponding Author: Prof. Mostafa Ghanei, Pulmonologist, Chemical Injury Research Center, Baqiyatallah University of Medical Sciences, Mollasadra St, Vanak Sq, Tehran, I.R. Iran, Tel/Fax: +98-21-8860067; E-Mail: mghaneister@gmail.com

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