High altitude pulmonary oedema: still a place for controversy?

High altitude pulmonary oedema (HAPE) is defined as acute respiratory failure following exposure to high altitude hypoxia, developing in a normal subject with no pre-existing pulmonary or cardiac disease. It is one of the manifestations of high altitude disease of which the simplest form is acute mountain sickness (AMS), generally limited to headache, nausea and insomnia during the first few days spent above 2500 m. In contrast to AMS, HAPE necessitates emergency management. It occurs 6–48 hours after rapid ascent to altitudes above 2500–4000 m. Severe exercise and cold exposure favour its occurrence. Subjects complain of shortness of breath disproportionate to the effort exerted, a dry followed by a productive cough, fatigue, and chest pain. Clinical examination reveals tachypnoea, tachycardia, mild fever, cyanosis, crepitant rales and rhonchi. The heart rate is generally more than 120 beats/min and the respiratory rate more than 20 breaths/min. There is no evidence of cardiac failure. When hospital facilities are available, radiographic examination shows diffuse infiltrates in two thirds of cases ("patchy distribution"), and prominence of pulmonary arteries. Patchy distribution is seen on chest radiography early in the disease, becoming bilateral and diffuse in the more advanced state. Clearing of these exudates lags a day or two behind clinical improvement. Electrocardiography shows tachycardia and signs of acute right ventricular overload. Doppler echocardiography may reveal tricuspid regurgitation or even tricuspid insufficiency. Haemodynamic parameters include a high pulmonary arterial pressure and low capillary wedge pressure. A mild leucocytosis may be found (rarely more than 14 000/mm³). The arterial oxygen saturation (SaO₂) ranges from 40% to 70%, and vital capacity and peak expiratory flow are decreased. Neurological signs are frequently associated with HAPE and can dominate the clinical picture with incapacitating headache, staggering gait, ataxia, projectile vomiting, emotional lability, diplopia, dislocation in time and place, coma, and death. The mortality rate varies between 4% and 11%, depending on the rapidity of descent or oxygen treatment. The main differential diagnoses are pneumonia, pulmonary embolism, and left heart failure. In a Colombian study of 13 cases of HAPE the initial diagnosis was bronchopneumonia in children and myocarditis or coronary insufficiency in adults. Excessive speed of ascent is a major contributing factor to HAPE. In a group of 166 cases 65% took less than two days to reach the altitude at which they developed HAPE, and HAPE occurred before the fifth day in 73% of cases. Mortality was higher in subjects treated by descent alone (12%) than in those treated with oxygen alone (5%), or both (3%); those not treated had a mortality rate of 44%.

Who can suffer from HAPE?
The risk of developing altitude sickness occurs in many human activities including climbing, trekking, skiing, touring, and working at high altitude. HAPE can occur in people residing at high altitude (in the Andean Altiplano and in Colorado) who return from a few days’ stay at a lower altitude ("re-entry" oedema). The incidence was 0.6% in adults and 8–9% in children in Peru in the 1970s.

Mountaineers climbing high peaks in the Himalayas, the Andes, and also in Europe (Mont-Blanc, Monte Rosa), North America (Colorado, Alaska), and Africa (Mount Kenya, Kilimanjaro) are prone to develop HAPE, especially when they do not take enough time to acclimatise themselves to the altitudes between 2500 and 4500 m. A typical example is an alpinist arriving at a base camp around 5000 m, excited by the idea of climbing the mountain, going up to a high altitude camp, expending a lot of energy, feeling tired and dyspnoeic, and found dead the next morning in his tent.

Cases are often observed in less severe conditions such as ski resorts in the Alps and Colorado, trekking in Nepal (Everest base camp, tour of Annapurna). The small number of cases in Europe is explained by less information about HAPE, the short duration of stay, and a relatively easy descent. A new group susceptible to HAPE are sedentary and elderly people touring places of cultural interest, such as Lhasa in Tibet (3500 m) or La Paz in Bolivia (3500–4000 m).

Soldiers fighting in high regions are at great risk. For example, the sino-Indian war at the beginning of the 1960s caused a large number of casualties from high altitude diseases among the Indian troops (not acclimatised) but not the Chinese, and promoted a series of studies by Indian physicians. Even today a war is being conducted between India and Pakistan on a contested border at 5000–6000 m with many casualties from high altitude disease.

Human activities at high altitude, such as astronauts at observatories at Mauna Kea, Hawaii (4200 m) or miners in Peru, Bolivia or North Chile (up to 5950 m) raise the question of detecting susceptible subjects and its significance in occupational medicine. Thousands of workers, resident at sea level, will work in newly developed Chilean mines over the next few years, spending one week working at around 4500 m and one week resting at low altitude. This is a new challenge for physicians involved in the selection and follow up of this population.

Who “discovered” HAPE?
AMS was first described in ancient texts from Asia and then by explorers in South America. The first scientific studies on AMS began in Europe in the 18th century, and became more extensive in the 19th century with the work of Paul Bert and the development of high altitude laboratories by Joseph Vallot on Mont Blanc and Angelo Mosso on Monte Rosa. Dr Jacottet died in the Vallot Observatory at the top of Mont Blanc in 1891 from acute respiratory distress; the necropsy revealed a normal heart and lungs with bilateral congestion and considerable oedema. The diagnosis was, at that time, pneumonia, as in most similar cases in high altitude expeditions until the 1970s. However, Egl-Sinclair established a clear relation between these observations and altitude sickness. Can it be considered as the first description of a link between lung disease at high altitude and mountain sickness?: “Is Dr Jacottet’s death not caused by the same disease [AMS]? It is my opinion. . . . The rapid evolution of the lung inflammation can only be explained by the influence of...
mountain sickness on the weakening of the heart and on the energy of the nervous system of this young man, formerly so robust.27

In 1913 Ravenhill, a medical officer in a mining company in Chile, described a “puna” (AMS) of “the cardiac type”, with dyspnoea, rales and cyanosis, which he defined as an acute heart condition coming on in a perfectly healthy man and disappearing rapidly with descent.28

Animal studies have been conducted since 1925 to understand better the effect of hypoxia on the lung morphology and function. Oedema, more often interstitial than clearly alveolar, has been reported by several authors, but the mechanisms of its formation are not known.29-33

A case of pulmonary oedema was described by Hurtado in 1937 but the patient probably had pre-existing cardiac disease since signs of “circulatory insufficiency” (peripheral oedema, dyspnoea, congestion of lung bases) were still present long after his acute episode had subsided.35

If we defined HAPE as a pulmonary oedema occurring at high altitude in a subject where signs of pneumonia and cardiac failure have been discarded, then the first clear description of this syndrome appeared in the 1950s and was published in Spanish in Peruvian publications. Seven cases were described between 1950 and 1952 in La Oroya (3750 m)34-35 and seven between 1953 and 1954 in Morocco (4500 m)36 in residents of high altitude returning after a stay at sea level. No explanation was suggested, except that, in most cases, no clinical, radiological or electrocardiographic signs of cardiac failure were observed. In 1957 an extensive paper published by Bardalez described HAPE as a well defined nosological entity and precise hypotheses for its pathophysiology were suggested.37 In 1960–1 three publications in English provided worldwide dissemination of the information to the medical public.38-39

Houston described the case of a 21 year old cross country skier at 12,000 ft in Colorado. Dr P D White, a distinguished cardiologist, advised him that elevation of the pulmonary artery pressure might play a part in the pathophysiology of HAPE, with the contribution of cold and exercise (Houston, personal communication). This was confirmed in the following years by catheterisation studies which also showed normal wedge pressures; cardiac failure is not a possible causative factor of HAPE.40-42 From the mid 1960s extensive research has been performed in hypobaric chambers, field expeditions, and by normobaric hypoxia with hypoxic gas mixtures. Many publications have come from Peru, India, North America and Europe, providing new insights into the anatomicopathology, haemodynamics, pathogenesis, clinical manifestations of HAPE. The information has spread into the climbing community, thanks to “climbing physicians” such as Charles Houston who climbed on Nanda Devi in 1936 and K2 in 1938 and dedicated a great part of his life to the research and prevention of HAPE.35 However, in spite of the comprehensive scientific data available, erroneous diagnoses are still made and pneumonia or cardiac failure are sometimes diagnosed in cases of respiratory failure occurring at high altitude. HAPE is often overlooked or mistaken for other illnesses.

Is there a common pathogenesis and an individual susceptibility in all altitude-related diseases?

The frequent association of HAPE with high altitude cerebral oedema (HACE), papilloedema, albuminuria, or peripheral oedema suggests a common pathogenesis in which a bodywide increase in capillary permeability plays a central part.43-44 The oedema is common, and the combination of high pressure and high flow in the brain could be responsible for AMS.47 The combination of hypoxia and overperfusion are necessary to produce pulmonary oedema in dogs.48 In fact, systemic hypertension is not associated with HACE or AMS, whereas pulmonary hypertension, when observed before treatment in the acute phase of the disease, is always present in HAPE. Inversely, HAPE is not always preceded by clinical signs of AMS. Severe cases of AMS with peripheral, pulmonary, and cerebral oedema have been associated with hyperventilation, weight gain and decreased urine output, suggesting a common pathogenesis involving fluid retention.49-51 Thus “high altitude (HA) illness” has been suggested to be a general problem of water handling leading to water retention and transfer of fluid from intravascular to interstitial and intracellular compartments. The “high-pressure high-flow” hypothesis is advocated: systemic hypertension and cerebral vasodilatation induce HACE, pulmonary hypertension and high cardiac output induce HAPE, and cutaneous vasodilatation and systemic hypertension induce peripheral oedema. However, four facts go against this hypothesis: (1) cerebral oedema can appear without exercise – that is, without systemic hypertension; (2) pulmonary oedema can occur without high flow – that is, without exercise; (3) there is no cutaneous vasodilatation at high altitude; and (4) severe normoxic exercise is associated with high pressure and high flow and does not produce pulmonary or cerebral oedema, even if exercise-induced arterial hypoxaemia occurs in healthy subjects at sea level.52 There are anecdotal reports of pulmonary oedema in marathon runners but these are associated with water intoxication. However, peripheral oedema has also been found to occur with prolonged exercise at low altitude.53

Experimental evidence of hypoxia-induced increase in capillary permeability was first shown by Landis in 1932.52 Thereafter, studies gave conflicting results. Some authors showed experimental evidence of increased lung vascular permeability,53-59 while others failed to show any increase in vascular permeability or oedema formation in the lung.60-65 Increased transvascular clearance of radiolabelled albumin was found in humans at high altitude, associated with proteinuria,66 which suggests a general increased permeability.67 The contradictions observed may be due to species differences, experimental design, or sensitivity of the method used. Pigs and rats seem more sensitive to pulmonary oedema than sheep or dogs.68 However, Viswanathan found a similar incidence of HAPE in dogs and rats and a higher incidence in mice.69 Experimental design seems to be important, particularly the duration and intensity of hypoxia and the speed of ascent. In studies where exposure to hypoxia was less intense (for a few minutes or hours) the animals are less likely to show pulmonary oedema.64 When exposure to hypobaric pressures is very abrupt, mechanical injuries are more likely to occur.65 With a longer duration of exposure of 24–48 hours animals are more likely to mimic humans exposed to high altitude and to develop pulmonary oedema.69 The sensitivity and specificity of the techniques used to evaluate the vascular leak can also be important. Some methods specifically explore vascular permeability59 while others give indirect indices of vascular leak such as lung water or lymph flow,64 and some explore both endothelial and epithelial resistance to fluid flux.65

A low ventilatory response to hypoxia has been related to susceptibility to AMS or HAPE. Cases of HAPE have been associated with blunted or absent ventilatory responses to hypoxia or CO2.70-72 When groups of subjects with a history of HAPE were compared with normal subjects a lower mean hypoxic ventilatory response (HVR) was found in the susceptible subjects; however, a few subjects had values in the normal range.73,74 When the literature on the association between HVR and AMS or HAPE is reviewed, 77% of susceptible subjects (n = 43)
Pathophysiology of HAPE (figure)

Diagnoses of heart failure or pulmonary infection were rapidly discarded when the early haemodynamic studies showed pulmonary hypertension with normal or low wedge pressures, the absence of clinical signs of heart failure, and the rapid reversibility of clinical and radiological signs when oxygen availability was restored. Thus, increased PAP was considered to be the main feature in the pathogenesis of HAPE. To explain alveolar oedema in the presence of precapillary vasoconstriction necessitated further assumptions. Pulmonary venous constriction and trans-arterial leakage were proposed but were not confirmed experimentally. Alterations in coagulation were favoured by some authors in view of necropsy findings and modifications of coagulation factors. However, it is not clear whether these alterations are the cause or the consequence of HAPE. The concept of “overperfusion” of unobstructed or unprotected areas of the lung was developed in the 1970s by Hultgren and coworkers. In the overperfusion concept the primary cause of HAPE would be a non-homogeneous vasoconstriction or thrombosis leading to the transmission of high blood flow and pressure to capillaries not protected by vasoconstriction. Arguments in favour of this hypothesis are: increased PAP, presence of thrombosis at necropsy, presence of oedema when blood flow is much increased in a region of the lung (pulmonary embolism, congenital absence of right pulmonary artery), experimental occlusion of lobar arteries, radiographic patchy distribution, and inhomo- geneity of hypoxic vasoconstriction. However, hydrostatic pressures in leaking capillaries have never been measured, patchy distribution of radiographic images are transitory and non-specific for this type of pulmonary oedema, and coagulation disorders probably do not precede HAPE. When breathing hypoxic mixtures, the PAP of HAPE-susceptible subjects is not always increased above that of normal subjects. In contrast, very high values of PAP have been found at high altitude in exercising subjects without HAPE. Moreover, physiological situations where PAP is much increased (such as severe exercise in normoxia) are not associated with HAPE. Thus, an increase in PAP is probably necessary but not sufficient to induce HAPE. The concept of uneven vasoconstriction is questioned by the findings of Vock et al concerning the evolution of radiographic images from “patchy” to “homogeneous” and clinical findings suggesting that structural abnormalities are not involved in the pathogenesis. However, this concept was recently supported by the studies of West and coworkers who showed that a hydrostatic stress, possibly comparable to that evaluated in HAPE, could break the lung endothelium and epithelium barrier in a model of an isolated rabbit lung. These alterations are less marked after reoxygenation. The disruption of the endothelium would expose the basal membrane and activate the release of inflammatory mediators such as thromboxane B2. This “mechanistic view” or “stress failure” hypothesis of HAPE is comparable with other diseases or manifestations such as Goodpasture’s syndrome with alteration of basal membrane, haemorrhages in race horses, and disorders in mechanically hyperventilated patients. The mechanical damage is the primary cause, after which permeability is increased by the release of mediators.

As the concepts of the physiopathology of HAPE have developed, another aspect has expanded in parallel which suggests that the primary insult is the alteration of the endothelium, either by a direct effect of hypoxia or by the release of mediators acting on inflammation processes, vascular permeability, and vasomotor responses. In favour of this “cellular view” or “high permeability” hypothesis are the many experimental studies that show a hypoxia-induced increase in peripheral or pulmonary vascular permeability, the frequent association with oedema in other organs with the absence of systemic hypertension, the composition of alveolar fluid with a high protein content and mediators of inflammation, the frequent occurrence of HAPE at rest and often during sleep, the rapid reversibility of HAPE when oxygen availability is rapidly restored, and the common finding of interstitial oedema and altered oxygen transfer in the lung of normal subjects at high altitude. However, as already mentioned, some studies have failed to show any hypoxia-induced increase in capillary permeability, and the composition of alveolar fluid is also compatible with the “stress failure” hypothesis, leading to a leakage of fluid and proteins into the interstitium. The fibrin-rich alveolar oedema found at nec-
ropsy supports the hypothesis of capillary damage or increased capillary permeability. Subclinical interstitial pulmonary oedema is a common finding in newcomers to high altitude, without HAPE. In 1978 Staub hypothesised that shear stress-induced alteration of endothelium leads to a leakage of fluid into the interstitium.

The effects of hypoxia on the endothelial cell permeability were investigated by Ogawa et al. Cultured bovine endothelium in monolayers were exposed to a Po2 of about 2 kPa. The permeability to macromolecules was increased in a time-dependent and dose-dependent manner. A significant increase was shown after 24–48 hours with a maximum at 72 hours. It was reversible within 48 hours of reoxygenation. These changes were associated with morphological alterations – namely, larger cells and the presence of intercellular gaps. Thrombomodulin activity (a cell surface anticoagulant cofactor) was inhibited by hypoxia and reversed by reoxygenation.

The effects of hypoxia on pulmonary vascular leakage was investigated in rats by Stelzel et al. Rats were exposed for 1–48 hours at 60 kPa. The authors showed an increase in transvascular protein leakage that was inhibited by glucocorticoid pretreatment, augmented by adrenal-ectomy, and independent of pulmonary arterial pressure. This phenomenon was not seen for short exposures (1–13 hours), was similar in normobaric or hypobaric hypoxia, and was associated with increased lung water and perivascular oedema cuffs on histological examination.

Humoral processes include the release of mediators by macrophages, neutrophils, platelets, endothelial cells or epithelial cells. However, the nature of the mediator(s) is still debated. Histamine and bradykinin have been suggested. Increased permeability of hypoxia-induced activation of phospholipase A2 led to the study of metabolites of arachidonic acid as mediators of hypoxic oedema (leukotrienes, prostaglandins, thromboxane). Some were increased in the alveolar fluid of HAPE subjects in a model of mechanical lung injury, and in plasma at high altitude. Release of atrial natriuretic peptide at high altitude is controversial. It could be released in response to increased pulmonary arterial or right atrial pressures and induce an increase in capillary permeability. Oxygen radicals can induce vascular injury after reoxygenation and may be involved in hypoxia-induced lipid peroxidation in rat lungs. Cytokines (TNF, IL-1, IL-6) and nitric oxide (NO) may also have a role but their specific action on capillary permeability remains to be established. Adhesion molecules present on the surface of endothelial cells and neutrophils have recently been shown to participate in the inflammatory process. Endothelial leucocyte adhesion molecule 1 (ELAM-1), intercellular adhesion molecule 1 (ICAM-1), and vascular cell adhesion molecule 1 (VCAM-1) are some of the molecules that can be found in plasma and serve as markers of endothelium or neutrophil activation. Mediators such as thrombin, histamine, and cytokines (TNF, IL-1) can activate the expression of adhesion molecules via a de novo protein synthesis. Lung vascular injury could be prevented by pretreatment with anti-adhesion molecule substances. Both experimental and human studies have shown a hypoxia-induced increase in plasma endothelin levels. In a study of 10 subjects plasma levels of endothelin were increased by 51% after one week of exposure to 6542 m. By contrast, moderate or severe exercise failed to stimulate endothelin release and the exercise-induced increase in plasma renin activity was inversely related to plasma endothelin levels. In a study at the Observatoire Vallot endothelin levels increased by 82% after five days at 4350 m; in addition, plasma levels of ELAM-1, but not ICAM-1, were also increased.

A third hypothesis was developed in the 1970s, by analogy with traumatic neurogenic pulmonary oedema. A sudden discharge in the adrenergic system would lead to pulmonary oedema, either by pulmonary venous constriction or left heart failure due to systemic hypertension. Neurogenic pulmonary oedema could also be directly mediated by a permeability defect. The only positive argument in favour of this theory is the beneficial effect of α blockers in HAPE subjects.

The conclusion drawn by Oelz et al in 1989 for the pathophysiology of HAPE is “permeability oedema in which hypoxic pulmonary hypertension is a crucial factor by enhancing the flow of liquid across the damaged endothelial barrier”.

### HAPE: vascular or epithelial leak?

It is important to consider that vascular permeability to fluid is a normal physiological process described by Starling’s equation. Moreover, many individuals develop pulmonary interstitial oedema or subcutaneous oedema at high altitude without HAPE. Fishman stressed the role of the alveolar rather than the endothelial barrier, and the role of lymphatic drainage in limiting interstitial oedema. Pulmonary interstitial oedema occurs with a small increase in vascular hydrostatic pressure. Compliance of lung vessels and interstitium is high, and only a 5–8% increase in lung weight is associated with peribronchovascular oedema. As much as a 35% increase in lung weight is necessary for alveolar oedema. Epithelial permeability is 10 times less than endothelial permeability which makes the epithelium a much stronger barrier than the endothelium. The question arises, however, whether the alveolar epithelium rather than the capillary endothelium is the limiting factor in the development of alveolar oedema.

Recent studies indicate that certain pneumocyte functions may be altered in hypoxia. Sodium transport by the alveolar epithelium represents an important mechanism for clearance of airspace fluid in acute lung injury. We have examined whether hypoxia affects Na-K-ATPase activity in alveolar epithelial cells. SV40 virus transformed rat type II alveolar epithelial cells were exposed to either hypoxia (5% O2 or normoxia) at 37°C for increasing durations (up to 48 hours) in the absence or presence of 10-5 M nifedipine. Na-K-ATPase activity was determined using ouabain-sensitive rubidium-86 influx (OsRb). Exposure to hypoxia for at least 12 hours induced a time-dependent decrease in OsRb. Incubation of normoxic cells with supernatant of hypoxic cells resulted in a 45% decrease of OsRb within one hour. Nifedipine prevented the hypoxia-induced decrease in OsRb. These results indicate that: (1) hypoxia induces a time-dependent decrease of Na-K-ATPase activity in alveolar type II cells; (2) this effect is most likely to be due to the release of a soluble factor; and (3) it is prevented by nifedipine. Autocrine alteration of the Na-K-ATPase activity in alveolar epithelial cells during hypoxia may reduce clearance of airspace fluid and contribute to the formation and/or maintenance of alveolar oedema.

### Treatment of HAPE

The complexity of the physiopathology is in sharp contrast to the simplicity of the treatment and evolution of the disease. Very few controlled studies have been conducted on the treatment of HAPE for obvious practical ethical reasons. The main treatment is to restore oxygen availability, either naturally by descending, or by inhaling oxygen or recompression in a portable hyperbaric chamber. Portable hyperbaric chambers have led to new practical possibilities for field care of HAPE. However, hyperbaric treatment...
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has a short beneficial effect on AMS but no controlled evaluation has been performed in patients with HAPE. It should not delay descent, which is the most efficient way to treat patients with HAPE.

The use of frusemide, first advocated by Indian authors, was rapidly abandoned because of lack of efficiency and the pathophysiological characteristics of HAPE. Blood volume and blood flow are already low, there is no left ventricular failure, and the risk of hypovolaemia and hypotension is high. Morphpine has frequently been used in the field but without controlled studies, and it is generally discarded because of its depressant effect on ventilation. Dexamethasone has been used but no controlled studies of its use in HAPE have been performed. The efficiency of acetazolamide, well established for AMS, is debatable for HAPE. Antibiotics are recommended because infection may develop in the waterlogged lung. Refractoriness to administration of 100% oxygen is rare in HAPE; however, in some cases the development of severe prolonged acute respiratory failure may initiate a state similar to the adult respiratory distress syndrome (ARDS).

The idea of using nifedipine in HAPE came from the effect of calcium antagonists on pulmonary hypertension in rats. In 1987 Oelz reported his own case of HAPE which occurred at 6900 m. He used nifedipine 20 mg sublingually first, followed by 20 mg slow release nifedipine. This allowed him to descend to 5400 m. Another episode of HAPE the following night was treated the same way and resolved, allowing a further descent to 4000 m the next morning. Obviously the nifedipine helped, but the descent itself may have played a part.

Nifedipine was given (20 mg sublingually followed by 20 mg slow release every six hours) to six subjects with HAPE at 4559 m. Clinical improvement, increased PaO2, decreased (A-a) DO2, decreased PAP (echo Doppler), and progressive clearing of oedema (radiography) was observed without side effects except persistent headache in four subjects. The decrease in PAP was probably responsible for the therapeutic effects, although other pharmacological effects of nifedipine cannot be ruled out. For example, nifedipine could also act in blocking the inflammatory response to hypoxia or pneumocyte reabsorptive function. The general use of nifedipine in HAPE-susceptible subjects should not be encouraged at present as it has potentially harmful side effects; however, it offers a potential emergency measure for patients when descent or evacuation is impossible.

The prophylactic administration of nifedipine (20 mg slow release every six hours) is effective in lowering PAP and preventing HAPE in susceptible subjects.

In 14 mountaineers ascending to 4559 m, with or without AMS but no HAPE, nifedipine decreased PAP and slightly decreased systemic arterial pressure, but had no effect on PaO2 and AMS; thus, nifedipine or isradipine cannot be recommended for prevention of benign AMS.

The rationale for the use of α blockers is based on neurogenic pulmonary oedema with massive adrenergic discharge. It was shown to be efficient in field situations, with or without oxygen.

Prevention depends on slow ascent for progressive acclimatisation, rest after ascent, clinical education, and early recognition of symptoms. The detection of HAPE-susceptible subjects with appropriate hypoxic tests should be encouraged.

Inhaled nitric oxide (NO) has recently been used to treat HAPE at 4559 m. Its vasodilator action on the pulmonary vasculature, associated with its rapid inactivation within the circulation by haemoglobin which prevents any peripheral action, are interesting features. However, it is not yet of practical use in the field.

New developments

There are two main barriers to a better understanding of HAPE: (1) the lack of a good animal model of hypoxia-induced alveolar oedema, and (2) the difficulty of performing controlled studies in patients suffering from HAPE in remote areas, far from hospital facilities. Non-invasive studies of the pulmonary circulation could help us to understand the time dependence of pulmonary hypertension with the appearance of HAPE. Is alveolar oedema the cause or the consequence of pulmonary hypertension? Molecular biology could also help by providing information about the eventual role of hypoxic sensing dysfunction. Hypoxic sensing can be seen in chemoreceptors, in EPO-producing cells, and in vessels producing vasodilatation in the lungs and vasodilatation in the periphery. HAPE is most probably sensed at the molecular level and upregulates the expression of several genes, especially coding for EPO, endothelin, or vascular endothelial growth factor (also known as vascular permeability factor). Studies on these molecular mechanisms may provide further insight into the pathophysiology of HAPE and in the susceptibility of individuals to develop the condition.

In-breeding studies in cattle have shown that the responsiveness of the pulmonary circulation to hypoxia is genetically coded. Anecdotal reports of a familial predisposition to HAPE need further investigations in humans. Genetic characteristics of HAPE could be explored with new generations of young alpinists.

Conclusions

The hypoxic challenge triggers important changes in both the pulmonary and the peripheral circulation systems. Increased interstitial or tissue oedema is the main feature of altitude-induced hypoxia. An exacerbation of the permeability properties of the endothelium could be directly mediated by hypoxia or by substances released locally. In the pulmonary circulation hypoxic arteriolar vasconstriction and pulmonary hypertension interfere with the increased vascular permeability. Interstitial oedema has only limited functional consequences, but when oedema develops within the alveoli or the brain severe consequences may result. The precise mechanisms by which lung interstitial oedema turns into alveolar oedema remain to be elucidated by further studies on epithelial function. Individual susceptibility to the condition also deserves further investigation so that better means of prevention and treatment can be developed.

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