High altitude in the heart of healthy and sick
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Summary
Today exposure to high altitudes is relatively common. Thus, basic knowledge is important also for physicians that are not specialised in this topic. Exercise capacity is limited at high altitude, mainly due to hypoxia. Additional factors (circumstances) may aggravate the problem. Still, day trips to altitudes up to 3000–3500 m are relatively safe even in patients with cardiovascular diseases, unless patients are not stable. Unstable patients are not suited for hypoxia exposure although direct evidence is largely lacking. Overnight stay at high altitude increases the risk. Also, patients with moderate to severe pulmonary hypertension, irrespective of the underlying cause, should limit exposure to altitudes of not more than 1500–2000 m.

With respect to flying, similar limitations apply. Moreover, circumstances of travel must be taken into account (e.g., limited access to medical help) and sufficient medication (in duplicate at different places) should be taken with the patient.

As discussed in the last part of this review, subjects that are susceptible to develop high altitude pulmonary edema should avoid rapid ascent to high altitude, or should be treated prophylactically. To what extent research in these subjects is relevant for pathophysiology of pulmonary hypertension at normoxia needs further investigation.

Key words: high altitude; hypoxia; cardiovascular disease; travel

Introduction
It might be argued that research of high altitudes is relevant for a very small proportion of subjects in clinical practice only. This assumption is, however, not true because of different reasons. Thus, trips to and also stays at high altitude are increasingly popular. Importantly, many people live at altitudes above 1500 m and relatives living at low altitude wish to visit them. Both healthy subjects and even more so patients with cardiovascular problems may suffer from serious problems when exposed. Moreover, airline travel is nowadays part of daily life and also patients with cardiac diseases want or have to fly. Finally, it may also serve as a model of pulmonary hypertension that may be extrapolated to low altitudes. Thus, several aspects of high altitudes are relevant to a significant proportion of the population.

Various factors are different at high altitudes that may cause problems such as higher ultra violet radiation, lower temperature and rapid changes of the environment. Most importantly, however, oxygen partial pressure is diminished with increasing altitude (table 1) [1], resulting in hypoxia. Some subjects may be susceptible to hypoxia even when healthy. Moreover, hypoxia may be less well tolerated by patients with cardiovascular and pulmonary diseases.

Cardiovascular adaptation to high altitudes
When exposed to reduced oxygen content, there are several changes that occur rapidly [2]. As a consequence, the oxygen content in the arterial blood decreases, i.e. hypoxia occurs (table 1). Because of the S-shape of the haemoglobin dissociation curve, patients will experience oxygen desaturation at elevations above 3000 m. During exercise, however, desaturation may occur at much lower altitudes, particularly in the elderly patient and in patients with cardiovascular and pulmonary diseases [3]. Total oxygen demand for a given workload, however, is independent of altitude. Thus, cardiac output must increase to maintain oxygen delivery for a given workload, given the reduced arterial oxygen content of the blood.

As a result of hypoxia, ventilation increases to compensate for the reduced arterial oxygen content in part. This results in hypocapnia and respiratory alkalosis as well as an increase in 2,3 di-phospho-glycerate (DPG). There is also an increase in sympathetic tone and circulating levels of both noradrenaline and adrenaline, possibly via stimulation of peripheral chemoreceptors. This results in an increase in cardiac output and coronary flow, but also myocardial work and myocardial ox-
Table 1
Pressure at different levels of altitude [1].

<table>
<thead>
<tr>
<th>Altitude in m (ft)</th>
<th>Barometric pressure (mm Hg)</th>
<th>Atmosphere pO₂ (mm Hg)</th>
<th>Alveolar pO₂ (mm Hg)</th>
<th>Arterial pO₂ (mm Hg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sea level</td>
<td>760</td>
<td>1013</td>
<td>159</td>
<td>110</td>
</tr>
<tr>
<td>609 (2000)</td>
<td>711</td>
<td>948</td>
<td>148</td>
<td>99</td>
</tr>
<tr>
<td>1523 (5000)</td>
<td>629</td>
<td>838</td>
<td>131</td>
<td>85</td>
</tr>
<tr>
<td>2438 (8000)</td>
<td>558</td>
<td>744</td>
<td>116</td>
<td>69</td>
</tr>
<tr>
<td>3000</td>
<td>523</td>
<td>679</td>
<td>109</td>
<td>53</td>
</tr>
<tr>
<td>5000</td>
<td>412</td>
<td>549</td>
<td>76</td>
<td>40</td>
</tr>
</tbody>
</table>

There are several factors that may aggravate hypoxia in both healthy and sick. In the latter, however, such aggravation may become clinically relevant since reserves are already reduced. Thus, among the most important aggravating factors are cardiovascular and pulmonary diseases. There are other factors that may be important particularly when patients are exposed to high altitudes. Thus, smoking is very relevant as the uptake of carbon monoxide further reduces O₂ uptake in the blood. Moreover, smoking results in a left-shift of the O₂-dissociation curve, reducing the peripheral availability of oxygen. In addition, alcohol use increases hypoxia, probably due to the reduction in respiratory drive [9].

Physical activity, particularly strenuous exercise reduces the arterial O₂ content as mentioned above (e.g., [8]). This is seen in all subjects, not only in subjects that are susceptible to hypoxia exposure or patients with cardiovascular or pulmonary diseases. Still, it is obviously more relevant in the latter groups. Medication may also influence O₂ saturation when exposed to high altitude, although studies addressing this im-
Exposure to high altitudes in patients with coronary heart disease

The activation of the sympathetic nervous system, together with the above mentioned dehydration, may theoretically activate thrombocytes and, as a consequence, increase the risk of thromboembolic events including acute coronary syndrome. However, based on epidemiological data and small pathophysiological studies, this does not seem to be the case up to altitudes of app. 4000 m [11], although this was obviously not tested in proper prospective trials.

In healthy subjects, coronary flow reserve is increased, even in subjects susceptible to high altitude pulmonary edema (HAPE) that show an exacerbated response of the pulmonary vasculature to high altitudes but not the coronary vasculature. Only immediately prior to HAPE, a reduced flow reserve was seen [12]. In patients with coronary artery disease, however, some although not all studies described a reduced coronary flow reserve at high altitude [13]. Whether this is of clinical relevance is debated. Still, exercise testing during hypoxia in stable patients with coronary artery disease showed little ST-deviation, at least if ischaemia was absent at low to moderate exercise at low altitudes. Also, exercise response did not differ from low altitudes in such patients [14]. Thus, heart rate x blood pressure product did not differ and the same lactate levels were achieved. Most previous, though small studies, showed ST-deviations at a similar heart rate x blood pressure product as at low altitude [11]. A small study even showed that intermittent exposure to hypoxia may reduce the ischaemia score in patients with stable coronary artery disease. These subjects had once a week 4 h exposure to hypoxia starting at simulated altitude of 2400 m with weekly increase of 300 m (total 14 sessions) [15]. Obviously, more studies are required before such an approach can be recommended to reduce ischaemic burden. Nevertheless, limited exposure to hypoxia in patients with stable coronary artery disease seems to be safe, but no experience exists in unstable patients and due to the above mentioned theoretical considerations, such patients may well be at significant risk.

Exposure to high altitudes in patients with heart failure

Patients with HF may tolerate exposure to high altitudes less well than healthy subjects. Obviously, reduced O2 may further reduce already limited exercise capacity, but relative reduction in peak VO2 may even be disproportionate compared to healthy subjects (fig. 2) [16]. The underlying cause of this is not yet precisely known. Patients with HF show an exacerbated respiratory response to exercise even at normoxia, expressed as pathologic increase in VE/VCO2 during
HF not showing pulmonary hypertension may well be exposed to high altitude (up to 3000 m, possibly 3500 m), although data are very limited.

**Risk of arrhythmias at high altitudes**

Again, direct evidence is very scarce and sufficiently supported recommendations cannot be made. However, arrhythmias do not seem to be a major problem in subjects exposed to high altitude. Theoretically, the activation of the sympathetic activity may increase the risk of (serious) arrhythmias. This might be further increased due to the fact that the alkalosis due to hyperventilation results in a shift of potassium to the intracellular space and thereby increases the risk of hypokalaemia. Moreover, pulmonary hypertension and the theoretical increased risk of acute coronary syndrome may contribute to the risk of serious arrhythmias. Clinically, there are reports of increased numbers of both ventricular and supraventricular ectopic beats, but there are only case reports of serious arrhythmias and sudden cardiac death during exposure to high altitude. Patients prone to serious arrhythmias, particularly ventricular tachycardias, however, should not be exposed to altitude above 3000–3500 m.

**Additional aspects when travelling, general recommendations**

Another important aspect when travelling to remote places is the limited access to medical treatment if required. Thus, medication should be carried during

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**Figure 2**

**Figure 3**
Algorithm if patients with cardiovascular diseases travel to high altitudes. (Adapted from [18]: Higgins JP, Tuttle T, Higgins JA. Altitude and the heart: is going high safe for your cardiac patient? Am Heart J. 2010;159:25–32. © Elsevier, reprint with kind permission.
travelling in duplicate, stored at different places. Availability of medical services should be checked prior to travel and decisions whether to travel should be based on this and the potential risk of individual patients.

In light of the lack of data, it is always better to be cautious in case of doubt. In general, patients with cardiac diseases should not go to high altitudes in an unstable disease state [18]. In addition, patients should be advised that they may become symptomatic at an earlier stage, as heart rate × blood pressure product reflects physical load and not workload itself. When stable and well compensated, risk is not significantly increased by the altitude-related hypoxia up to app. 3000 m for patients with HF and 3500 m for patients with stable coronary artery disease; however, patients have to account for the particular circumstances of a stay at high altitudes. A stress test prior to exposure may help to risk stratify patients, as suggested by several small studies in patients with coronary artery disease prior to ascent [11, 18]. Symptoms or ischaemia must not occur at light to moderate exercise. Above 3500 m, there is a considerable decrease of exercise capacity and therefore, in addition to a stable disease, preserved left ventricular function and exercise capacity clearly above normal are required for a safe stay. Even then, patients with cardiovascular heart disease should generally be advised not to go higher than 4500 m. Patients with pulmonary hypertension are very sensitive to hypoxia and even an altitude of 1500 m may cause problems [19]. This limit also applies to unstable patients in general. It is important to note that this does not only refer to patients with pulmonary hypertension class 1 according to WHO classification, but also secondary to other diseases including HF. Figure 3 and table 2 show general recommendations for safe travel to high altitudes in patients with different diseases [18].

Recommendations for flying

When flying in commercial aircraft, cabin pressure is higher than the pressure of the actual altitude. Nevertheless, cabin pressure is lower than pressure at low altitude. Thus, pressure usually corresponds to an altitude between 5000 ft and 7000 ft (i.e., 1524 m and 2134 m), never exceeding 10000 ft (3048 m). There are,

<table>
<thead>
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<th>Table 2</th>
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<tr>
<td>Advice for exposure of high altitude (and air-travelling) in cardiac patients.</td>
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<tr>
<td>Coronary artery disease</td>
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<td>Safe altitude if no or little symptoms</td>
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<td>Limited exposure to max. 2000 m</td>
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<td>Limited exposure to max. 1500 m</td>
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<td>Specific recommendations</td>
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<td>Level of evidence</td>
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<td>Intracardiac shunting</td>
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<td>HAPE susceptibles</td>
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However, additional potential problems related to commercial flights. Thus, immobility, together with the increased risk of dehydration, increases the thrombotic risk. Dehydration occurs because humidity is very low during flights (progressive decrease during flights down to 5–10% relative humidity), since the absolute content of water in the air outside the aircraft is very limited. Moreover, subjects may be stressed during flights with the accompanied increase in sympathetic tone. Patients that are handicapped may have difficulties to board and leave an aircraft, but may also be very limited during flights. Also, possibilities to provide care in case of emergencies in an aircraft are very limited and it may take hours to bring a patient to hospital. Table 2 summarises recommendations for travelling to high altitude, which may also be used (as applicable) for aircraft travelling.

These aspects must be considered when patients want to fly. Patients who do not tolerate altitudes above 2000 m should not fly (e.g., patient with clinically relevant pulmonary hypertension [19]). If flying is absolutely necessary, oxygen supply during the flight is required and airlines should be contacted as early as possible. In general, patients who are not stable are unable to fly (i.e., unstable angina, first two weeks after uncomplicated infarction, six weeks in complicated infarction, first two weeks post coronary catheter interventions [PCI] and at least 3 weeks post bypass surgery, HF NYHA III+; similar recommendations for patients with congenital heart diseases. Cave pulmonary hypertension) [18, 20]. Intake of alcohol may reduce tolerability [9].

Despite the potentially increased risk, severe complications are rarely seen [21, 22]. Thus, death rate was as little as 0.31 per million passengers of which the majority (77%) were not known to have health problems (i.e., 399 of 515 deaths; in additional 62 deaths no information was available on history) [21]. Moreover, only 25% of health care problems among commercial air travelers occurred during flight and required only rarely unscheduled landings (7 in one year at an international airport in the US) [22]. Another study found more acutely ill patients during flight compared to stay at the airport, but with a rate of 0.0002%, it was very low. Cardiac arrest occurred in 0.00015% of passengers in aircraft [23]. Although these are rather old data, it is very unlikely that this picture has changed significantly since. It is likely that precautions taken are, at least in part, responsible for the very low complication rate, indicating that the measures taken are probably efficient, but well required.

**High altitude pulmonary edema (HAPE) – relevant for lowlanders?**

HAPE is a serious problem, accompanied even with mortality when untreated, among subjects staying at altitudes >2500 m without acclimatisation (i.e., ascent within <24h). With increasing altitude, the risk increases and is between 5% and 10% at an altitude of 4500 m in the general population, but 60–70% in those known to be susceptible (HAPE-susceptible). HAPE usually occurs after an overnight stay at high altitudes. It is accompanied by excess pulmonary arterial and to a lesser extent pulmonary venous vasoconstriction. Since vasoconstriction is inhomogeneous, it leaves areas of the pulmonary vasculature unprotected [24]. The individual vascular response to hypoxia might be genetically determined since we found a correlation of the increase in pulmonary artery pressure between fathers and their kids [25].

If HAPE occurs, the person concerned should be brought to lower altitudes as quickly as possible. Oxygen should be given and patients treated with high doses of nifedipine or alternatively other agents that showed to prevent HAPE if given prophylactically. The most important measure to prevent HAPE is acclimatisation. Thus, subjects should not increase more than 300 m/day (sleeping altitude) above 2500 m altitude. This is a general recommendation applicable not only to HAPE-susceptibles, but often not really feasible. Thus, HAPE-susceptibles often require medical prevention which encompasses nifedipine 3 × 20 mg and as alternatives dexamethasone 2 × 8 mg, tadalafil 2 × 10 mg, or inhaled salmeterol 5 × 125 µg, starting the day before ascent [26]. All these measures successfully reduce the risk of HAPE, but the studies were too small to know which of the interventions is most effective.

Research of HAPE-susceptibility may also be of interest to improve understanding of pathophysiology of pulmonary hypertension. Thus, we found that the function of the left ventricle, both systolic and diastolic, as well as perfusion are not altered even when pulmonary pressure is significantly elevated (i.e. >60 mm Hg
systolic pulmonary artery pressure). Moreover, it may shed further light in the limitation of exercise capacity when pulmonary pressure is elevated [8]. More importantly, it may highlight the potential importance of exercise-induced pulmonary hypertension, which is believed to be a relatively rare disease. We found, however, that HAPE-susceptibles show slightly, but significantly dilated right-sided cardiac cavities as compared to matched control subjects (fig. 4) [27]. Since elevation of pulmonary pressure was found to be elevated already at intensities corresponding to daily life activities (increase in systolic pulmonary arterial pressure at 40% of peak exercise 23 ± 6 versus 11 ± 5 mm Hg in non-susceptibles, p <0.001) [27] and the relative commonness of HAPE-susceptibility as mentioned above, exercise-induced might be a significantly underestimated problem. Thus, patients with heart failure and preserved ejection fraction were found to have (slightly) increased pulmonary artery pressures at rest [28]. It may be speculated that there is a link between these two entities which would give new opportunities for treating these patients. Obviously, further research is required to investigate this aspect and to see if HAPE-susceptibility may also be of importance in diseases associated with hypoxia (e.g., COPD). Taken together, susceptibility to HAPE may well be of relevance in various circumstances at low altitude, but further research is required to specifically address this important issue.

Conclusion

Nowadays, exposure to high altitudes is relatively common. Thus, basic knowledge is important also for physicians that are not specialised in this topic. Exercise capacity is limited at high altitude, mainly due to hypoxia. Additional factors (circumstances) may aggravate the problem. Day trips to altitudes up to 3000–3500 m are probably relatively safe even in patients with cardiovascular diseases, unless patients are not stable. Unstable patients are not suited for hypoxia exposure although direct evidence is largely lacking.

Subjects that are susceptible to develop HAPE should avoid rapid ascent to high altitude, or should be treated prophylactically. To what extent research in these subjects is relevant for pathophysiology of pulmonary hypertension at normoxia needs further investigation.

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