Pulmonary vascular complications of chronic liver disease: Pathophysiology, imaging, and treatment

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Abstract:
To review the pathogenesis of pulmonary vascular complications of liver disease, we discuss their clinical implications, and therapeutic considerations, with emphasis on potential reversibility of the hepatopulmonary syndrome after liver transplantation. In this review, we also discuss the role of imaging in pulmonary vascular complications associated with liver disease.

Key words:
Hepatopulmonary syndrome, portopulmonary hypertension, pulmonary arteriovenous shunts, Yttrium-90 microsphere embolization hepatocellular carcinoma

Our objective was to review current knowledge and advances made in understanding pulmonary vascular complications of liver disease between 2000 and 2010. The MEDLINE data base was explored through PubMed, and bibliographies of 229 papers were reviewed. The selected papers included case studies and series reporting results from patients with the vascular complication of chronic liver disease (CLD). We added our own radiological and clinical experience during this period. As the success of liver transplantation (OLT) is the only available definitive treatment for CLD and its success is dependent on the presence or absence of pulmonary vascular complications associated with CLD; our primary search only included studies emphasizing these findings. Of importance are the hepatopulmonary syndrome (HPS) and portopulmonary hypertension (PoH). HPS is defined as a triad of liver disease, IPVD, and a defect in oxygenation. The HPS is characterized by a defect in arterial perfusion mismatch.[3] Hypoxia occurs as a result of alteration in the production or clearance of chemical mediators causing intrapulmonary vasodilatation (IPVD) and significant arteriovenous shunting (AVS) through the lungs. A major complication of IPVD is the ventilation-perfusion mismatch.[5] Hypoxia occurs as a result of inability of oxygen to diffuse through the markedly dilated lung capillaries. The capillaries are known to dilate to 15–500 µm (n = 8–15 µm) in HPS.[3] Imbalance in the endothelin receptor response, pulmonary microvascular remodelling, and genetic predisposition is thought to lead to IPVD.[6]

HPS has three major components, i.e. liver disease, IVPD, and a defect in oxygenation. Increased alveolar-arterial gradient occurs in HPS while breathing room air.[13] The major defect that underlies HPS is hypoxemia secondary to IPVD associated with AVS. AVS occurs through...
Due to pulmonary capillaries dilatation oxygen encounters difficulty in diffusing into the centre of the larger capillaries. Increased cardiac output and the associated reduced transition time of blood through the pulmonary vascular bed also impair diffusion, leading to a diffusion–perfusion defect or alveolar capillary oxygen disequilibrium.\cite{11}

Early investigators attributed IVPD in HPS to inadequate synthesis or metabolism of pulmonary vasoactive substances by a failing liver.\cite{7,8} However, it has been shown that nitric oxide, a potent vasodilator, is exhaled in larger quantities in patients with cirrhoses related HPS and normalizes following OLT.\cite{8} A single case report supports this theory where NG-LNAME, a nitric oxide production inhibitor, transiently improved HPS.\cite{14} Similarly intravenous methylene blue given to patients with HPS experienced improvements in gas exchange accompanied by an increase in pulmonary vascular resistance.

Gram-negative intestinal endotoxemia in PoH associated with increased nitric oxide release and tumor necrosis factor have been proposed as a cause of excessive pulmonary vasodilatation.\cite{15} Improvement in arterial oxygenation has been reported following antibiotic therapy.\cite{16}

A recent study has thrown doubt on vasodilatation being the sole cause of the pathophysiology underlying HPS. It has been shown that administration of inhaled l-NAME in HPS patients did not improve V/Q mismatch and arterial hypoxemia.\cite{17} This study suggests that vasodilatation alone is unlikely to explain the more than 10-fold increase in capillary diameters in HPS especially as there is little smooth muscle to relax in normal capillaries.

**Clinical presentation**

Typically HPS presents with dyspnea of insidious onset, platypnea and orthodeoxia.\cite{18} Physical examination may reveal dyspnea associated with cyanosis in 90% of all cases. The presence of clubbing has the highest positive predictive value (75%) and dyspnea the highest negative predictive value (100%) for HPS. Spider nevi are a common clinical feature of patients with HPS with significant relationship between cutaneous spider angiomata and systemic and pulmonary vasodilatation suggesting that spider nevi may represent a cutaneous marker for intrapulmonary vascular dilatation.\cite{19,20}

Two types HPS have been described: Type I lesions are associated with vascular dilatations at the precapillary level close to the normal gas exchange units of the lung and Type II with focal larger dilatations amounting to AVS distant from the gas exchange units. Supplementary oxygen improves Type I HPS PaO2 but not Type 2 HPS.\cite{1}

**Prognosis**

HPS is associated with an increase risk of death, worse functional status, and quality of life in patients evaluated for OLT.\cite{21} The median survival time in cirrhotic patients with HPS has been reported as 10.6 months compared to 40.8 months in cirrhotic patients without HPS. The leading cause of death is hemorrhagic shock secondary to gastrointestinal bleeding. The degree of arterial hypoxemia affects survival. Survival is worse with lower baseline PaO2 (≤ 50 mmHg).\cite{22}

**Diagnosis**

Diagnosis is usually based on imaging and arterial gas analysis. Arterial gases are analysed initially to identify elevated alveolar-arterial differences in O2 or hypoxemia in suspected HPS. There is controversy in the use of cut-off values of PaO2.\cite{23} A variety of cut-off points have been used resulting in a wide range of prevalence of HPS. Schenk and colleagues suggested that arterial hypoxemia defined as a PaO2 <70 mmHg or below the age-related threshold predicted the presence of HPS with high probability in the absence of intrinsic cardiopulmonary diseases.\cite{23} If PaO2 levels are abnormal a chest radiograph (CXR) and pulmonary function tests are used to help exclude other cardiopulmonary abnormalities. However, the demonstration of AV shunting despite the presence of other cardiopulmonary abnormalities by radiological means can achieve the diagnosis of HPS. In a cirrhotic patient suspected of HPS, it is important that other cardiopulmonary causes of hypoxia be excluded such as pulmonary atelectasis, ascites, chronic obstructive pulmonary disease, and hepatic hydrothorax.

A definitive diagnosis of HPS can be made by demonstration of pulmonary vasodilatation associated with functional arteriovenous shunting. Imaging studies that can identify such shunts include contrast echocardiography and perfusion scintigraphy with 99mTc, which are usually carried out following analysis of arterial gases to identify elevated alveolar-arterial differences in O2 or hypoxemia.

In the setting of HPS administration of 100% oxygen can be used as a noninvasive test to quantify pulmonary AVS. This is an important parameter to measure as demonstration of AVS suggests that the hypoxemia may not resolve following OLT. In response to breathing 100% oxygen if the PO2 levels rise to ≥600 mmHg, the presence of AVS is unlikely, but if levels fall to exceed 500 μmHg, the presence of a shunt cannot be ruled out. If PO2 levels fail to rise to above 150–200 μmHg, shunt is most probably the main mechanism of hypoxemia.\cite{24}
The presence of a ventilation–perfusion mismatch can be achieved by several means. The two methods available are 99Tc-MMA perfusion imaging or contrast echocardiography (CEE).\[24\] 99mTc-MMA perfusion scanning is a more sensitive procedure as it allows quantification of the degree of intrapulmonary shunting. 99mTc-MMA >20 µm in diameter are entrapped in pulmonary vasculature and undergo decay.\[25\] In patients with right to left shunt, the 99mTc-MMA enters the systemic circulation and distributed to systemic organs. Normally, less than 5% of the isotope can be quantitated over the brain. In HPS, there is >6% uptake in the brain.

Pulmonary angiography (PA) is an invasive procedure, which uses iodinated contrast media and is usually reserved in patients with an increase in the PaO2 to less than 300 mmHg. However, presently PA is rarely performed in the diagnosis of HPS but has a role when CTA suggests an AVM when PA may be used when the AVM appears amenable to embolization. PA is also able to differentiate between Type I and II HPS.\[29\] Type I HPS is depicted as normal to advanced diffuse ‘spider-like’ or spongy appearance.\[30\]

Most patients being cared for in hospital setting would have undergone a conventional CXR, which is not only useful in the diagnosis of HPS but is also crucial for excluding other causes of hypoxemia. A CXR in HPS shows bibasilar nodular or reticulonodular opacities in 5–13.8% of patients with CLD and 46–100% of patients with HPS.\[30\] These opacities are shown to represent dilated lung vessels on standard computed tomography [Figures 5–7]. High-resolution CT is useful in...
excluding pulmonary fibrosis or emphysema as the cause of these opacities. CT can identify decreased bronchoarterial ratio, which is highly specific for HPS. However, a multimodality approach is necessary to depict cases of liver origin. Suga and associates used breath-hold pulmonary perfusion SPECT-CT images to characterize CT manifestations of intrapulmonary
It is important that larger right to left shunts through pulmonary arteriovenous malformations are excluded by cross-sectional imaging when testing for HPS and when consideration is being given for hepatic radioembolization with Yttrium-90 glass microspheres. These coronal and sagittal reconstructions of contrast enhanced CT show a large arteriovenous malformation at the left lung base arteriovenous communications causing right-to-left shunt in two patients with HPS. In both patients, whole body scans depicted systemic organs thus confirming the existence of right-to-left shunt. Breath-hold SPECT-CT fusion images show that perfusion defects are predominantly located at subpleural reticulo-nodular opacities and/or dilated vessels in the lung base.

To summarise: the diagnostic flow chart of HPS generally starts with (1) arterial blood gas analysis performed in the supine position, (2) a CXR follows to rule out causes of hypoxemia other than HPS, (3) pulmonary function tests to include spirometry and DLCO. Spirometry is generally normal but may show a mild restrictive pattern due to pleural effusion or ascites. DLCO depicts decreased diffusion ability of lungs due to intrapulmonary vasodilatation. (4) 99mTc-MMA perfusion scanning (CEE if local expertise is available), (5) CT is reserved for patients when pulmonary AVM is suspected as the cause of AV shunt, but may be performed to look at the liver when the chest can be included into the protocol, (6) pulmonary angiogram is carried out as a prelude to vascular intervention when CT angiography has demonstrated a pulmonary AVM.

**Therapeutic measures**

No effective medical treatment is available, but better knowledge of the pathophysiology should improve this situation in the future. Medical management, transjugular intrahepatic portosystemic shunts (TIPS), and pulmonary arterial coil embolization have been described as temporizing measures until OLT is performed. Several therapeutic trials in HPS have shown poor results such as somatostatin analogues, cyclooxygenase inhibitors, and immunosuppressive agents such as corticosteroids and cyclophosphamide. Almitrine bimesylate a selective pulmonary vasoconstrictor has achieved anecdotal success. Inhaled prostanooid iloprost could improve quality of life in patients with HPS waiting for OLT and post-surgery until the resolution of the hypoxemia. A small study using garlic (*Allium sativum*) powder capsules daily for a minimum of 6 months resulted in a modest improvement in arterial blood gas profile. Suprahepatic inferior vena cava obstruction in association with HPS responds to percutaneous balloon cavaloplasty experiencing intrapulmonary shunt reversal. OLT provides the only definitive treatment of HPS. Over 85% of patients with HPS undergoing OLT show complete reversal of HPS or experience a significant improvement in hypoxemia. This reversal may take up to year. However, patients with HPS undergoing OLT have a significant mortality of 71% at 1 year.

Hepatic radioembolization with Yttrium-90 glass microspheres for treatment of inoperable primary liver cancer was first reported by Cao et al. It is presently considered a worthwhile therapeutic approach because of encouraging rate of response or stabilization. Initial work up includes assessment of the degree of AV shunt [Figures 1–4,8]. It is important that bigger intrapulmonary shunts are excluded by cross-sectional imaging. The same statement applies when dealing with HPS [Figure 9].

**Portopulmonary Hypertension**

**Pathophysiology**

PPH is defined by criteria obtained by right heart
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catheterization in patients with PoH and include: Elevated mean pulmonary artery pressure (>25 mmHg at rest, >30 mm Hg with exercise); increased pulmonary vascular resistance (>240 dynes s/cm²); and normal pulmonary artery occlusion pressure (<15 mmHg) or an elevated transpulmonary gradient. The risk of developing PH increases with the duration of PoH without any clear relation to the degree of PoH, hepatic failure, or amount of blood shunted. PH occurs in approximately 2% of patients with PoH due to either cirrhosis or extra hepatic lesions. Primary PH has been reported with extrahepatic PoH, portacaval shunt, and noncirrhotic portal fibrosis.

The cause of PPH is not well understood but may represent a combination of embolic and plexogenic factors. It is postulated that embolic material enters the lungs via portosystemic venous shunts associated with PoH [Figure 10]. Thromboembolic PH is a complication of congenital portosystemic venous shunt, and may be latently present in patients with PH of unknown etiology. Plexogenic PH may develop as a result of plexogenic arteriopathy due to vasoactive agents. Vasoactive substances bypass the liver via the portosystemic shunts or are ineffectively cleared by the compromised liver. Endothelial damage with vascular remodeling due to excessive pulmonary blood flow and smooth muscle proliferation adds further to vasoconstriction. Genetic variation in estrogen signalling and cell growth regulators is associated with the risk of PPH, which may explain the mechanism for its development in certain patients with severe liver disease. PoH either precedes or is diagnosed concurrently with PH, supporting the hypothesis that in PH, the pulmonary circulation may be exposed to vasoactive substances not metabolized or produced by the cirrhotic liver, which possibly induce vasoconstriction or direct toxic damage to the pulmonary arteries.

Clinical presentation
Early stage of PPH is generally asymptomatic or patients may have symptoms of the underlying liver disease. It is therefore imperative to have a high index of suspicion and screen for PH. With advancing disease exertional dyspnea is the most frequent presenting symptom of PPH (81%); other symptoms, such as syncope, chest pain, and fatigue, are seen in a third of the patients. Auscultation may reveal an accentuated pulmonary component of the second heart sound and a systolic murmur. Hemodynamic findings included severe pulmonary hypertension with normal pulmonary capillary wedge pressure and cardiac output.

Prognosis
Patients with PPH carry high perioperative mortality and have a mean survival of 15 months. As PPH increases perioperative death, screening for PPH is mandatory in patients undergoing evaluation for OLT.

Diagnosis
Lung function tests in PPH are not generally helpful although diffusing capacity may be reduced. B-type natriuretic peptide testing is helpful in monitoring the severity of disease and the efficacy of treatment in PH; but its role in PPH has not been defined. EKG findings in PPH include right atrial or ventricular enlargement and a right bundle branch pattern. A CXR may show prominent pulmonary arteries and cardiomegaly in advance PH. Standard echocardiography is an excellent tool to detect signs of PH. Findings suggestive of PPH include elevation of right ventricular systolic pressure (RVSP). Patients with an estimated RVSP greater than 50 mmHg undergo right heart catheterization.

Right heart catheterization (RHC) is regarded as a definitive test for confirmation of PPH, which allows accurate measurement of pulmonary artery pressures, pulmonary artery occlusion pressure, cardiac output, and pulmonary vascular resistance. RHC has the advantage of excluding other treatable causes of PH.

A CXR may be normal in the early stages. As the disease progresses, prominent pulmonary arteries and cardiomegaly are seen.
advances the central pulmonary arteries become prominent with a right ventricular enlargement and tapering of the peripheral vessels. At CT, PH can be diagnosed when the diameter of the main pulmonary artery is greater than or equal to 29 mm accompanied by a segmental artery-to-bronchus ratio greater than 1:1 in three of four pulmonary lobes or/and the ratio of the main pulmonary artery diameter to the aortic diameter is greater than 1, particularly in patients less than 50 years of age[52,53] [Figures 11–13]. HRCT may show a mosaic pattern of lung attenuation is a nonspecific finding common to all types for PH but is relatively specific for thromboembolic PH. Radionuclide lung perfusion scans may be normal or show diffuse patchy perfusion defects in severe PH.

**Therapeutic measures**

Optimal treatment for PPH remains elusive. Mild PPH with pulmonary artery pressure <35 mmHg is generally not treated. Moderate-to-severe PPH attempts are made to improve symptoms and lower pulmonary artery pressures and pulmonary vascular resistance. Anticoagulant therapy in PPH is generally not considered because of gastroesophageal varices and coagulation abnormalities related to liver disease, but anticoagulation should be considered in the absence of contraindications. Diuretics provide standard therapy in patients with PH and PPH, for the underlying liver disease, especially if ascites or peripheral edema is present. Oxygen therapy is used to treat hypoxemia. β-Blockers are used in some patients as prophylaxis of variceal bleeding. However, the use of β-Blockers in PPH is controversial as a significant worsening of exercise capacity has been reported after its use.[54]

A variety of vasodilators have been used in PPH. However, most of the studies of the use of vasodilator drugs have been used in PH from other causes. Prostanoids have been used successfully to lower pulmonary pressures in PPH. Epoprostenol has short half-life, a complicated delivery regime and is associated with several complications but has been shown to improve exercise capacity in PPH, although a survival advantage has not been documented to date.[55] Iloprost has similar effect and is used by inhalation but is very short-acting.[55] Bosentan is an endothelin receptor blocker, which is approved by FDA for the treatment of PH and PPH but is associated with mild hepatic dysfunction. However, irreversible hepatic toxicity is uncommon. Larger studies in the use of Bosentan in PPH are required. Ambrisentan has recently been approved for the treatment of PH. No studies are presently available for its use in PPH. Mild hepatic dysfunction may occur.[56] Sildenafil has been shown to increase cardiac output and decrease pulmonary artery pressures and pulmonary vascular resistance without serious adverse events in PH. Recent case series of 14 patients shows that sildenafil might be effective in PPH.[57]

OLT is not the treatment of choice for PPH, but after optimal hemodynamic and clinical improvement with medical therapy, OLT can be considered an option in selected patients.[58] PPH increases the risk of intraoperative and immediate postoperative complications of OLT. OLT should not be undertaken in patients with mean pulmonary artery pressure > 50 mm Hg whilst patients with mean pulmonary artery pressure between 35 and 50 mm Hg also have an increased mortality rate but may benefit from prolonged treatment for PH. Sildenafil use to treat PPH prior to OLT has met with some success.

PPH is an uncommon complication of ESLD in children. Epoprostenol can bridge PPH patients to OLT. OLT leads to

![Figure 11: A CXR of a 41-year-old man with cirrhosis and PPH show enlarged hila and peripheral pruning of the vessels secondary to PH. An axial contrast enhanced CT through the pulmonary arteries of another patient with PPH showing aneurismal dilatation of the main pulmonary artery and the right pulmonary artery with peripheral pruning due to PH](image1)

![Figure 12: A CXR on a patient with PPH secondary to cirrhosis shows a dilated pulmonary conus and aneurismal dilatation of the right pulmonary artery. The image on the left is coronal reconstruction of contrast enhanced CT on the same patient, which depicts a probable re-canalized thrombus underlying the aneurismal dilatation of the right pulmonary artery](image2)

![Figure 13: Pulmonary angiography on a patient with PPH showing dilatation of the pulmonary arteries and pruning of the peripheral vessels](image3)
rapid resolution of HPS and PPH and currently represents the only successful treatment for these children.  

Most successful OLT in PPH has been reported from unrelated cadaver donors. One successful case of living-related liver transplantation in a patient with PPH has been published.

Conclusions

The major pulmonary vascular complications of CLD include HPS and PPH. The pathogenesis, imaging, treatment, and prognostic implications of these complications are discussed. In the context of ongoing uncertainty about causation and treatment of the HPS, future studies must focus on better understanding the pathogenesis of the HPS, predicting reversibility after OLT, and identifying other treatment options. It is emphasized that appropriate knowledge of the pathogenesis of vascular complications and imaging of CLD would assist the clinician in initiating the appropriate treatment at the most appropriate time.

References

1. Krowka MJ, Cortese DA. Hepatopulmonary syndrome: An evolving perspective in the era of liver transplantation. Hepatology 1990;11:138-42
2. Kennedy TC, Knudson RJ. Exercise-aggravated hypoxemia and orthodeoxia in cirrhosis. Chest 1997;72:305-9
3. Krowka MJ, Cortese DA. Hepatopulmonary syndrome: Current concepts in diagnostic and therapeutic considerations. Chest 1994;105:1528-37.
4. Abrams G, Jaffe CC, Hoffer PB, Binder HJ, Fallon MB. Diagnostic utility of contrast echocardiography and lung perfusion scan in patients with hepatopulmonary syndrome. Gastroenterology 1995;109:1283.8.
5. Castro M, Krowka MJ. Hepatopulmonary syndrome: A pulmonary vascular complication of liver disease. Clin Chest Med 1996;17:35-48.
6. de Macêdo LG, Lopes EP. Hepatopulmonary syndrome: An update. Sao Paulo Med J 2009;127:223-30
7. Bruix J, Bosch J, Kravetz D, Mastai R, Rodés J. Effects of prostaglandin inhibition on systemic and hepatic hemodynamics in patients with cirrhosis of the liver. Gastroenterology 1985;88:430-5
8. Cremona G, Higenbottam TW, Mayoral V, Alexander G, Demonchneaux E, Borland C, et al. Elevated exhaled nitric oxide in patients with hepatopulmonary syndrome. Eur Respir J 1995;8:1883-5
9. Davis HH 2nd, Schwartz DJ, Lefrak SS, Susman N, Schainker BA. Alveolar-capillary oxygen disequilibrium in hepatic cirrhosis. Chest 1978;73:507-11
10. Krowka M, Wiseman G, Burnett O. Hepatopulmonary syndrome: A prospective study of relationships between severity of liver disease, PaO2 response to 100% oxygen, and brain uptake after TC 99mMMAA lung scanning. 2000. p. 615-624
11. Hedenstierna G, Soderman C, Eriksson LS, Wahren J. Ventilation perfusion inequality in patients with non-alcoholic liver cirrhosis. Eur Respir J 1991;4:711-7.
12. Hopkins WE, Waggoner AD, Barzilai B. Frequency and significance of intrapulmonary right-to-left shunting in end-stage hepatic disease. Am J Cardiol 1992;70:516-9.
13. Whyte MK, Hughes JM, Peters AM, Ussov W, Patel S, Burroughs AK. Analysis of intrapulmonary right to left shunt in the hepatopulmonary syndrome. J Hepatol 1998;29:85-93.
14. Brussino L, Bucca C, Morello M, Scappaticci E, Mauro M, Rolla G. Effect on dyspnea and hypoxemia of inhaled NG-nitro-L-arginine methyl ester inhepatopulmonary syndrome. Lancet 2003;362:43-4.
15. Zhang HY, Han DW, Wang XG, Zhao YC, Zhou X, Zhao HZ. Experimental study on the role of endotoxin in the development of hepatopulmonary syndrome. World J Gastroenterol 2005;11:567-72.
16. Rabiller A, Nunes H, Lebec D, Tazi KA, Wartski M, Dulmet E, et al. Prevention of gram-negative translocation reduces the severity of hepatopulmonary syndrome. Am J Respir Crit Care Med 2002;166:514-7.
17. Gómez FP, Barberà JA, Roca J, Burgos F, Gistau C, Rodríguez-Roisin R. Effects of nebulized N(G)-nitro-L-arginine methyl ester in patients with hepatopulmonary syndrome. Hepatology 2006;43:1084-91.
18. Seward JB, Hayes DL, Smith HC, Williams DE, Rosenow EC 3rd, Reeder GS, et al. Platypnea-orthodeoxia: Clinical profile, diagnostic work-up, management and report of 7 cases. Mayo Clin Proc 1984;59:221.
19. Lima BL, Franca AV, Pazin-Filho A, Araújo WM, Martinez JA, Maciel BC, et al. Frequency, clinical characteristics, and respiratory parameters of hepatopulmonary syndrome. Mayo Clin Proc 2004;79:42-8
20. Rodríguez-Roisin R, Roca J, Agusti AG, Mastai R, Wagner PD, Bosch J. Gas exchange and pulmonary vascular reactivity in patients with liver cirrhosis. Am Rev Respir Dis 1987;135:1085-92
21. Fallon MB, Krowka MJ, Brown RS, Trotter JF, Zacks S, Roberts KE, et al. Impact of hepatopulmonary syndrome on quality of life and survival in liver transplant candidates. Gastroenterology 2008;135:1168-75.
22. Swanson KL, Weiner RH, Krowka MJ. Natural history of hepatopulmonary syndrome: Impact of liver transplantation. Hepatology 2005;41:1122-9.
23. Schenf P, Fuhrmann V, Madl C, Funk G, Lehr S, Kandel O, et al. Hepatopulmonary syndrome: Prevalence and predictive value of various cut offs for arterial oxygenation and their clinical consequences. Gut 2002;51:853-9.
24. McAdams HP, Erasmus J, Crockett R, Mitchell J, Godwin JD, McDermott VG. The hepatopulmonary syndrome: Radiologic findings in 10 patients. AJR Am J Roentgenol 1996;166:1379-85.
25. El-Shabrawl MH, Omran S, Wageeh S, Isa M, Okasha S, Mohnen NA, et al. 99mTc-pertechnetium-macroaggregated albumin perfusion lung scan versus contrast enhanced echocardiography in the diagnosis of the hepatopulmonary syndrome in children with chronic liver disease. Eur J Gastroenterol Hepatol 2010;22:1006-12.
26. Shub C, Tajik AJ, Seward JB, Dines DE. Detecting intrapulmonary right-to-left shunt with contrast echocardiography: Observations in a patient with diffuse pulmonary arteriovenous fistulas. Mayo Clin Proc 1976;51:814-6.
27. Karabulut A, Ilhumur K, Yalcin K, Toprak N. Hepatopulmonary syndrome and right ventricular diastolic functions: An echocardiographic examination. Echocardiography 2006;23:271-8.
28. Vedrine JM, Duperret S, Bizollon T, Magnin C, Motin J, Trepo C, et al. Comparison of transoesophageal and transthoracic contrast echocardiography for detection of an intrapulmonary shunt in liver disease. Chest 1997;111:1236-40.
29. Scott VL, Dodson SF, Kang Y. The hepatopulmonary syndrome. Surg Clin North Am 1999;79:23-41.
30. Castro M, Krowka MJ. Hepatopulmonary syndrome: A pulmonary vascular complication of liver disease. Clin Chest Med 1996;17:35-48.
47. Hervé P, Lebrec D, Brenot F, Simonneau G, Humbert M, Sitbon O, et al. Pulmonary arterial hypertension in portal hypertension. Eur Respir J 1998;11:430-2.

48. Fallon MB, Krowka MJ, Brown RS, Trotter JF, Peter I, Tighiourt H, et al. Genetic risk factors for portopulmonary hypertension in patients with advanced liver disease. Am J Respir Crit Care Med 2009;179:835-42.

49. Robalino BD, Moodie DS. Association between primary pulmonary hypertension and portal hypertension: Analysis of its pathophysiology and clinical, laboratory and hemodynamic manifestations. J Am Coll Cardiol 1991;17:492-8.

50. Mandell MS, Groves BM. Pulmonary hypertension in chronic liver disease. Clin Chest Med 1996;17:37-33.

51. The International Primary Pulmonary Hypertension Study (IPPHS). Chest 1994;105:375-415.

52. Tan RT, Kuzo R, Goodman LR, Siegel R, Haasler GB, Presberg KW. Utility of CT scan evaluation for predicting pulmonary hypertension in patients with parenchymal lung disease. Chest 1998;113:1290-6.

53. Ng CS, Wells AU, Padley SP. A CT sign of chronic pulmonary arterial hypertension: The ratio of main pulmonary artery to aortic diameter. J Thorac Imaging 1999;14:270-8.

54. Provencher S, Herve P, Jais X, Lebrec D, Humbert M, Simonneau G, et al. Deleterious effects of beta-blockers on exercise capacity and hemodynamics in patients with portopulmonary hypertension. Gastroenterology 2006;130:120-6.

55. Kuo PC, Johnson LB, Plotkin JS, Howell CD, Bartlett ST, Rubin LJ. Continuous intravenous infusion of epoprostenol for the treatment of portopulmonary hypertension. Transplantation 1997;63:604-6.

56. Anderson JR, Nawarskas JJ. Pharmacotherapeutic management of pulmonary arterial hypertension. Cardiol Rev 2010;18:148-62.

57. Reichengerger F, Voswinckel R, Steveling E, Enke B, Kreckel A, Olschewski H, et al. Sildenafil treatment for portopulmonary hypertension. Eur Respir J 2006;28:563-7.

58. Porres-Aguilar M, Zuckerman MJ, Figueroa-Casas JB, Krowka MJ. Portopulmonary hypertension: State of the art. Ann Hepatol 2008;7:321-30.

59. Iqbal CW, Krowka MJ, Pham TH, Freece DK, El Youssef M, Ishitani MB. Liver transplantation for pulmonary vascular complications of pediatric end-stage liver disease. J PediatrSurg 2008;43:1813-20.

60. Swanson KL, Krowka MJ. Screen for portopulmonary hypertension, especially in liver transplant candidates. Cleveland Clin J Med 2008;75:121-36.

61. Sulica R, Emre S, Poon M. Medical management of portopulmonary hypertension and right heart failure prior to living-related liver transplantation. Congest Heart Fail 2004;10:192-4.

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