Review article

The role of routine pulmonary imaging before hyperbaric oxygen treatment

Connor TA Brenna^{1,2}, Shawn Khan², George Djaiani³, Jay C Buckey Jr.⁴, Rita Katznelson^{1,3}

¹ Department of Anesthesiology and Pain Medicine, University of Toronto, Toronto, ON, Canada

² Faculty of Medicine, University of Toronto, Toronto, ON, Canada

³ Department of Anesthesia, University Health Network, Toronto, ON, Canada

⁴ Department of Medicine, Geisel School of Medicine at Dartmouth, Hanover, NH, USA

Corresponding author: Dr Rita Katznelson, Toronto General Hospital, 200 Elizabeth St, Toronto, ON, M5G 2C4, Canada <u>rita.katznelson@uhn.ca</u>

Keywords

Arterial gas embolism; Lung; Pneumothorax; Pulmonary barotrauma; Radiological imaging; Risk assessment

Abstract

(Brenna CTA, Khan S, Djaiani G, Buckey JC Jr, Katznelson R. The role of routine pulmonary imaging before hyperbaric oxygen treatment. Diving and Hyperbaric Medicine. 2022 30 September;52(3):197–207. <u>doi: 10.28920/dhm52.3.197-207</u>. <u>PMID: 36100931</u>.)

Respiratory injury during or following hyperbaric oxygen treatment (HBOT) is rare, but associated pressure changes can cause iatrogenic pulmonary barotrauma with potentially severe sequelae such as pneumothoraces. Pulmonary blebs, bullae, and other emphysematous airspace abnormalities increase the risk of respiratory complications and are prevalent in otherwise healthy adults. HBOT providers may elect to use chest X-ray routinely as a pre-treatment screening tool to identify these anomalies, particularly if a history of preceding pulmonary disease is identified, but this approach has a low sensitivity and frequently provides false negative results. Computed tomography scans offer greater sensitivity for airspace lesions, but given the high prevalence of incidental and insignificant pulmonary findings among healthy individuals, would lead to a high false positive rate because most lesions are unlikely to pose a hazard during HBOT. Post-mortem and imaging studies of airspace lesion prevalence show that a significant proportion of patients who undergo HBOT likely have pulmonary abnormalities such as blebs and bullae. Nevertheless, pulmonary barotrauma is rare, and occurs mainly in those with known underlying lung pathology. Consequently, routinely using chest X-ray or computed tomography scans as screening tools prior to HBOT for low-risk patients without a pertinent medical history or lack of clinical symptoms of cardiorespiratory disease is of low value. This review outlines published cases of patients experiencing pulmonary barotrauma while undergoing pressurised treatment/testing in a hyperbaric chamber and analyses the relationship between barotrauma and pulmonary findings on imaging prior to or following exposure. A checklist and clinical decision-making tool based on suggested low-risk and high-risk features are offered to guide the use of targeted baseline thoracic imaging prior to HBOT.

Introduction

Hyperbaric oxygen treatment (HBOT) is generally very safe, but adverse events may occur during treatment.¹ Changes in atmospheric pressure during HBOT may cause pulmonary barotrauma (PBt) during the decompression phase of the treatment.^{2,3} Isolated case reports have documented several pressure-change-related respiratory complications with HBOT, including arterial gas embolism (AGE), tension pneumothorax (PTX), and pneumomediastinum.^{4–6} While uncommon, these adverse events are associated with significant morbidity and mortality.

PULMONARY COMPLICATIONS DURING HYPERBARIC OXYGEN TREATMENT

Pulmonary barotrauma during HBOT is rare. Our combined five-year experience (2016–2021) of three North American

HBOT referral centres in Toronto, Canada (University Health Network and Rouge Valley Medical Centre) and Lebanon, NH, USA (Dartmouth-Hitchcock Medical Center), comprising 62,040 treatments performed on 2,250 patients, includes only a single case of PBt. This equates to an incidence of 0.0016% per treatment, or 0.044% per patient.

To review the utility of pre-treatment screening for predicting or preventing PBt during HBOT, we searched for articles describing patients undergoing pressurised treatment/testing in a hyperbaric chamber who had significant findings on pulmonary imaging either before hyperbaric exposure (i.e., pre-existing blebs, bullae, cysts) or afterwards (i.e., barotrauma, gas emboli). The search included several major databases (MEDLINE-Ovid, Embase, Cochrane CENTRAL, and CINAHL) and is detailed in <u>*Appendix 1</u>. A total of 1800 articles were screened independently by two authors (CB and SK) to identify relevant reports, which are described in Tables 1 and 2.

Our search identified 11 reports of respiratory complications after HBOT/hyperbaric exposure with relevant radiological findings as specified above. For those reports where the patients were receiving HBOT, one detailed 126 patients undergoing mechanical ventilation and concurrent HBOT (for a variety of indications), of whom six experienced patient-ventilator asynchrony while in the hyperbaric chamber.7 An additional six single-case studies documented a heterogeneous group of patients aged 5-80 (one female and five males) for whom HBOT was complicated by tension PTX,^{6,8} pulmonary oedema,⁹ pneumomediastinum,¹⁰ acute pulmonary embolism,11 and AGE.12 A final report described a survey of 98 HBOT centres, reporting a combined incidence of PBt of 0.00045%.13 For those cases that involved hyperbaric air exposure (e.g., pressure tolerance testing), one case series described two otherwise healthy individuals who sustained AGE while undertaking routine pressure tolerance testing in a hyperbaric chamber,¹⁴ while another report described a single case of AGE during decompression from a 'dry dive' in a patient with previously undiagnosed pulmonary sarcoidosis.⁵ A final, single case reported the discovery of a bronchopulmonary sequestration determined to contraindicate diving but not HBOT.15 Isolated case reports of underwater divers and passengers on commercial airline flights describe otherwise asymptomatic adults experiencing fatal complications, such as air emboli, when exposed to variable changes in ambient pressure.14-19

Approximately half of the case reports described patients with pre-existing pulmonary comorbidities such as acute respiratory distress syndrome (ARDS), chronic obstructive pulmonary disease (COPD), or pulmonary sarcoidosis, pointing to possible associations with the risk of PBt during HBOT. None of the identified studies reported a significant impact of pre-treatment pulmonary screening on the decision to proceed with HBOT. In fact, in some cases pulmonary pathology was identified prior to HBOT but did not deter treatment (presumably because the centres had previously treated patients with similar pulmonary pathologies, without incident).

CURRENT PRE-HYPERBARIC OXYGEN TREATMENT SCREENING PRACTICES FOR AIRSPACE ABNORMALITIES

The hyperbaric medicine community wants to identify features that predict respiratory injury during treatment, to prevent this adverse event for those at increased risk. Pulmonary bullae, blebs, or cysts – emphysematous pockets of air within the lung parenchyma,^{20,21} may be among these features.¹³ Less commonly, congenital respiratory anomalies

such as bronchogenic cysts and/or bronchopulmonary sequestration can be identified on imaging studies and may be associated with elevated risk of barotrauma with rapid changes in atmospheric pressure.^{15,22,16-19}

Airspace abnormalities are remarkably common in the general population (Table 3). Emphysematous changes and air trapping, once thought to represent high-risk features for pressure-related respiratory complications, are frequently present in individuals without lung disease.²³ Airspaces within the visceral pleural or the subpleural lung itself are classically delineated as blebs or bullae on the basis of diameter (smaller or larger than 1 cm, respectively).²⁴ The prevalence of pulmonary blebs among adults without known pulmonary disease has been reported in two cohort studies, one quoting 6.0% using diagnostic thoracoscopy²⁵ and the other reporting 24.6% using postmortem computed tomography (CT).²⁶ Similarly, pulmonary bullae have been reported in 2.3-5.3% of asymptomatic patients,^{26,27} often coincident with blebs. Other emphysematous changes can be found in the lungs of 14.2–16.1% of adults.^{27,28} A variety of reports have described incidental findings of pulmonary nodules in 0.4-12.8% of adults,²⁸⁻³³ and one quoted as many as 48.4% in a population of cardiac patients imaged using CT angiography.²⁷ Other pulmonary pathologies identified in patients who have experienced barotrauma during HBOT or air dives include pulmonary fibrosis⁴ and sarcoidosis,⁵ which occur in $0.1\%-0.8\%^{34,35}$ and $0.03-0.09\%^{36,37}$ of the general population, respectively. Airspace disease may be even more common among patients undergoing assessment for HBOT than in the general population. For example, the main applications for HBOT are in treating radiation injury for patients who received radiation treatment for head and neck cancers, and these patients are likely to have a smoking history and hence potentially also some degree of COPD.

Patients with these pulmonary aberrations are often clinically asymptomatic, although they have an increased baseline risk of developing a PTX when the volume of gas in these spaces increases, commensurate with a decrease in atmospheric pressure (e.g., when an airplane is ascending to altitude or a hyperbaric chamber is being depressurised).³⁸ Given the increased compliance of these intrathoracic air pockets, patients with these findings who are subjected to pressure changes during HBOT are thought to be at a heightened risk of barotrauma.¹³

Some HBOT centres require a routine chest X-ray (CXR) prior to initiation of treatment to identify patients with pulmonary bullae, blebs, or PTX.^{39,40} Certain centres also pursue additional investigations, such as CT imaging or spirometry, to characterise the nature of pre-existing respiratory disease and assess the risk of injury, while others routinely use CT chest imaging before HBOT.⁸ Other

investigations which may be available to providers evaluating patients for the presence of absence of gas trapping include whole body plethysmography⁴¹ and ventilation scans using xenon,⁴² nitrogen, or helium,⁴³ although these tests may not be available in all centres and the evidence supporting their use in pre-HBOT screening is currently limited. Despite the common practice of obtaining CXR or CT imaging as a screening tool prior to HBOT, the basis for this approach remains unclear. Presently, specific guidance on the use of pulmonary imaging prior to HBOT is not provided by the Undersea and Hyperbaric Medicine Society, the Canadian Undersea and Hyperbaric Medical Association, or the European Committee for Hyperbaric Medicine.

SENSITIVITY AND SPECIFICITY OF PRE-TREATMENT PULMONARY IMAGING

Chest X-ray is the mainstay of pre-HBOT pulmonary screening tools, largely because of its relatively low cost and minimal radiation dose.^{20,44} Its diagnostic sensitivity for minor airspace abnormalities (including bullous and bleb disease) is low,⁴⁴ and relevant pathology may go unrecognised despite screening. CXRs are valuable tools for the diagnosis of pathologies such as consolidation and pleural effusion,44 but have poor interrater reliability and limited specificity for pathologies which do not vary greatly in density.20 The sensitivity of a CXR for even moderately severe and severe emphysematous changes is only 41%,⁴⁵ and minor airspace abnormalities like bullae and blebs can easily be missed. For example, one study describes three cases of pulmonary air cysts missed on CXR, and subsequently found on chest CT scans, in divers who had experienced PBt.⁴⁶ CXR is similarly limited in its ability to detect PTX, with a pooled diagnostic sensitivity of only 52%.47

In fitness-to-dive assessments CXR had a false negative rate of 32% for the identification of relevant intrapulmonary pathology.⁴⁸ The routine use of CXR for the detection of blebs, bullae, cysts, and other airspace disease may not add more to the pre-dive assessment than the individual's medical history.^{46,48} This is likely also true for pre-HBOT screening in patients without any risk factors. In patients for whom a significant concern for pulmonary pathology exists, CT imaging has superior diagnostic accuracy.⁴⁴

High-resolution CT has been proposed as a substitute for CXR in pre-HBOT screening, particularly in subjects with clinical indications.^{8,12,14,48} The radiation exposure associated with high-resolution CT varies dramatically based on imaging parameters but, if performed conservatively, is comparable to a CXR.⁴⁹ While CT is a superior diagnostic tool for airway abnormalities such as pulmonary cysts,²¹ it frequently identifies findings of unknown medical significance.^{23,48} One study conducted in the emergency department setting noted that 33.4% of the general population

had some form of incidental findings on CT imaging, such as pulmonary nodules.⁵⁰ Another, conducting postmortem CT chest scans in a sample of the general population (ages 21-71, without lung disease) reported a 33.8% prevalence of small bullae and/or blebs.²⁶ Incidental, clinically insignificant CT findings may be more prevalent in older patients,⁵¹ and complicate the potential role of CT imaging in 'clearing' patients for HBOT. Additionally, while CT provides information on the presence and size of any relevant pulmonary pathology, it cannot provide guidance on whether the structure can equalise pressure during compression or decompression. While size is an important consideration (larger bullae have higher wall stress and are more likely to rupture than small ones), the relevant consideration for HBOT is whether the structure communicates with the bronchial tree during pressure changes.

CLINICAL INTERPRETATION OF PULMONARY FINDINGS

Because of the shortcomings of imaging modalities available for pre-HBOT screening, how to estimate the risk associated with potential findings is unclear. Many of the studies outlined in Table 1 report respiratory complications of HBOT despite adherence to pre-HBOT imaging protocols and unremarkable imaging studies prior to treatment.^{5,7–10,14} The difficulty associated with interpreting incidental imaging findings is highlighted by two case reports detailing patients whose pre-HBOT imaging identified bullous or bleb disease, but who nonetheless proceeded with HBOT and sustained respiratory complications.^{6,12}

Without clear evidence to discriminate abnormalities representing an elevated risk for PBt from incidental morphology, the utility of pre-HBOT imaging is limited. In a survey of practice patterns among 98 HBOT centres, a majority of centres reported choosing to proceed with treatment for patients in whom pulmonary blebs or bullous lesions were radiologically identified.13 Of those centres which did not, 54% screened patients with a history of lung disease using CXR, while a minority screened those with known pathology using CT, high-resolution CT, or spirometry.¹³ Some of the surveyed centres reported taking additional precautions when treating patients with identifiable blebs or bullous lesions (such as slower compression/decompression rates, pressure limits, and bronchodilator administration).¹³ The applicability of the survey to current practice can be challenged given its age, low response rate (36.8%), and methodological limitations. But among its 98 responding centres, imaging results seldom influenced treatment decisions in a meaningful way.¹³ Nonetheless, PBt was still remarkably infrequent among the surveyed centres, with a reported incidence of 0.00045% or nine instances from approximately 2,000,000 HBOT sessions.13

Table 1

Previous reports of pulmonary complications during hyperbaric oxygen exposures; data are reported as raw numbers unless otherwise noted. *number of responding hyperbaric centres, not numbers of patients; ARDS - acute respiratory distress syndrome; AGE - arterial gas embolism; CO - carbon monoxide; COPD - chronic obstructive pulmonary disease; CT - computed tomography; CXR - chest radiography; DCS - decompression sickness; F - female; HBOT - hyperbaric oxygen treatment; M - male; N/A - not applicable; NR - not reported; PBt - pulmonary barotrauma; PCT - prospective cohort trial; PTX - pneumothorax; PVD - peripheral vascular disease; US - ultrasound

| imaging | Relevant commentary on pre- exposure imaging | CXR is limited in identifying small or anterior PTX with certainty. Chest US and tomodensitometry are both better, and physicians should perform more relevant, non- invasive tests. | Sensitivity of the CT scan in the detection of blebs and bullae is 88%. A bleb or bullae that was not detected on CT may be the reason for PTX in the reported patient. | NR | Routine pre- and post-HBOT CXR in intubated patients may prevent or minimise adverse outcomes related to pneumomediastinum | NR |
|---------------|---|--|---|--|--|---|
| Pulmonary | Impact on exposure | Did not inform treatment decision | Did not inform treatment decision | Did not inform treatment decision | Did not inform treatment decision | N/A |
| | Pulmonary imaging modality | Imaging modality not specified | CXR and CT prior to HBOT – no bullae or blebs. After the 7th session, CT showed total right lung collapse with left mediastinal shift. | Pre-HBOT CXR unremarkable. Repeat CXR after 2nd HBOT session noted pulmonary oedema. | Post-intubation CXR normal. CXR after HBOT found occult pneumo-mediastinum | No pre-HBOT screening. CXR after HBOT and intubation bilateral alveolar perivascular infiltrates. |
| HBOT exposure | Respiratory complications | Patient- ventilator asynchrony (n = 6) | Tension PTX $(n = 1)$ | Pulmonary oedema $(n = 1)$ | Pneumo- mediastinum $(n = 1)$ | Acute pulmonary embolism (n = 1) |
| | Sessions | Mean = 1 | 7 | 7 | 1 | NR |
| | Exposure indication | DCS, CO poisoning, AGE, soft tissue infection, chronic wounds | Lower extremity wounds | CO poisoning | CO poisoning | Diabetic foot ulcer |
| Population | Patient comorbidities | ARDS (23%), mechanical ventilation (100%) | ARDS | ΥΝ | NR | Ischaemic cardio- myopathy, diabetes, PVD |
| | Age, Sex | Mean Age: 57 Sex: M = 78 F = 48 | Age: 28 Sex: M | Age: 31 Sex: M | Age: 5 Sex: M | Age: 80 Sex: M |
| | u | 126 | 1 | - | 1 | 1 |
| Citation, | country, and study design | Bessereau et al. (2017) ⁷ France PCT | Cakmak et al. (2015) ⁸ Turkey Case report | Cho et al. (2018) ⁹ Japan Case report | Jaeger et al. (2013) ¹⁰ USA Case report | Obiagwu et al. (2015) ¹¹ USA Case report |

Table 1 continued.

tolerance testing or a dry dive experience. AGE – arterial gas embolism; CT – computed tomography; CXR – chest radiography; HBOT – hyperbaric oxygen treatment; M – male; N/A – not applicable; PBt – pulmonary barotrauma Previous reports of pulmonary complications during hyperbaric air exposures; data are reported as raw numbers unless otherwise noted. The exposure indication in all cases was routine pressure

| Citation, | | Popu | lation | Hypei | rbaric exposure | | Pulmonary ir | aging |
|---|---|----------------------------------|--|-------|------------------------------|--|---|--|
| country, and study design | u | Age, Sex | Comorbidities | и | Respiratory complications | Pulmonary imaging modality | Impact on exposure | Relevant commentary on pre-exposure imaging |
| Buschmann et al. (2010) ¹⁴ South Africa Case series | 7 | Mean Age: 30 Sex: M = 2 | N/A | 1 | AGE $(n = 2)$ | Case 1: CXR on day 1 was normal. A CT chest on day 2 noted a 2.5 cm right basal subpleural bleb/bulla. Case 2: CT (chest) on day 8 was normal. | Did not inform treatment decision | CXR is commonly performed but based on weak evidence. CT more sensitive, but cost may not be justified with an overall low incidence of PBt/AGE and an unclear relationship between findings and pulmonary risk of barotrauma. Lung compliance rather than anatomical lesions (blebs/bullae), may guide risk. |
| Tan et al. (2020) ¹⁵ Singapore Case report | 1 | Age: 26 Sex: M | N/A | N/A | N/A | Lateral pre-exposure CXR revealed a left lower lobe pulmonary nodule. A chest CT then diagnosed a cavitary left lower lobe (intralobar) broncho-pulmonary sequestration. | Did not inform treatment decision | Bronchopulmonary sequestrations and other air-filled parenchymal lesions should contraindicate diving (but the patient was still considered eligible for HBOT). Although this case supports routine use of lateral CXR in pre-diving health screening, its marginal utility should be weighed against costs (financial, radiation exposure, and false positive rates). |
| Tetzlaff et al. (1999) ⁵ Germany Case report | 1 | Age: 46 Sex: M | Pulmonary sarcoidosis (discovered after hyperbaric exposure) | 1 | AGE $(n = 1)$ | CXR was normal four years pre-exposure. Post-exposure CXR showed bilateral middle and upper lobe infiltrates. CT showed scarring in both lungs. | Did not inform treatment decision | Case illustrates a potential risk of PBt during hyperbaric exposure, even in asymptomatic subjects with normal imaging. Authors emphasise careful evaluation of spirometry and CXR in patients undergoing hyperbaric exposure. |

Table 3

High-risk features in the general population; prevalence of high-risk features for pulmonary complications of hyperbaric oxygen treatment, including pulmonary blebs and bullae, other emphysematous changes, pulmonary fibrosis, and sarcoidosis. Data are reported as percentage of study population or number per 100,000 patients. CAD – coronary artery disease; CXR – chest radiography; CT – computed tomography; NR – not reported

| High-risk feature | Study population | Screening method | Prevalence | Citation |
|------------------------------|--|---------------------------|-----------------------------|--|
| Bulmonory blobs only | Dutch population without pulmonary disorders | Post-mortem CT imaging | 24.6% (32/130) | de Bakker et al. $(2020)^{26}$ |
| r unitoliar y blebs only | Young healthy adults | Thoracoscopy | 6.0% (15/250) | Amjadi et al. (2007) ²⁵ |
| Pulmonary blebs and bullae | Dutch population without pulmonary disorders | Post-mortem CT imaging | 6.9% (9/130) | de Bakker et al. $(2020)^{26}$ |
| Pulmonary bullae | Dutch population without pulmonary disorders | Post-mortem CT imaging | 2.3% (3/130) | de Bakker et al. $(2020)^{26}$ |
| only | Patients with CAD | CT angiography | 5.8% (10/171) | Yorgun et al. (2010) ²⁷ |
| Other emphysematous | Adult trauma patients | Spiral CT | 14.2% (297/2092) | Barrett et al. $(2009)^{28}$ |
| changes | Patients with CAD | CT angiography | 16.4% (28/171) | Yorgun et al. (2010) ²⁷ |
| | Adult trauma patients | Spiral CT | 10.9% (229/2092) | Barrett et al. $(2009)^{28}$ |
| | General population | CT angiography | 12.8% (33/258) | Gil et al. (2007) ²⁹ |
| | Cardiac patients | Electron- beam CT | 4.8% (65/1356) | Horton et al. $(2002)^{30}$ |
| Incidental pulmonary nodules | Cardiac patients | Electron- beam CT | 0.44% (8/1812) | Hunold et al. $(2001)^{31}$ |
| | Cardiac patients | Cardiac CT | 2.4% (4/166) | Haller et al. (2006) ³² |
| | Cardiac patients | Cardiac CT | 6.6% (33/503) | Onuma et al. (2006) ³³ |
| | Cardiac patients | CT angiography | 48.5% (83/171) | Yorgun et al. (2010) ²⁷ |
| Dulman en finacia | General Population (Quebec, Canada) | NR | 0.08% (76/100,000) | Tarride et al. $(2018)^{34}$ |
| Pulmonary librosis | General population, ages 16–84 (USA) | NR | 0.0099% (9.85/100,000) | Raghu et al. (2016) ³⁵ |
| Samaidasis | General adult population (USA) | NR | 0.88% (29,372/3,340,000) | Baughman et al. $(2016)^{36}$ |
| Sarcoldosis | General population, ages 20–69 (USA) | CXR or histology | 0.03% (259/830,891) | Rybicki et al. (1997) ³⁷ |

Considering the relatively high incidence of otherwisebenign pulmonary lesions in the general population and the low incidence of pulmonary complications following HBOT, we can conclude that many patients with pulmonary abnormalities are routinely undergoing HBOT without any observed complications. A patient with relevant pulmonary abnormalities is more likely to have unremarkable pretreatment CXR imaging than they are to experience iatrogenic pulmonary complications during HBOT. Based on an incidence for PBt during HBOT of 0.00045%,¹³ the number needed to treat (NNT) to prevent one case of barotrauma would be 2,222 if there was a perfectly sensitive and specific tool to identify patients certain to experience that complication. In reality, the NNT must be much higher to account for both the limitations in CXR sensitivity and the unknown likelihood that an identified abnormality will predispose to barotrauma. In current practice, if patients identified as having radiological risk factors for PBt are not actually excluded from HBOT, the NNT of CXR is infinity.

Table 4

Low-risk features of patient history and physical exam which may be reassuring of low-risk for pulmonary complications following hyperbaric oxygen treatment. ARDS – acute respiratory distress syndrome; COPD – chronic obstructive pulmonary disease; HBOT – hyperbaric oxygen treatment PTX – pneumothorax

| Possible low-risk features |
|--|
| No history, symptoms, or physical exam findings of asthma, COPD, pulmonary fibrosis, sarcoidosis, PTX, or ARDS |
| Unremarkable thoracic imaging, if previously performed and available for review |
| Previous HBOT without incident |
| History of scuba diving or air travel without incident |
| Age < 40 years |
| Non-smoker |

Based on the current evidence, we suggest that thousands of patients would have to undergo pulmonary imaging – with its own associated costs and risks – to prevent one from undergoing HBOT and developing PBt. The process would also exclude patients who would not have otherwise sustained barotrauma, and could also include some who would suffer it nonetheless. Given the challenges in applying imaging findings to the clinical determination of which patients are safe to endure hyperbaric conditions, we suggest that pre-HBOT imaging adds very little to a thorough history and physical exam in low-risk populations.

RISK STRATIFICATION BEYOND PRE-TREATMENT IMAGING

Pre-HBOT imaging can (sometimes) provide information on whether a patient has intrathoracic anatomical abnormalities, but it can offer little guidance on whether those abnormalities are likely to cause problems in the hyperbaric chamber. We instead draw on common features of patients reported as having experienced PBt during HBOT in the scientific literature^{4–12,14,15} to suggest a checklist of possible clinical indicators of relatively low risk for pulmonary complications of HBOT (for whom imaging may have the least to offer). These features include: the absence of pre-existing obstructive lung diseases, restrictive lung diseases, PTX, or ARDS; a history of HBOT, scuba diving, or air travel without incident; age younger than 40 years; and non-smoking (Table 4). When prior thoracic imaging is available, especially if it is recently performed, it should be reviewed. The risk factors for spontaneous pneumothoraces or emphysematous lung changes (e.g., younger age, male sex, low body mass index, pulmonary infection, and cigarette and/or marijuana smoking),^{46,52–54} which are themselves predictors of PBt during HBOT, may also indirectly inform hyperbaric exposure risk. Based on the available evidence, and clinical experience, a practical clinical risk tool is provided in <u>*Appendix 2</u> (in the form of a questionnaire) to support clinicians' and patients' decisions to pursue or forego chest imaging prior to HBOT.

ROUTINE PULMONARY SCREENING BEFORE OTHER VOCATIONAL OR RECREATIONAL HYPERBARIC EXPOSURES

While the present article focuses on pulmonary screening prior to HBOT, its findings are applicable to medical assessments preceding other hyperbaric exposures. Pulmonary barotrauma occurring in divers is well described,^{55,56} and the risk can be extrapolated to others working in environments prone to rapid atmospheric compression and decompression, such as caisson or compressed air workers. The incidence of PBt in these groups has not been clearly defined, but reports of affected divers have identified several risk factors including airway obstruction, pre-existing respiratory disease or structural parenchymal abnormality (e.g., bullae or blebs), or a reduced mid-expiratory flow at 25% of vital capacity.^{57–59}

Despite the risk of PBT associated with compression and decompression in these contexts, whether pulmonary imaging is required as part of the standard medical assessment of prospective commercial or recreational divers remains controversial. Recognising the low yield of a screening CXR, the guidelines of some national organisations (such as the UK Health and Safety Executive)⁶⁰ and many major sources of knowledge in the field suggest that CXR is not a requirement unless justified by heightened individual risk.^{61,62} Others, in contrast, have suggested that there is a role for routine CXR screening for all prospective divers,63 or at least for professional divers/diving instructors.46,64 When pulmonary screening is warranted by local policy or a high index of suspicion for PBT-predisposing factors, highresolution CT imaging has been advocated as a potential tool for the initial examination of divers,^{46,65} although this is not currently practical in many settings.

The pre-HBOT risk stratification checklist presented in Table 4 overlaps with, and can be supplemented by, the known risk factors identified for PBt among divers such as pre-existing respiratory disease and blebs/bullae.^{57,58} However, the precise risk profiles of HBOT and other hyperbaric exposures may differ. For example, compression/

decompression injury during diving typically involves much faster pressure change and relates largely to nitrogen, which is inert and less soluble, while oxygen (in HBOT) is more soluble and metabolically consumed. These differences may help explain the relative rarity of AGE during HBOT, which we found reported in only two case studies.^{4,12}

LIMITATIONS

The core limitation of this review is its susceptibility to publication bias. Cases where pulmonary complications were avoided via the identification of bullae or blebs on pre-HBOT imaging are almost certainly under-reported in the literature, although survey data suggest most centres do not consider CXR findings of bullae or blebs to be an absolute contraindication to HBOT, and routinely proceed with treatment – with a very low overall incidence of PBt.¹³ This core limitation could be overcome in the future by using an international multicentre hyperbaric oxygen treatment registry⁶⁶ designed to collect and analyse outcomes and complications related to HBOT exposures.

Conclusions

This review highlights the limitations of routine pulmonary imaging as a screening tool prior to HBOT. Reports of PBt during HBOT often describe patients with known pre-existing pulmonary pathology (e.g., asthma, COPD, pulmonary fibrosis, sarcoidosis, PTX, or ARDS) or occult intrathoracic abnormalities (e.g., bullous lesions or blebs). Abnormalities which might be considered to increase the risk of pulmonary complication during HBOT are common, even among otherwise healthy individuals without any pulmonary disease. Importantly, normal pre-HBOT CXR does not preclude patients from developing barotrauma. The use of routine imaging prior to HBOT does not provide a reliable way to reduce the risk of iatrogenic injury in low-risk populations. In high-risk patients or when clinical findings are unclear (e.g., unable to rule out a PTX), high-resolution CT imaging may be a superior test for the identification of airway or parenchymal lung disease in carefully selected patients. The presence of an abnormality on CT scan, however, does not provide a dependable measure of whether the lesion might rupture or leak with changes in atmospheric pressure. Ultimately, the provider will need to use clinical judgement when determining how to proceed for patients deemed high-risk for respiratory complications of HBOT.

A thorough approach to patients' past medical histories and physical examinations are more relevant steps in assessing the risk for iatrogenic respiratory complications related to HBOT. Further research is needed to characterise how specific features of patients' demographic and past medical history may influence the risk of iatrogenic lung injury during HBOT. This review suggests that, for lowrisk individuals, HBOT can proceed without pre-treatment chest imaging.

References

- Hadanny A, Meir O, Bechor Y, Fishlev G, Bergan J, Efrati S. The safety of hyperbaric oxygen treatmentretrospective analysis in 2,334 patients. Undersea Hyperb Med. 2016;43:113–22. PMID: 27265988.
- 2 Hamilton-Farrell M, Bhattacharyya A. Barotrauma. Injury. 2004;35:359–70. doi: 10.1016/j.injury.2003.08.020. PMID: 15037370.
- 3 Hillman K. Pulmonary barotrauma. Clinics in Anaesthesiology. 1985;3:877–98. doi: 10.1016/S0261-9881(21)00097-5.
- 4 Wolf HK, Moon RE, Mitchell PR, Burger PC. Barotrauma and air embolism in hyperbaric oxygen therapy. Am J Forensic Med Pathol. 1990;11:149–53. doi: 10.1097/00000433-199006000-00009. PMID: 2343842.
- 5 Tetzlaff K, Reuter M, Kampen J, Lorr C. Hyperbaric chamber-related decompression illness in a patient with asymptomatic pulmonary sarcoidosis. Aviat Space Environ Med. 1999;70:594–7. <u>PMID: 10373052</u>.
- 6 Unsworth IP. Case report. Pulmonary barotrauma in a hyperbaric chamber. Anaesthesia. 1973;28:675–8. doi: 10.1111/J.1365-2044.1973.TB00555.X. PMID: 4759876.
- 7 Bessereau J, Aboab J, Hullin T, Huon-Bessereau A, Bourgeois JL, Brun PM, et al. Safety of hyperbaric oxygen therapy in mechanically ventilated patients. Int Marit Health. 2017;68:46–51. doi: 10.5603/IMH.2017.0008. PMID: 28357836.
- 8 Cakmak T, Battal B, Kara K, Metin S, Demirbas S, Yildiz S, et al. A case of tension pneumothorax during hyperbaric oxygen therapy in an earthquake survivor with crush injury complicated by ARDS (adult respiratory distress syndrome). Undersea Hyperb Med. 2015;42:9–13. PMID: 26094299.
- 9 Cho K, Minami T, Okuno Y, Kakuda Y, Tsutsumi T, Kogame T, et al. Convulsive seizure and pulmonary edema during hyperbaric oxygen therapy: a case report. J Med Invest. 2018;65:286–8. doi: 10.2152/JMI.65.286. PMID: 30282875.
- 10 Jaeger NJ, Brosious JP, Gustavson RB, McFarlin CA, Gearhart W, Zamboni WA, et al. Pneumomediastinum following hyperbaric oxygen therapy for carbon monoxide poisoning: case report. Undersea Hyperb Med. 2013;40:521–3. <u>PMID:</u> 24377195.
- 11 Obiagwu C, Paul V, Chadha S, Hollander G, Shani J. Acute pulmonary edema secondary to hyperbaric oxygen therapy. Oxf Med Case Reports. 2015;2015:183–4. doi: 10.1093/omcr/ omv002. PMID: 25988073. PMCID: PMC4370014.
- 12 Rivalland G, Mitchell SJ, van Schalkwyk JM. Pulmonary barotrauma and cerebral arterial gas embolism during hyperbaric oxygen therapy. Aviat Space Environ Med. 2010;81:888–90. doi: 10.3357/asem.2783.2010. PMID: 20824998.
- 13 Toklu AS, Korpinar S, Erelel M, Uzun G, Yildiz S. Are pulmonary bleb and bullae a contraindication for hyperbaric oxygen treatment? Respir Med. 2008;102:1145–7. doi: 10.1016/j.rmed.2008.03.012. PMID: 18571913.
- 14 Buschmann DK. Arterial gas embolism during pressure tolerance testing in a hyperbaric chamber: a report of two cases. Aviat Space Environ Med. 2010;81:1133–6. doi: 10.3357/asem.2842.2010. PMID: 21197859.
- 15 Tan TXZ, Li AY, Sng JJ, Lim M, Tan ZX, Ang HX, et al. A diver's dilemma a case report on bronchopulmonary sequestration. BMC Pulm Med. 2020;20:121. doi: 10.1186/S12890-020-1159-1. PMID: 32366303. PMCID: PMC7199314.
- 16 Closon M, Vivier E, Breynaert C, Duperret S, Branche P,

Coulon A, et al. Air embolism during an aircraft flight in a passenger with a pulmonary cyst: a favorable outcome with hyperbaric therapy. Anesthesiology. 2004;101:539–42. doi: 10.1097/00000542-200408000-00037. PMID: 15277939.

- 17 Arnaiz J, Marco de Lucas E, Piedra T, Arnaiz Garcia ME, Patel AD, Gutierrez A. In-flight seizures and fatal air embolism: the importance of a chest radiograph. Arch Neurol. 2011;68:661– 4. doi: 10.1001/archneurol.2011.85. PMID: 21555644.
- 18 Raymond LW. Pulmonary barotrauma and related events in divers. Chest. 1995;107:1648–52. doi: 10.1378/ chest.107.6.1648. PMID: 7781361.
- 19 Zaugg M, Kaplan V, Widmer U, Baumann PC, Russi EW. Fatal air embolism in an airplane passenger with a giant intrapulmonary bronchogenic cyst. Am J Respir Crit Care Med. 1998;157:1686–9. doi: 10.1164/ajrccm.157.5.9706040. PMID: 9603155.
- 20 Martini K, Frauenfelder T. Advances in imaging for lung emphysema. Ann Transl Med. 2020;8:1467. doi: 10.21037/ ATM.2020.04.44. PMID: 33313212. PMCID: PMC7723580.
- 21 Raoof S, Bondalapati P, Vydyula R, Ryu JH, Gupta N, Raoof S, et al. Cystic lung diseases: Algorithmic approach. Chest. 2016;150:945–65. doi: 10.1016/j.chest.2016.04.026. PMID: 27180915. PMCID: PMC7534033.
- 22 St-Georges R, Deslauriers J, Duranceau A, Vaillancourt R, Deschamps C, Beauchamp G, et al. Clinical spectrum of bronchogenic cysts of the mediastinum and lung in the adult. Ann Thorac Surg. 1991;52:6–13. doi: 10.1016/0003-4975(91)91409-0. PMID: 2069465.
- 23 Mets OM, van Hulst RA, Jacobs C, van Ginneken B, de Jong PA. Normal range of emphysema and air trapping on CT in young men. AJR Am J Roentgenol. 2012;199:336–40. doi: 10.2214/AJR.11.7808. PMID: 22826394.
- 24 Hansell DM, Bankier AA, MacMahon H, McLoud TC, Müller NL, Remy J. Fleischner Society: glossary of terms for thoracic imaging. Radiology. 2008;246:697–722. doi: 10.1148/radiol.2462070712. PMID: 18195376.
- 25 Amjadi K, Alvarez GG, Vanderhelst E, Velkeniers B, Lam M, Noppen M. The prevalence of blebs or bullae among young healthy adults: a thoracoscopic investigation. Chest. 2007;132:1140–5. doi: 10.1378/chest.07-0029. PMID: 17890475.
- 26 de Bakker HM, Tijsterman M, de Bakker-Teunissen OJG, Soerdjbalie-Maikoe V, van Hulst RA, de Bakker BS. Prevalence of pulmonary bullae and blebs in postmortem CT imaging with potential implications for diving medicine. Chest. 2020;157:916–23. <u>doi: 10.1016/j.chest.2019.11.008</u>. <u>PMID: 31759963</u>.
- 27 Yorgun H, Kaya EB, Hazirolan T, Ateş AH, Canpolat U, Sunman H, et al. Prevalence of incidental pulmonary findings and early follow-up results in patients undergoing dual-source 64-slice computed tomography coronary angiography. J Comput Assist Tomogr. 2010;34:296–301. <u>doi: 10.1097/</u> <u>RCT.0b013e3181c1d0e4. PMID: 20351524</u>.
- 28 Barrett TW, Schierling M, Zhou C, Colfax JD, Russ S, Conatser P, et al. Prevalence of incidental findings in trauma patients detected by computed tomography imaging. Am J Emerg Med. 2009;27:428–35. doi: 10.1016/j.ajem.2008.03.025. PMID: 19555613.
- 29 Gil BN, Ran K, Tamar G, Shmuell F, Eli A. Prevalence of significant noncardiac findings on coronary multidetector computed tomography angiography in asymptomatic patients. J Comput Assist Tomogr. 2007;31:1–4. doi: 10.1097/01. rct.0000233125.83184.33. PMID: 17259825.

- 30 Horton KM, Post WS, Blumenthal RS, Fishman EK. Prevalence of significant noncardiac findings on electron-beam computed tomography coronary artery calcium screening examinations. Circulation. 2002;106:532–4. doi: 10.1161/01. cir.0000027136.56615.de. PMID: 12147531.
- 31 Hunold P, Schmermund A, Seibel RM, Grönemeyer DH, Erbel R. Prevalence and clinical significance of accidental findings in electron-beam tomographic scans for coronary artery calcification. Eur Heart J. 2001;22:1748–58. <u>doi: 10.1053/ euhj.2000.2586. PMID: 11511125</u>.
- 32 Haller S, Kaiser C, Buser P, Bongartz G, Bremerich J. Coronary artery imaging with contrast-enhanced MDCT: extracardiac findings. AJR Am J Roentgenol. 2006;187:105– 10. doi: 10.2214/AJR.04.1988. PMID: 16794163.
- 33 Onuma Y, Tanabe K, Nakazawa G, Aoki J, Nakajima H, Ibukuro K, Hara K. Noncardiac findings in cardiac imaging with multidetector computed tomography. J Am Coll Cardiol. 2006;48:402–6. doi: 10.1016/j.jacc.2006.04.071. PMID: 16843193.
- 34 Tarride JE, Hopkins RB, Burke N, Guertin JR, O'Reilly D, Fell CD, et al. Clinical and economic burden of idiopathic pulmonary fibrosis in Quebec, Canada. Clinicoecon Outcomes Res. 2018;10:127–37. doi: 10.2147/CEOR.S154323. PMID: 29503576. PMCID: PMC5826203.
- 35 Raghu G, Weycker D, Edelsberg J, Bradford WZ, Oster G. Incidence and prevalence of idiopathic pulmonary fibrosis. Am J Respir Crit Care Med. 2006;174:810–6. <u>doi: 10.1164/</u> rccm.200602-163OC. PMID: 16809633.
- 36 Baughman RP, Field S, Costabel U, Crystal RG, Culver DA, Drent M, et al. Sarcoidosis in America. Analysis based on health care use. Ann Am Thorac Soc. 2016;13:1244–52. doi: 10.1513/AnnalsATS.201511-7600C. PMID: 27509154.
- 37 Rybicki BA, Major M, Popovich J, Maliarik MJ, Iannuzzi MC. Racial differences in sarcoidosis incidence: a 5-year study in a health maintenance organization. Am J Epidemiol. 1997;145:234–41. doi: 10.1093/oxfordjournals.aje.a009096. PMID: 9012596.
- 38 de Menezes Lyra R. Etiology of primary spontaneous pneumothorax. J Bras Pneumol. 2016;42:222–6. doi: 10.1590/ S1806-37562015000000230. PMID: 27383937. PMCID: PMC5569604.
- 39 Dey D, Kochhar H, Rao K, Venkatesh S, Verma R. The rationale of carrying out specific investigations prior to hyperbaric oxygen therapy. Indian J Aerosp Med. 2013;57:56–65.
- 40 Heyboer M 3rd. Hyperbaric oxygen therapy side effects where do we stand? J Am Coll Clin Wound Spec. 2018;8:2–3. doi: 10.1016/j.jccw.2018.01.005. PMID: 30276115. PMCID: PMC6161636.
- 41 Sylvester KP, Clayton N, Cliff I, Hepple M, Kendrick A, Kirkby J, et al. ARTP statement on pulmonary function testing 2020. BMJ Open Respir Res. 2020;7:e000575. doi: 10.1136/bmjresp-2020-000575. PMID: 32631927. PMCID: PMC7337892.
- 42 Suga K. Technical and analytical advances in pulmonary ventilation SPECT with xenon-133 gas and Tc-99m-Technegas. Ann Nucl Med. 2002;16:303–10. doi: 10.1007/ BF02988614. PMID: 12230089.
- 43 Chiumello D, Cressoni M, Chierichetti M, Tallarini F, Botticelli M, Berto V, et al. Nitrogen washout/washin, helium dilution and computed tomography in the assessment of end expiratory lung volume. Crit Care. 2008;12:R150. doi: 10.1186/cc7139. PMID: 19046447. PMCID: PMC2646315.
- 44 Candemir S, Antani S. A review on lung boundary detection in

chest X-rays. Int J Comput Assist Radiol Surg. 2019;14:563– 76. <u>doi: 10.1007/S11548-019-01917-1</u>. <u>PMID: 30730032</u>. <u>PMCID: PMC6420899</u>.

- 45 Thurlbeck WM, Simon G. Radiographic appearance of the chest in emphysema. AJR Am J Roentgenol. 1978;130:429– 40. doi: 10.2214/AJR.130.3.429. PMID: 415543.
- 46 Toklu AS, Kiyan E, Aktas S, Cimsit M. Should computed chest tomography be recommended in the medical certification of professional divers? A report of three cases with pulmonary air cysts. Occup Environ Med. 2003;60:606–8. doi: 10.1136/ oem.60.8.606. PMID: 12883024. PMCID: PMC1740603.
- 47 Ding W, Shen Y, Yang J, He X, Zhang M. Diagnosis of pneumothorax by radiography and ultrasonography: a meta-analysis. Chest. 2011;140:859–66. <u>doi: 10.1378/ chest.10-2946. PMID: 21546439</u>.
- 48 Wingelaar TT, Bakker L, Nap FJ, van Ooij PJAM, Endert EL, van Hulst RA. Routine chest X-rays are inaccurate in detecting relevant intrapulmonary anomalies during medical assessments of fitness to dive. Front Physiol. 2021;11:613398. doi: 10.3389/fphys.2020.613398. PMID: 33488401. PMCID: PMC7816860.
- 49 Foley SJ, McEntee MF, Rainford LA. Establishment of CT diagnostic reference levels in Ireland. Br J Radiol. 2012;85:1390–7. doi: 10.1259/bjr/15839549. PMID: 22595497. PMCID: PMC3474022.
- 50 Thompson RJ, Wojcik SM, Grant WD, Ko PY. Incidental findings on CT scans in the emergency department. Emerg Med Int. 2011;2011:624847. doi: 10.1155/2011/624847. PMID: 22046542. PMCID: PMC3200145.
- 51 Copley SJ, Wells AU, Hawtin KE, Gibson DJ, Hodson JM, Jacques AET, et al. Lung morphology in the elderly: comparative CT study of subjects over 75 years old versus those under 55 years old. Radiology. 2009;251:566–73. doi: 10.1148/radiol.2512081242. PMID: 19401580.
- 52 Stern EJ, Webb WR, Weinacker A, Muller NL. Idiopathic giant bullous emphysema (vanishing lung syndrome): imaging findings in nine patients. AJR Am J Roentgenol. 1994;162:279–82. doi: 10.2214/AJR.162.2.8310909. PMID: 8310909.
- 53 Ohtsuka Y, Homma Y, Ukita H, Masaki Y, Doi I, Ohe M, et al. Clinical characteristics of idiopathic interstitial pneumonia (IIP) with bullae. Intern Med. 1994;33:6–9. doi: 10.2169/ internalmedicine.33.6. PMID: 8180446.
- 54 Johnson MK, Smith RP, Morrison D, Laszlo G, White RJ. Large lung bullae in marijuana smokers. Thorax. 2000;55:340–2. doi: 10.1136/thorax.55.4.340. PMID: 10722775. PMCID: PMC1745720.
- 55 Raymond LW. Pulmonary barotrauma and related events in divers. Chest. 1995;107:1648–52. <u>doi: 10.1378/ chest.107.6.1648</u>. <u>PMID: 7781361</u>.
- 56 Harker CP, Neuman TS, Olson LK, Jacoby I, Santos A. The roentgenographic findings associated with air embolism in sport scuba divers. J Emerg Med. 1993;11:443–9. doi: 10.1016/0736-4679(93)90248-6. PMID: 8228108.

- 57 Tetzlaff K, Reuter M, Leplow B, Heller M, Bettinghausen E. Risk factors for pulmonary barotrauma in divers. Chest. 1997;112:654–9. doi: 10.1378/chest.112.3.654. PMID: 9315797.
- 58 Goffinet CMJ, Simpson G. Cerebral arterial gas embolism in a scuba diver with a primary lung bulla. Diving Hyperb Med. 2019;49:141–4. <u>doi: 10.28920/dhm49.2.141-144</u>. <u>PMID:</u> <u>31177521</u>. <u>PMCID: PMC6704005</u>.
- 59 Russi EW. Diving and the risk of barotrauma. Thorax. 1998;53:S20–S24. <u>doi:10.1136/THX.53.2008.S20</u>. PMID: 10193343. PMCID: PMC1765901.
- 60 UK Health and Safety Executive (HSE). The medical examination and assessment of commercial divers (MA1), 4th ed. 2015. p. 12. [cited 2022 Aug 6]. Available from: <u>https://www.hse.gov.uk/pubns/ma1.pdf</u>.
- 61 Edmonds C, Bennett M, Lippmann J, Mitchell SJ. Diving and subaquatic medicine, 5th ed. Boca Raton (FL): CRC Press; 2015. p. 563–9. doi: 10.1201/B18700.
- 62 Bove AA, Davis JC. Diving medicine, 4th ed. Philadelphia (PA): Saunders; 2003. p. 537.
- 63 Krzyżak J, Korzeniewski K. Medical assessment of fitness to dive. Part I. Int Marit Health. 2021;72:36–45. <u>doi: 10.5603/</u> <u>MH.2021.0005</u>. <u>PMID: 33829471</u>.
- Godden D, Currie G, Denison D, Farrell P, Ross J, Stephenson R, et al. British Thoracic Society guidelines on respiratory aspects of fitness for diving. Thorax. 2003;58:3–13. doi: 10.1136/thorax.58.1.3. PMID: 12511710. PMCID: PMC1746450.
- 65 Reuter M, Tetzlaff K, Warninghoff V, Steffens JC, Bettinghausen E, Heller M. Computed tomography of the chest in diving-related pulmonary barotrauma. Br J Radiol. 1997;70:440–5. doi: 10.1259/BJR.70.833.9227223. PMID: 9227223.
- 66 Harlan NP, Ptak JA, Rees JR, Cowan DR, Fellows AM, Kertis JA, et al. Development of an international, multicenter, hyperbaric oxygen treatment registry and research consortium: protocol for outcome data collection and analysis. JMIR Res Protoc. 2020;9:e18857. doi: 10.2196/18857. PMID: 32579537. PMCID: PMC7459436.

Conflicts of interest and funding

The Multicenter Registry for Hyperbaric Oxygen Therapy (International Registered Report Identifier RR2-10.2196/18857) received support from the Dartmouth-Hitchcock Medical Center Department of Medicine Scholarship Enhancement in Academic Medicine (SEAM) Award Program. No conflicts of interest were declared.

Submitted: 19 April 2022 Accepted after revision: 5 August 2022

Copyright: This article is the copyright of the authors who grant Diving and Hyperbaric Medicine a non-exclusive licence to publish the article in electronic and other forms.