Administration of Alpha-1 Antitrypsin in Haemodialysis

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Abstract
The deficit of alpha-1-antitrypsin (AATD) is the most common potentially fatal hereditary disease in adulthood, causing the onset of emphysema, various liver diseases and favouring the development and progression of tumours and systemic vasculitis. Treatment is replacement rate. The dosing schedule alpha1-antitrypsin (α-1AT) has been modified over time by infraestrutura centers dedicated to it. In the beginning, a weekly administration agreed, but given the saturation of the hospitals and the deterioration of the quality of life of patients having to go to hospital every week, it was decided to increase the interval to every 15-21 days.

Keywords: Alpha-1 Antitrypsin; Human serum pro tease; Perfusion; Nephelometry

Introduction
The α-1AT, 52KD glycoprotein, is the most abundant inhibitory human serum protease [1] has significant anti-inflammatory properties (blocked cytotoxicity conducted by neutrophils and stimulating stetitis IL-8, IL-6, IL1β), TNF α- and other cytokines), increasing their activity in inflammatory, tumoral or infectious processes [2]. Its deficit (plasma levels below 35% of normal) is transmitted with a recessive autosomal pattern and leads to the development of emphysema and chronic liver (cholestasis, cirrhosis), and predisposes to the onset and progression of tumours and systemic vasculitis [3,4], therefore it is considered life threatening.

The only treatment currently available, in addition to the usual support of any bronchial condition, is symptomatic substitute, not “curable” and consists of intravenous (IV) of purified human α-1AT, which maintains its enzyme activity in plasma and in broncho alveolar lavage. Lung activity correlates directly with their plasma concentration, allowing monitor treatment [1].

The administration schedule α-1AT has been changing over time as the infraestrutura centers dedicated to it. Thus, in the eighties, with the saturation of the hospitals and the deterioration of the quality of life of patients having to go to hospital every week, it was decided to increase the interval to every 15-21 days.

To date there is no literature on administration in hemodialysis patients, case presented.

Clinical Case
44 patients with a family history of AAT (ZZ phenotype) with hepatic impairment (centrilobular cholestasis with portal hypertension) and pulmonary (panlobular widespread emphysema/panacinar, bulbosa dystrophy and bronchiectasis colonized by Pseudomonas aeruginosa conditioning recurrent respiratory infections). Moderate OSA, rejecting treatment with nasal CPAP intolerance. In 2006 it presents criteria for chronic airflow obstruction (FVE1 26-28%) and dyspnea on minimal exertion, requiring home oxygen. α-1AT replacement therapy with purified human (Prolactina®), Talecris Bio therapeutics) from diagnosis in 1998, in addition to tiotropium, salmeterol-fluticasone, acetylcysteine and nebulized colistin since April 2011. From the point of view nephrology, has advanced chronic kidney disease secondary not biopsied chronic glomerulonephritis (nephrotic syndrome), on hemodialysis since December 2007. Guideline dialysis: 4 hours/session, 3 sessions/week, through a FAV humerobasilica reaching a Qb 450 ml/min and dialysis polyacrylonitrile 1.65 m², CUF 50 ml/h/mmHg).

Prior to the start of hemodialysis Prolactina® administration took place in “day hospital” the pattern of 400 ml (10 grams) to spend 2 hours every 21 days, with blood drug levels far below what is considered therapeutic or protective (Figure 1).

Since the start of renal replacement treatment in our unit, the intradialysis administration was decided by drug infusion pump postfilter a maximum rate of 4 ml/min. To do this, we ensure that Prolactina® molecule (52 kDa) not spread through the dialyzer: plasma samples pre and post-dialysis were taken and in the effluent of ultrafiltration (Table 1).

In January 2008, we start with the usual pattern of pulmonary at 180 mg/kg/21 days, but to persist infraterapeuticos levels frequency to fortnightly (April 2008: 120 mg/kg/15 d) is increased and last week (July 2008: 60 mg/kg/week), reaching levels in rank order.

The sample for measurement Prolactina® predialysis levels taken before the infusion (Cmin trough levels), considering the normal range by nephelometry between 90-200 mg/dl. The drug preparation is performed in hospital pharmacy, by reconstituting the powder solution perfusion solution. Each vial contains 1000 mg human α-1AT and each milliliter of reconstituted solution of 25 mg protein. This solution should be administered within three hours of its preparation at a rate not exceeding 0.08 ml/kg/min.

The evolution to 7 years since initiating treatment intradialysis has been very satisfactory. The patient is with marked improvement, without dyspnea or need for home oxygen since 2012 and performing...
daily activities without difficulty. He has presented few bronchial exacerbations (an average of two per year) without requiring hospitalization or increased baseline bronchodilator treatment. Imaging techniques have not shown progression centrilobular emphysema or bronchiectasis. Pulmonary function tests show no worsening (Figure 1). The only adverse event related to treatment was the presence on a single occasion of grade I dyspnea associated with infusion faster than usual, giving the endentecer up.

We present a patient with homozygous AAT that, to clarify the situation of renal replacement therapy with hemodialysis, he increased the frequency of administration of the α-1AT (intradialysis once weekly administration as we confirmed molecule was not dialyzed), thereby entering levels drug in the therapeutic range, while allowing a slowing in the progression of the disease.

Discussion

AAT deficiency is the second most important cause of lung disease of genetic nature. The only specific treatment that is currently available replacement with the purified protein from human plasma, indicated in homozygous patients with airflow obstruction on spirometry, obtaining maximum benefit in those with forced expiratory volume (FEV1) between 30-65% [1-6]. It is not curative but their periodical administration has been associated with clinical improvement, especially in moderate stages [8,9]. Although there is some discrepancy in clinical trials for improving lung function, its efficacy has been demonstrated in numerous experimental and observational studies, especially by reducing the incidence of chest infections and exacerbations [2,10-12]. Its mechanism of action is based on its neutralizing neutrophil elastase activity, inhibiting tissue destruction in tissue bronchioloalveolar [2]. It does not happen just in liver tissue, which does not confer on protection [2,3]. In addition, he is credited ability to inhibit replication of genetic nature. The only specific treatment that is currently available replacement with the purified protein from human plasma, indicated in homozygous patients with airflow obstruction on spirometry, obtaining maximum benefit in those with forced expiratory volume (FEV1) between 30-65% [1-6]. It is not curative but their periodical administration has been associated with clinical improvement, especially in moderate stages [8,9]. Although there is some discrepancy in clinical trials for improving lung function, its efficacy has been demonstrated in numerous experimental and observational studies, especially by reducing the incidence of chest infections and exacerbations [2,10-12]. Its mechanism of action is based on its neutralizing neutrophil elastase activity, inhibiting tissue destruction in tissue bronchioloalveolar [2]. It does not happen just in liver tissue, which does not confer on protection [2,3]. In addition, he is credited ability to inhibit replication of viruses and bacteria [1].

Its rate of infusion makes a hospital stay of at least two hours. This was in the 90s a saturation of day centers in which the product was administered together with a deterioration of the quality of life of patients who had to go to hospital every week, which created different patterns treatment described above, which spread the frequency of its administration up to 30 days [1,2,10-12], although sheet will continue to recommend the weekly, to be the best documented [1,13,14] and which allows better pharmacokinetic levels. Therefore, using the patient should go to their scheduled dialysis three days a week and since no therapeutic levels were achieved with monthly administration of α-1AT sessions, we decided to administer in our unit shortening intertherapy first fortnightly and then weekly period, obtaining very good clinical, functional and analytical results.

Clinical markers at our disposal to monitor the effectiveness of substitution treatment are mainly [13-15] 1) FEV1, 2) TAC density, 3) frequency of respiratory infections, 4) sputum markers. All of them are considered to FEV1 as the main predictor of survival in these patients, who at 2 years is almost 100% until the FEV1 reaches 33%, and thereafter, decreases exponentially and poses 50% when FEV1 is 15% predicted [11]. The current goal in the treatment of AAT deficiency is the increase in the level of AAT in plasma and interstitial lung both above the protection threshold, whereby said plasma levels are used as a guide for replacement therapy. Minimum or trough concentration at steady state (Cmin in the pre-administration of the next dose time is measured. It is considered a Cmin of 50 mg/dl the right to provide a good clinical, functional and analytical results.

By the molecular weight of the product (52 KD), should not spend the hemodialysis membrane [16]. To ensure this premise, serial measurements in the ultrafiltration without finding significant concentrations of the molecule were performed. However, the suspension of plasma protein is not the same as in the ultra-filtrate and nephelometry is ready to blood plasma, so that it cannot safely ensure that a minimum amount of the product is not dialyzing. If so, in our case, this amount should not be significant since it achieve and maintain therapeutic levels.
We observed in our patient, since entering dialysis and thus the increasingly frequent administration α-1AT how the same levels increased progressively to be therapeutic (Figure 1). We objectify also a correlation between plasma levels of these improved with a decrease in the frequency of acute respiratory infections especially with stability over time of the operating parameters and a clear improvement of the general condition of the patient. The latter cannot be fully adjudicated by Pseudomonas aeruginosa. AATD mortality is high, dated in some studies 30% at 5 years, the main predictor FEV1 below 50% [17].

Another drawback of the administration of this treatment outside the hemodialysis session is repeated puncture of peripheral veins, often considering channeling central access type reservoir to precisely reduce the hemodialysis session is repeated puncture of peripheral veins, often considering channeling central access type reservoir to precisely reduce the hemodialysis session. Treatment outside the hemodialysis session is repeated puncture of peripheral veins, often considering channeling central access type reservoir to precisely reduce the hemodialysis session.

Conclusion

According to the recommendations of the Spanish and American Societies of Pneumology, in the absence of conclusive studies linking clinical efficacy pharmacokinetic measures, the choice of α1Antitritpsina administration schedule should be individualized and emerge from compromise between biochemical efficacy, expectations and availability of patients and the hospital. We therefore conclude that the administration should be weekly in patients undergoing hemodialysis, since this scheme maintains higher levels of drugs in blood with good safety profile and tolerance, improves quality of life and availability of patients and the hospital. We therefore conclude that the administration should be weekly in patients undergoing hemodialysis, since this scheme maintains higher levels of drugs in blood with good safety profile and tolerance, improves quality of life and presumably the same effect on survival patient.

References