Activity of alpha amylase in tracheobronchial secretions of patients without morbid salivary aspiration

Abstract

Background: The normal value of alpha-amylase in humans is unknown.

Objective: To determine the normal values of alpha-amylase activity in humans.

Material and methods: From October 2009 to June 2011 we studied 111 patients referred to thoracic service to be submitted to bronchoscopy. The patients were positioned in supine position, performed local antisepsis and anesthetized with 2% lidocaine. Thereafter we introduced a needle into catheter by puncturing the cricothyroid membrane using a 14G cateter. Finally we introduced ten milliliters of saline and immediately aspirated with the maximum power of the vacuum system. The samples were sent for alpha-amylase activity determination by CNPG method.

Results: The activity of alpha-amylase of tracheal aspirate ranged from 24 to 10.000 IU/L), and a mean 1914IU/L. The levels had no statistical differences according age, sex, race, smoking history and the lung diseases.

Conclusion: we could define the probably physiologic levels of amylase activity in human beings.

Keywords: alpha-amylase, lung, salivary aspiration, aspiration pneumonia, transtracheal puncture

Introduction

There is no reference in previously published material on the normal value of alpha-amylase in tracheobronchial secretions. The amylase in the lungs of humans under normal conditions probably has two components-one is from local production and the other from physiological micro aspiration. Takano in 1938 demonstrated that the blood of theirs right heart had major amylase concentration than the left chambers, which the plausible explanation is the passage of amylase from lung circulation to general circulation. Sano studying activity of enzymes in lung tissue of humans by histochemical techniques found that the enzyme most frequently present on his specimens was amylase, principally of salivary type and discovered salivary glands acinus in lung parenchyma. In 1985 Nandapalan studying alpha-amylase activity in laryngectomized patients without salivary fistula found a alpha-amylase activity range from 35 to 1025, mean 428, standard ad deviation 367, and median of 295 IU/L in tracheobronchial secretions. His group considered these values as the normal levels of this enzyme in normal human beings. We disagree with that conclusion because it has long been demonstrated that normal people microaspirate saliva. This is the main reason of our project was approved by the Ethics and Research Commission at the University of Juiz de Fora- Minas Gerais-Brazil; UFMG, federal university of minas Gerais- Belo Horizonte- Brazil

Objective

To determine the normal levels of alpha-amylase in tracheobronchial secretions of patients with very low possibility of morbid aspiration, based on known risk factors. There is a possibility that alpha-amylase could be a marker of tracheobronchial aspiration of saliva and may be considered a tool for assessing aspiration of patients with oropharyngeal dysphagia. The main application of this tool could be in weaning from mechanical ventilation, allowing us to diagnose neurogenic dysphagia with major accuracy than the card test. Which is very subjective and as rule not used in clinical paractice. Currently, an objective tool to diagnose oropharyngeal dysphagia before extubation does not exist. By comparing saliva and tracheobronchial amylase activity, it is possible to diagnose oropharyngeal dysphagia and take measures to reduce saliva secretion and prevent weaning failure, thereby reducing morbidity, costs and mortality. This project was approved by CEP_HU_UFJF, Number: 0129/2009. The ideal procedure to define the physiologic parameters of lung amylase should use healthy volunteers ; but this probable would not be approved by the Committee on ethics in human beings, because the procedures have some side effects and a little, but possible, risk of complications.

Material and methods

From October 2009 to June 2011, we prospectively evaluated 111 patients without clinical signals of hypersecretion who underwent transtracheal puncture before undergoing bronchoscopy. The study project was approved by the Ethics and Research Commission at the
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Juiz de Fora Federal University (number: 0129/2009), Minas Gerais, Brazil. All patients signed an informed consent term. Inclusion criteria were patients without hyper secretion in the tracheobronchial tree, and who were referred to the Thoracic Surgery Service for diagnostic flexible bronchoscopy. Patients with any risk factors for aspiration owing to neurologic or muscle degenerative disease, acute cerebral ischemic event, and surgery or radiotherapy of the cervical region, age higher than 65 years was not included in this study.

All patients were asked to lie in a supine position with light cervical hyperextension. The antisepsis was performed at the anterior cervical region using a 70% alcoholic solution. All patients underwent sedation and local anesthesia. Anesthesia was administered intravenously with a combination of diazepam and meperidine to achieve a sedation level of 2-3 in the Ramsay scale. The local anesthesia comprised the skin, subcutaneous, and cricothyroid membrane levels and was performed using 0.5 to 1ml of 2% lidocaine solution (Xylestesin®).

We performed the puncture of the cricothyroid membrane with a 25/7G needle, and 10 ml of lidocaine (2%) without a vasoconstrictor was injected into the tracheobronchial tree. Lidocaine was allowed to take effect by waiting for 2 minutes and spreading into the tracheobronchial tree. This technique of anesthesia eliminates the cough reflex and leaves no free lidocaine in the major airways. The transtracheal puncture was then performed using an intravenous catheter passing into a 14G needle (BioCat®-sao Paulo-Brazil). After the puncture, the needle was positioned about 45° in the cranial-caudal direction. With the catheter introduced 5-10 cm in the tracheal lumen, we performed an aspiration with the vacuum system at maximum power. In the event of no recurrence of secretion, the patient was confirmed as non-hypersecretory. For such cases, we proceeded with the infusion of 10ml saline solution, and immediately the catheter was connected to the vacuum system, through which the specimen was aspirated. The vacuum system was a conventional system with a pressure from 30 to 40 cm Hg. When the volume of the aspirate was at least 100µl, it was collected as a sample of saliva (bottle 1) and the aspirate (bottle 2).

The high viscosity of both fluids was responsible for the negative results in 19 patients. After diluting the samples, this problem was solved. After these procedures the patients were submitted to video-naso-laringo-bronchoscopy and evaluated for morbid aspiration. In case of positivity for aspiration the patient should be excluded from the study. This fact never had occurred. Activity of α-amylase was measured using α-2-chloro-p-nitrophenyl-α-maltotrioside1,4 which is hydrolised by α-amylase releasing 2-chloro p nitrophenol that can be measured by photometry. We used LabMax240 (Labtest®- Amylase CNPG liquiform, Lagoa Santa, Minas Gerais, Brazil) for measuring of α-amylase in both sample fluids. SPSS software version 13 for Windows was used to compile the survey data. We used, KS test for normality evaluation, γ-squared test for categorical variables, Student T test for means and normal distributed, Pearson correlation for analyzing the relationship between salivary and tracheobronchial amylase, descriptive statistics and Kruskal-Wallis test to compare amylase activity among the various radiological diagnosis.

Results

The population studied had a mean age of 48.7 years (standard deviation of 14.7 years), 77 male 77 (77%), 72 (65%) white and active smoking. There were no statistically significant differences in age, gender, race or smoking history (Table 1 & Table 2).

Table 1 Main patient's characteristics

| Number of patients percentage (%) | Mean(age) | P value |
|-----------------------------------|-----------|---------|
| White 72(65%)                     |           |         |
| Race                               |           |         |
| Non-White 33(30%)                 |           | 0.159   |
| Missing 6(5%)                     |           |         |
| Sex                                |           |         |
| Male 77(70%)                      |           | 0.139   |
| Female 34(30%)                    |           |         |
| Smoking                            |           |         |
| Yes 7(68.5%)                      |           | 0.222   |
| No 32(28.8%)                      |           |         |
| Missing 3(2.7%)                   |           |         |
| Age(Years)                         | 48.7      | 14.4    |

Table 2 Alpha amylase activity

| Mean(age) | Median | SEM  |
|-----------|--------|------|
| 19(4)IU/ml| 1056IU/ml| 240IU/ml |

Discussion

The α-amylase activity in the human tracheobronchial tree seems to have two origins. One is from local production and the other is from physiological microaspiration. Nandapalan in 1995 published two works on this-one on laryngectomized patients and the other on tracheotomized patients. It was shown that the human lung produces amylases and thought that this should be the normal value of α-amylase activity in humans. We disagree with this information because we have shown that physiological microaspiration occurs. Therefore, evaluating patients without the signs of morbid aspiration to obtain normal values of α-amylase in the lungs needs to be performed. This is important because α-amylase could be a useful tool in the diagnosis of morbid aspiration. The gold standard for diagnosis aspirations are videofluoroscopy and videendoscopy. However, both methods have limitations. Videofluoroscopy is only useful in patients with some degree of cognition to obey the order made by the speech therapy specialist for treating oropharyngeal dysphagia. Videendoscopy has advantages and disadvantages in comparison with videofluoroscopy; for example, videendoscopy does not evaluate the oral phase of dysphagia, and, akin to videofluoroscopy, is not worldwide economically viable. Transtracheal aspiration or puncture (TTA, TTP) has been used widely because of the original publication, although it is not an ideal procedure because transfixing the anterior wall of the trachea, as in the original method carries major risk of haemorrhagic complications. The recommended name for this technique would be, “transcricothyroid membranes puncture” owing the fact it is an avascularity structure and with less chance of thyroid gland penetration, therefore with less chance of bleeding. The more appropriate term for the technique should be, “transcricoid puncture (TCP).” However, the descriptor “transcricoid” does not exist, and all the research on the current subject must use the terms “transtracheal aspiration and puncture.” It is worth noting that our pilot study of 33 patients was critical in modifying the classical
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Conflict of interest
The author declares no conflict of interest.

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Conclusions
We have defined the normal values of amylase in tracheobronchial secretions of human beings. The alpha-amylase in the human lung has probably two components: one produced locally and the other due to normal physiologically aspiration.

Acknowledgements
None.