Pulmonary Hypertension and Erythropoietin

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\textbf{Key Words}
Chronic hemodialysis · Chronic renal failure · Erythropoietin · Hypertension · Nitric oxide · Vascular access, hemodialysis

\textbf{Abstract}
Numerous uremic patients on hemodialysis have pulmonary hypertension attributable to the presence of arteriovenous fistulas, vascular calcification, and endothelial dysfunction due to alterations in the balance between vasoconstrictive and vasodilatory substances. For these reasons, the effects of recombinant human erythropoietin, a drug widely used in patients on dialysis, on the pulmonary circulation were studied. Some authors maintain that recombinant human erythropoietin has an antihypertensive effect, while others have observed that this hormone induces a reduction in pulmonary arterial pressure due to its vasoactive and stimulatory effects on endothelial and smooth muscle cell precursors.

\textbf{Pulmonary Hypertension (PH): The Problem}
It is widely known that uremic patients with end-stage renal disease on chronic hemodialysis having an arteriovenous fistula are at an increased risk of atherosclerosis and cardiovascular diseases, including PH. PH occurs in 40% of the patients, its instrumental signs being unequivocal and exposing the patients to an increased morbidity and mortality [1, 2]. Uremia affects the lung with pathological alterations that worsen, as the disease progresses. Particularly noteworthy are anomalies in the transport of respiratory gases, alterations in diffusive mechanisms, diminution in ventilatory processes at rest, and functional alterations in the respiratory muscles [3, 4]. These patients, moreover, almost always present a significantly higher cardiac ejection fraction, low hemoglobin and hematocrit levels, and an increase in the pulmonary arterial pressure (PAP). The PAP usually tends to normalize, whenever arteriovenous fistula closure is undertaken, e.g., when the patient undergoes renal transplantation [5]. Chronic hypoxia elicits a number of physiological responses that result in PH. Hypoxia causes active pulmonary vasoconstriction as well as a structural remodeling of the pulmonary arterial vasculature. Both of these responses diminish the luminal diameter of the small pulmonary arteries, increasing vascular resistance and contributing to the development of PH [6]. In table 1, we have summarized the most common pathological lung alterations in uremic patients.

\textbf{PH in Hemodialysis Patients: The Reason}
From a pathophysiological point of view, an increase in the cardiac output under normal conditions cannot in itself cause PH, considering the enormous capacity of the small circulation. However, it is also true that in patients...
Table 1. The most common pathological lung alterations in hemodialysis patients

<table>
<thead>
<tr>
<th>Number</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>Anomalies in the transport of respiratory gases</td>
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<td>2</td>
<td>Diminution in the ventilatory processes at rest</td>
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<tr>
<td>3</td>
<td>Functional alteration in the respiratory muscles</td>
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<tr>
<td>4</td>
<td>Increased uremia-mediated vascular pulmonary resistance</td>
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<tr>
<td>5</td>
<td>Vascular calcifications with thickening of pulmonary vessels, alveolar interstitium, and wall of the respiratory tree</td>
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<tr>
<td>6</td>
<td>Increase of the vascular tone of the small pulmonary circulation, mediated by an attenuated production of nitric oxide</td>
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with end-stage renal disease this circulation may not be able to bear the increase in the cardiac load induced by an arteriovenous fistula. Not only does the increase in the ejection fraction contribute to this, but so does a reduced vasodilatory response cause the histological modifications associated with uremia which are vascular calcifications with thickening of the pulmonary vessels and endothelial dysfunction [7]. It is, therefore, possible that the pulmonary circulation of patients on chronic dialysis is the site of excessive vasoconstriction. Hormonal-metabolic alterations associated with uremia could be the cause of the increased vascular pulmonary resistance [8]. PH involves vasoconstriction and obliteration of the lumen of small vessels in the lungs by plexiform lesions, resulting in an increased resistance to flow. The proposed mechanisms for the formation of plexiform lesions include dysregulation of endothelial growth and angiogenic responses to local triggers [9]. Moreover, the phenomenon of extraosseous calcifications is very common in patients on chronic hemodialysis and involves different anatomical regions, including heart, kidney, and stomach [10, 11]. At the pulmonary level, calcium is deposited in the alveolar interstitium and in the wall of the respiratory tree, the main paths to the bronchioles, as well as in the depth of the pulmonary vessels. In their study, Yigla et al. [12], utilizing bone scintigraphy with $^{99m}$Tc-methylene diphosphonate, showed calcifications at the pulmonary level in about 40% of their hemodialysis patients, without finding a direct correlation between pulmonary uptake and PAP values. Although the consequences of calcifications of aorta and coronary vessels are known, it is equally evident that this phenomenon also concerns the small peripheral vessels. However, no in-depth studies have been conducted to clarify this aspect.

Also, in the absence of phenomena of vascular calcification, it is known that uremic patients have anomalies in the endothelial function, a mechanism through which they are exposed to a higher risk for the development of early atherosclerosis. It is known that vascular tone and its function are physiologically regulated at the local level by a balance between vasodilation, induced principally by prostacyclin and nitric oxide, and vasoconstriction, mediated above all by thromboxane A$_2$ and endothelin-1 [13]. Nitric oxide also inhibits platelet activation, adhesion, and aggregation and modulates the interaction between leukocytes and endothelial cells [14]. Endothelin-1, apart from its vasoconstrictive action, also has an important mitogenic activity, and its hematological levels are increased during the course of PH [15]. However, in their recent study, Nakhoul et al. [16] demonstrated that the endothelin-1 concentration, although statistically significantly elevated in patients on hemodialysis as compared with healthy subjects, is not correlated with PH; in the same study [16], the authors found, on the other hand, that the concentrations of nitric oxide metabolites (NO$^+$, NO$^3$) had a different behavioral pattern. Although the values of patients on hemodialysis were similar to those of normal individuals, the subgroup of patients with PH in a predialytic stage had levels of circulating nitric oxide that were lower than those in patients without PH. During the dialysis session, moreover, the levels of these metabolites increased significantly, and this increase was more marked in patients without PH. Given the vasodilatory and antimitogenic properties of nitric oxide, this study suggests that the attenuated production of this substance, basal and induced by the hemodialysis session, in patients with PH can be responsible for an increase in vascular tone in the small circulation. Moreover, the temporary normalization of PAP and cardiac output in hemodialyzed patients who undergo arteriovenous fistula closure or renal transplantation appears to indicate that the excessive pulmonary flow could be involved in the pathogenesis of this disease.

**PH: The Erythropoietin (EPO) Role**

It is, therefore, reasonable to ask ourselves whether traditional drugs prescribed for patients on dialysis, including anticoagulants, diuretics, digoxin, calcium antagonists, vasodilators, and antiproliferative agents, can reduce PH. On the contrary, other drugs, commonly taken by these patients, could play an important role in the genesis of PH. In this context, of particular interest is recombinant human EPO, a drug with an endothelium-mediated activity and which is utilized by all patients on dialysis.
The data reported in the literature appear, however, to suggest that chronic EPO treatment has an antihypertensive action, at least under experimental conditions. In a recent study, Sato et al. [17] reported that the endogenous EPO/EPO receptor system prevents the development of PH during chronic hypoxia in mice in vivo through the recruitment of epithelial progenitor cells, suggesting the therapeutic importance of this system for the treatment of PH.

These data are particularly significant in animals made genetically deficient of the EPO receptor at the level of the cardiovascular system. Also the mobilization of the endothelial progenitor cells and their recruitment by the pulmonary endothelium appear to be compromised. Hypoxia, moreover, would increase the expression of the EPO receptor on pulmonary endothelial cells of control rats, whereas in knockout rats it would activate the endothelial enzyme nitric oxide synthase. Finally, the EPO/EPO receptor system would play a protective role in the pulmonary endothelium, inhibiting the development of PH induced by hypoxia.

Weissmann et al. [18] have studied the pulmonary vascular tree in transgenic rats that had overexpression of EPO under conditions of both normoxia and chronic hypoxia. Notwithstanding the increase in hematocrit, the rats maintained in normoxia did not develop right ventricular hypertrophy. The numbers of vessels with a diameter of 51–95 μm and >155 μm were found to be increased, while the percentage of small vessels (30–50 μm) was diminished. Also vascular pulmonary resistance, hypoxic vasoconstriction, measured in isolated perfused lungs, and vasoconstriction induced by a thromboxane A2 mimetic (U-46619) tended to be reduced. Moreover, after chronic hypoxia, the vascular resistance and the vasoconstrictive responses to hypoxia were also significantly reduced in transgenic rats as compared with controls and the grade of muscularization of the pulmonary vessels as well. According to these experimental data in animals, the congenital overexpression of EPO would appear to have an antihypertensive pulmonary effect, both at structural and functional levels.

An important question is the role played by polycythemia and EPO administration. In animal models, Walker et al. [19] did not observe any significant effect of recombinant EPO induced polycythemia on the PAP. These authors hypothesized that it is the local shear stress that induces release of nitric oxide, offering a protective effect against the development of PH.

This mechanism could be ineffective in hemodialysis patients. In fact, hemodialysis patients show low hematocrit levels despite EPO treatment, exerting shear stress, stimulating nitric oxide release.

Other authors [20] have administered EPO to mice with a different sensitivity to chronic hypoxia and have demonstrated that the drug does not increase the right ventricle/body weight ratio (index of ventricular hypertrophy) as well as the systolic pressure peak in hypoxia. In another study [21], transgenic mice with constitutio

nal overexpression of the EPO gene, in a way, however, independent of oxygen, presented polycythemia notwithstanding the fact that they had a normal arterial pressure. However, the PAP was found to be increased in vivo, and it decreased, on the other hand, in isolated perfused lungs. The pulmonary vascular apparatus was characterized by a high production of prostacyclin, a strong expression of endothelial nitric oxide synthase, and a reduced thickness of the smooth muscle sheath of the pulmonary vessels. The fact that these transgenic and polycythemic animals developed a marked PH in vivo but not in vitro suggests the pathogenic role of the blood hyperviscosity in this experimental model. Polycythemia, therefore, would, in this sense, play a role independent of the development of PH. The lungs of these transgenic mice, moreover, have shown a mechanism of adaptation to the high PAP: an increase in the local synthesis of substances with a vasodilatory action and a reduction in the vascular smooth muscle tone. On studying the blood flow of the human pretibial muscle by means of 133 Xe clearance, Buemi et al. [22] found an EPO-induced reduction in post-ischemic vasodilation. Moreover, in anesthetized rats subjected to hemorrhagic shock by intermittent bleeding, EPO administration significantly increased the mean arterial pressure, the survival time, and the percentage of animals surviving with respect to untreated controls [23]. EPO also seems to be able to inhibit acetylcholine-induced cutaneous vasodilation and can stimulate the endothelial release of endothelin and inhibit nitric oxide, a potent vasodilator [24]. A possible explanation for the diverging results following EPO treatment can be different administration modalities. Experimental models used chronic EPO treatment, while in human models infusion of single acute doses leads to enhanced PAP.

Regarding the pulmonary resistance, in a previous study, Allegra et al. [25], using a catheter placed in the pulmonary artery, performed a hemodynamic evaluation in 2 patients with hypovolemic shock. EPO administration caused an increase in the pulmonary vascular resistance and in the mean PAP. Moreover, Allegra et al. [26], in 10 patients with chronic cor pulmonale and marked signs of right ventricular hypertrophy and a pul-

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Pulmonary pressure >35 mm Hg, have demonstrated that an intravenous bolus infusion of recombinant human EPO beta (70 U/kg) had an effect on the vascular pulmonary resistance. Such observations were recently made in vivo in animal models also by Satoh et al. [17]. The action of EPO on the pathophysiology of chronic cor pulmonale may, however, not be limited to its vasomotor effect on the pulmonary circulation; it may also impact on pulmonary vascular remodeling. It is well known that EPO can influence the proliferation process of cell types other than erythroid precursors, such as endothelial progenitor cells and smooth muscle cells [27, 28]. In EPO-transgenic mice, the relative lung weight is increased, the interalveolar septa have a greater thickness, and this is accompanied by interstitial and perivascular fibrosis [29]. Foster et al. [30] have demonstrated that developmental lung growth involves paracrine EPO signaling with parallel upregulation of the EPO receptor.

It may, therefore, be suggested that the chronic increase in endogenous EPO concentrations, present in patients with chronic obstructive bronchopathy, contributes to the thickening of the arterial wall of the pulmonary vessels, typical of chronic pulmonary heart disease.

The data obtained appear to confirm the importance of the role played by EPO in the pulmonary circulation, but also appear to partly contradict other reported data cited above. A significant increase in the indices of vascular pulmonary resistance has, in fact, been observed immediately after EPO administration. In table 2, we have summarized animal (preclinical) and human models showing experimentally the controversial influence of EPO in establishing a PH status.

### Table 2. Controversial influence of EPO in establishing a PH status

<table>
<thead>
<tr>
<th>Antipulmonary hypertension</th>
<th>Pulmonary hypertensive effect</th>
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<tbody>
<tr>
<td><strong>Animal Models</strong></td>
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<td>Remodeling operated by endothelial progenitor cells [17]</td>
<td>In vivo effect on the PAP of polycythemia [21]</td>
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<tr>
<td>Structural adaptation to hematological viscosity enhancement [18, 20]</td>
<td>Acute EPO administration in shocked rats [23]</td>
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<td></td>
<td>Interstitial and perivascular fibrosis [29]</td>
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<td>Upregulation of EPO receptor in the lung vessels [30]</td>
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<td><strong>Clinical Models</strong></td>
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<td></td>
<td>Acute EPO administration in chronic cor pulmonale [26]</td>
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### EPO and PH: An Open Question

Uremic patients represent a particular model of PH correlated with hormone-metabolic alterations characteristic of renal disease itself. As compared with the healthy population, in fact, important phenomena of vascular calcification and lacerations are observed in the balance of vasoconstrictive substances, like thromboxane A₂ and endothelin-1, and vasodilator substances, like prostacyclins and nitric oxide. The use of substances with a vasoactive action, like EPO, therefore, continues to be a controversial issue. Together with the protective role of the hormone, evidenced in different studies using animal models, an increase in the pulmonary vascular resistance has been found in humans which is probably correlated not only with the vasomotor effect but also with the vascular remodeling due to stimulation of the EPO receptor. In the light of these data, further in-depth studies to be performed in humans are required; they should be conducive to regulating the use of this hormone in patients at an increased risk of developing PH, such as uremic patients.

### References


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