Pathobiology of Pulmonary Arterial Hypertension and Right Ventricular Failure

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Abstract

Pulmonary arterial hypertension (PAH) is no longer an orphan disease. Three different classes of drugs for the treatment of PAH are being used and an increasing number of patients are being treated with a single drug or a combination therapy. During the last twenty-five years, new insights into the pathobiology of PAH have been gained. The classical mechanical concepts of pressure, flow, shear stress, RV wall stress and impedance, have been complemented with the new concepts of cell injury and repair and interactions of complex multicellular systems. Integrating these concepts will become critical, as we design new medical therapies in order to change the prognosis of the patients with this fatal group of diseases. This review intends to summarize recent pathobiological concepts of PAH and RVF derived mainly from human studies, which reflect the progress made in the understanding of these complex group of pulmonary vascular diseases.

Keywords: BMPR2, genetics, inflammation, growth factors
Pulmonary hypertension (PH), like systemic hypertension, is not a single disease but a group of diseases, which share the defining element of a mean pulmonary arterial pressure ≥25 mmHg. PH has been classified and divided into 5 groups(1). The World Health Organization group 1, or pulmonary arterial hypertension (PAH), includes diseases characterized by elevated pulmonary arterial pressure and elevated blood flow resistance due to a precapillary pulmonary microangiopathy. Whereas a quarter of a century earlier idiopathic PAH (IPAH) was a disease of young women, data from the Registry to Evaluate Early and Long Term PAH Disease Management (REVEAL) now indicate that – at least in the United States – the phenotype of PAH has significantly changed. Today, up to 79% of patients are women close to menopausal age, with higher predominance of females (3.6:1 female:male ratio) and a large percentage of obesity(2). However, there is an apparent discrepancy in the PAH phenotype between US and European registries(3-5), suggesting perhaps the presence of currently unidentified environmental factors that could contribute to the development and progression of the disease.

PAH is no longer an orphan disease(6). In the last 15 years, three classes of drugs have been developed and an increasing number of patients are being treated with a single drug or a combination therapy (7,8). The advent of PAH-specific therapies has improved the survival of PAH patients in comparison to historic registries. However, in spite of treatment, the mortality remains unacceptably high, as the survival for incident patients continues to be as low as 54.9% at 3 years after diagnosis (95% CI, 41.8 to 68.0)(9).

Although the structural changes that explain the “fixed pulmonary hypertension” of PAH patients have been described in recent decades (10-12) and major changes to the “classic” pathobiology of PAH have been described(13,14), current PAH-specific drugs are predominantly vasodilators (although some antiproliferative properties have been reported in cultured normal lung cells)(15). Furthermore, the mechanisms underlying right ventricular failure (RVF) are incompletely understood(16,17). This review intends to put in perspective pathobiological concepts of PAH and RVF derived mainly from human studies, which reflect the progress made in the understanding of this complex group of pulmonary vascular diseases.

Genetic influences and epigenetics in PAH
The pathogenesis of PAH is based on one or more “hits”, and genetic factors conferring a predisposition for the development or progression of severe PAH may be required (Figure 1)(18-20). Mutations of the Bone Morphogenic Type II Receptor (BMPR2) gene, a member of the transforming growth factor (TGF-β) superfamily, were first described in patients with familial/hereditary PAH (HPAH)(18). Although BMPR2 mutations in HPAH have a high prevalence (70%), limited numbers of patients with IPAH have mutations(19), and only 20% of the carriers ever develop PAH during their lifetime(19). BMPR2 mutation carriers at the time of PAH-diagnosis are younger (20,21) and less likely have a vasoreactive component(22). A large number of mutations have been reported and the majority translate into a loss of function (23) or reduced BMPR2 expression(24). It has been reported that the disease appears to be more severe when patients carry a truncating BMPR2 mutation(25) however, this appears not to be the case in the French patients (21). Mechanistic studies indicate that BMPR2 mutations are permissive but not necessary for the development of severe PAH. BMPR2 may be one of the guardians of lung vessel homeostasis, as gene knockout and silencing experiments have clearly demonstrated that both apoptosis and cell proliferation of pulmonary vascular smooth muscle and endothelial cells are controlled by BMPR2(26). Intact BMPR2 signaling may be necessary for the execution of a normal lung vascular wound-healing program, preventing apoptosis-induced compensatory cell proliferation. BMPR2 loss makes cells more susceptible to apoptosis(27), however, vascular cell-apoptosis alone, is insufficient for angioproliferation to occur. BMPR2 signaling appears to define the cellular identity during reparative responses(28). BMP signaling is regulated at many different levels and each level could potentially contribute to abnormal BMPR2 function, without necessarily involving a mutation in the BMPR2 gene(26).

For example, mutations in the type I TGF-β receptor, ALK-1(29,29-31), have been observed in patients with severe PAH occurring in families with hereditary hemorrhagic telangiectasia. Moreover, although infrequent, mutations in the BMPR2-downstream mediators, the smad proteins, have also been described in PAH patients(32). In the aggregate the data indicate that BMP/TGF-β signalling plays an important role in the maintenance of the normal lung arteriolar structure.

Epigenetic mechanisms influence gene expression via modifications of the chromatin, histones and regulatory micro RNAs(33). At present, there is no firm evidence that PAH has an epigenetic component(34). Downregulation of BMPR2 expression has been explained by activation of a
STAT3/miRNA-17-92 microRNA axis in normal human lung endothelial cells after interleukin-6 exposure(35). Interestingly, mice overexpressing IL-6 develop severe pulmonary hypertension(36) and, unlike other PH mice models, also develop angioobliterative vascular remodelling and robust RV hypertrophy(37). Overexpression of miR-17 also increases proliferation of human pulmonary artery smooth muscle cells and inhibition with an specific miR-17 antagomir ameliorated PH in two experimental models(38). Another microRNA, miRNA-204, has been found downregulated in pulmonary artery smooth muscle cells isolated from patients with PAH(39). Upregulation of miR204 seems to induce an apoptosis-resistant phenotype in smooth muscle cells.

**The paradigm shift: From vasoconstriction to cell growth.**

Terms like “cell phenotype switch,” “apoptosis-resistance” and “angiogenic niche” have introduced a new vocabulary which can be used to explain the pathobiology of PAH(40). The early pioneers of PH research focused their attention on hypoxic vasoconstriction(41), and because the arachidonic acid-derived prostacyclin is a powerful inhibitor of hypoxic pulmonary vasoconstriction(42), it was thus intuitive to establish prostacyclin infusion as the first treatment for severe PAH(43). Since then, rapid progress has been made. Our knowledge of the cellular and molecular components involved in the pathobiology of PAH has expanded and these new insights are likely to change our future approach to the treatment of PAH. We now regard the plexiform pulmonary vascular lesions as a product of an angioproliferative process leading to vascular occlusion (angioobliteration)(13,40) and we appreciate that these lesions are composed of a multitude of cell types, among them, actively dividing and phenotypically abnormal apoptosis-resistant endothelial cells with a monoclonal origin(44), progenitor cells(45) and immune cells(46-48). Immunohistochemistry, using antibodies directed against cell surface epitopes and nuclear markers of cell activation, identifies different cell phenotypes with their expressed genes and proteins(40). Hallmarks of cancer in angioproliferative PAH have been described, in particular angiogenesis(13), metabolic changes(49) and apoptosis-resistant cells(50). Based also on the therapy refractoriness of PAH, a hypothesis of “quasi-malignancy” has been proposed(40): apoptosis may be the initiating event followed by the selection of proliferating apoptosis-resistant cells –some of which may be stem cells (Figure 1). PAH plexiform lesions also show decreased expression of the tumor-suppressor protein
peroxisome proliferator activated receptor (PPAR) gamma(51), while exhibiting increased expression of β-catenin, CXCR4 (a receptor that mediates metastatic cell homing), and survivin, among other(40). Survivin, an inhibitor of apoptosis, is expressed in essentially all cancers, and cultured cells isolated from the lungs of PAH patients exhibit a markedly increased expression of survivin. Moreover, experimental gene therapy targeting this protein suppresses proliferation and ameliorates PAH in rats(52). Another hallmark of cancer is the metabolic switch towards glycolysis (Warburg effect, perhaps as a defense against oxidative damage). Pulmonary arterial endothelial cells from IPAH patients cultured in vitro, demonstrate reduced oxygen consumption, reduced mitochondrial respiration and increased glycolytic metabolism(49).

The “quasi-malignancy“ disease model also accommodates the concept of a cancer stem cell/angiogenic niche(40). Precursor cells can be transported into the pulmonary vascular adventitia via vasa vasorum(53), stem cells can divide and migrate from a niche at the media-adventitia border zone(54) and pulmonary microvascular endothelial cell stem cells of small arteries(55) may proliferate to the point of lumen obliteration. It is presently unclear whether stem- and precursor cells are friends or foes. For example, autologous endothelial progenitor cells transplantation therapy for PAH has reported positive results(56). However, this early trial has been criticized(57) because the cells utilized for the clinical trial were not “true” progenitors. Conversely, anti c-Kit (a stem cell marker) treatment for PAH with the tyrosine kinase inhibitor imatinib mesylate, has demonstrated anecdotal positive results(58). Stem cell-based gene therapy is currently being investigated in a clinical trial (NCT00469027).

Growth factors and receptor tyrosine kinases

Many potent cell growth factors are overexpressed in IPAH vascular lesions, and it is likely that these growth factors and their receptor tyrosine kinases contribute to cell growth and vessel obliteration. The Platelet Derived Growth Factor (PDGF) has been extensively studied in PAH(59), and it has been shown that PDGF can induce the proliferation and migration of SMCs and fibroblasts. PDGF has been proposed as a key mediator in the progression of several fibroproliferative disorders. It was postulated that PDGF receptor blockers could reverse PAH, based on the case report of a patient with end-stage IPAH, treated with imatinib mesylate. Subsequently, a clinical safety trial evaluating imatinib mesylate as a treatment of
advanced forms of PAH was performed(60). The data showed a significant decrease in pulmonary vascular resistance (PVR) and a moderate increase in cardiac output, but no significant change in pulmonary artery pressures, leading to the conclusion that imatinib mesylate may have a limited role in the treatment of IPAH patients. Other tyrosine kinase inhibitors for PAH are being studied(61,62). It is important that the tyrosine kinase inhibitors imatinib, sorafenib and nilotinib were shown in animal studies to be acute pulmonary vasodilators(63). However, the vasodilator effect in clinical studies of PAH was not prominent. This is an example, that identifying highly specific molecular targets in PAH animal models that do not reproduce the salient features of human PAH, may be misleading and the drug effect may be lost in translation(37,64). Paradoxically, dasatinib, a multi-tyrosine kinase inhibitor, has been associated with the development of PAH(65,66).

Inflammation and Immune Response

The association of autoimmune disorders with PAH and the presence of antinuclear and antiphospholipid antibodies in the serum of patients with IPAH has been appreciated for many years(67). There is more evidence that lymphoid neogenesis occurs in IPAH; macrophages, mast cells, T- and B- lymphocytes, plasma cells and anti-endothelial cell antibodies are present in and around the complex pulmonary vascular lesions in IPAH patients(13,48). Serum levels of IL-1 and IL-6 are high in IPAH patients(68), and serum IL-6 levels negatively predict patient survival(69). However, whether inflammation and aberrant immune responses in IPAH are cause or consequence remains unknown. Likely PH occurs when an inflammatory pulmonary arteriolar injury is not resolved by (normally) protective, innate anti-inflammatory mechanisms. Regulatory T-cells (Tregs) control not only other T-cells but also regulate monocytes, macrophages, dendritic cells, natural killer cells and B-cells(70), and recent evidence suggests that decreased Treg cell number or function may favour the development of PH(71). For example, conditions associated with PAH, such as HIV, systemic sclerosis, systemic lupus erythematosus, Hashimoto’s thyroiditis, Sjogren’s Syndrome and the antiphospholipid syndrome are characterized by abnormal CD4\(^+\) T-cell number and function(72-77). At the experimental level, athymic rats, lacking T-cells, develop pronounced PH after vascular injury with a vascular endothelial growth factor receptor blocker. The lungs in these animals are populated by infiltrating macrophages, mast cells and B cells, similar to human PAH.
lesions(71,78). Most importantly, PH is prevented by immune reconstitution of Tregs prior to the induction of vascular injury(71). All together, it is a possibility that aberrant Treg-cell function in the face of vascular injury can result in heightened innate and adaptive immune responses that could initiate and/or propagate the development of PH (79)(Figure 2). Despite the increasing evidence suggesting a role for immune deregulation in PAH, there is only limited anecdotal evidence suggesting that PH associated with connective tissue disorders –such as systemic lupus erythematosis and systemic sclerosis or viral infections, such as human herpes virus-8 – can respond to glucocorticoids or targeted B cell depletion(80,81). Sanchez et al(82) and Jais et al(83) reported on treatment of patients with mixed connective tissue disease - or systemic lupus erythematosus – associated PH, where corticosteroid and cyclophosphamide has been used as first-line drugs. The effectiveness of B cell depletion is currently being put to test with an NIH-trial examining the effectiveness of rituximab – a chimeric monoclonal antibody against the protein CD20, primarily expressed by B-cells – for systemic sclerosis-associated PH (NCT01086540).

It is becoming apparent that circulating factors can likely amplify lung vascular injury, attract immune cells and/or repair cells which respond to a variety of chemotactic stimuli, suggesting perhaps, a systemic disease component contributing to the development or progression of PAH. In the “modern era” of PAH treatments, where standard vasodilation therapies have failed to reverse or stop the progression of PAH, novel targets such as discrete immune pathways hold promise. However, new drugs and clinical trials will require to assess which patients may respond to anti-inflammatory treatment strategies.

The Right Ventricle-Lung Circulation Axis

Patients with severe PAH die from right ventricular failure (RVF). The interactions between the remodelled lung circulation and the right ventricle (RV) are only partially understood(16). RVF is commonly seen as a consequence of chronic RV pressure overload and a developing imbalance between RV oxygen demand and supply, sometimes culminating into RV ischemia. Indeed, increased pulmonary arterial pressure increases RV wall tension and oxygen consumption, and at the same time is associated with compromised coronary blood flow. However, and remarkably, patients with the Eisenmenger physiology generate a degree of RV afterload similar to that of patients with IPAH, but have a much better
survival and only develop overt RVF until a late stage of their disease (84). We also do not fully understand why some patients, at the time of PAH diagnosis, present with signs and symptoms of RVF while other patients with the same degree of RV afterload present in NYHA functional class I with no signs of RVF. Based on mechanistic animal studies, it has been proposed that, in addition to pressure overload, other factors, perhaps products of the “sick lung” circulation, contribute to RV dysfunction (85,86). Similar to left heart failure, it has been postulated that RVF is characterized by an abnormal energy metabolism (87). Positron emission tomography studies have demonstrated increased accumulation of the radiopharmaceutical glucose analog $^{18}$F-2-Deoxy-2-Fluoro-D-Glucose (18-FDG) in the RV of PAH patients, in comparison to normal subjects (88,89). Although increased, 18-FDG uptake does not seem to correlate with disease severity. A recent study showed no difference in 18-FDG uptake between NYHA class II and class III PAH patients, even when there was a clear-cut difference in cardiac output (90). Moreover, this study found no correlation between RV 18-FDG uptake and RV mechanical efficiency. It is important to remember that 18-FDG uptake studies have limitations and that the physiological interpretation is not straightforward. Tracer uptake can be influenced by multiple factors (91). Moreover, 18- FDG avidity represents the total amount of 18-FDG present in the tissue at the time of image acquisition (92), which is not necessarily equal to the total amount of 18-FDG absorption by the cell. Instead, the scan also captures the 18- FDG present in the blood and intercellular spaces. Perhaps, the most prominent limitation of 18-FDG studies interpretation is explained by its chemical structure. 18-FDG lacks the functional hydroxyl group that is required for the enzyme phosphoglucose isomerase (the second enzymatic step of glycolysis) to function properly (93). Briefly, 18-FDG cannot be further metabolized inside the cells. Thus, 18-FDG uptake studies are too crude to identify a “metabolic switch” in the failing RV. Whereas, experimental studies have demonstrated that RVF exhibits increased expression of glycolysis-related genes (94) and increased enzymatic glycolysis rates (95), a complete characterization of RV “metabolic remodeling” in human PAH is still lacking.

Noninvasive imaging technologies, in particular echocardiography and cardiac magnetic resonance imaging (CMRI), are now providing a more complete assessment of the structural and functional changes of the failing and non-failing RV in the setting of chronic PAH. Whereas changes in longitudinal shortening (tricuspid annulus plane systolic excursion, TAPSE) have been the standard measure to evaluate RV
dysfunction (96), other changes in RV morphology have been shown to correlate with survival. For example, a large RV volume and impaired left ventricular filling, assessed by CMRI, are strong independent predictors of mortality and treatment failure (97). It has also been reported that decreased RV-longitudinal shortening and free-wall motion at the time of diagnosis do not significantly change over time, whereas RV-transverse shortening continues to decline over time in non-survivor PAH patients (98). RV-transverse shortening could be particularly valuable for the monitoring of treatment success or failure, as we move forward with clinical trials for RV-targeted therapies. Measurement of RV free wall longitudinal deformation (strain) and the rate of deformation by speckle-tracking strain echocardiography, has been shown to identify patients with greater severity of RV dysfunction(99). Most importantly, changes in RV strain correlate with significantly with functional class, and can predict mortality. Lastly, the prognostic importance of RVF in patients with PAH has recently been revisited. Van de Veerdonk et al have reported that although elevated PVR (in treated PAH patients) is associated with higher mortality, the RV ejection fraction (measured by CMRI) is the strongest predictor of survival, even after a response to vasodilator therapy has been established(100). Other noninvasive imaging tools to evaluate RV dysfunction in PAH patients have been described. For a more detailed review on noninvasive imaging tools in PAH, please refer to (101-104).

**Challenges of PAH treatment**

The future of the medical management of PAH faces multiple obstacles. 1) In spite of improved screening tools, PAH is not discovered during its early stage and treatment is delayed. Data from the REVEAL registry indicate that the lag-time from the onset of symptoms to right heart catheterization has not changed when compared to the data published based on the first NIH-registry(105). 2) Although improvements in survival have been demonstrated in clinical trials, the overall patient survival in the “modern era” remains low and the survival of “incident” PAH patients appears to be similar to that of the idiopathic pulmonary hypertension patients enrolled in the first NIH registry (8). Indeed, the concept of vasodilator drug treatment for PAH has become controversial in the last decade (for reasons discussed in some detail above). In one meta-analysis, Macchia et al called PAH “a clinical condition looking for new drugs and research methodology” (106) and stated “whether current therapies of PAH improve long-term
survival has never been demonstrated in a randomized clinical trial (RCT)”(107). However, Macchia’s meta-analysis was criticized due to the fact that many of the studies assessed were underpowered to determine mortality(108). A later meta-analysis, by Galiè et al (which included 23 RCTs performed exclusively in PAH patients) reported that a reduction of 43% in overall-mortality was observed in the groups randomized to active treatments, when compared with those randomized to placebo. However, the average duration of the trials assessed was limited (14.3 weeks) and, with the exception of the first epoprostenol trial(43), none of the trials defined survival as a primary end-point. Finally, 3) Future therapeutic strategies face another critical paradox: While the remodelled lung circulation in PAH is characterized by angiogenesis, apoptosis-resistance and cell proliferation, the failing RV suffers from ischemia(109), capillary rarefaction and cardiomyocyte apoptosis(86), conditions that could be worsened by pharmacotherapy targeting pulmonary vascular remodelling. One example could be the tyrosine kinase inhibitors, which are known to be cardiotoxic(110).

Conclusions

PAH is increasingly recognized not as one disease but as a group of diseases where genetic susceptibility renders patients vulnerable to a chronic pulmonary microangiopathy. It appears that increased afterload of the RV is insufficient to explain right heart failure, as it is best illustrated by the better prognosis and longer survival of patients with Eisenmenger physiology. Further explorations of the molecular mechanisms of cell reprogramming in PAH, in both the lung vessels and the heart, should lead to a better understanding of this group of diseases. The classical pathophysiology of PAH which operates with the mechanical concepts of pressure, flow, shear stress, RV wall stress and impedance, should be complemented with the new pathobiological concepts of cell injury and repair and interactions of complex multicellular systems (Figure 4). Precursor-and stem cells derived from the bone marrow and resident lung vascular stem cells should also be considered as active players in the process of pulmonary vascular wound healing “gone awry”.

Many questions remain: Is there an early vasoreactive disease phase in PAH? Can the inflammatory disease component – discussed above – be treated in most patients? Do patients with reactive PAH (treatable with calcium-channel blockers) and PAH patients with NYHA class I, that do not progress after
many years, represent pathogenetically-unique subgroups? The former a vasospastic variant, the latter a variant characterized by a robust, afterload – and stress – resilient right ventricle? Can prevention of RV dysfunction prolong life?

As new treatment strategies appear on the horizon, clinical trialists should consider investigating the mechanisms of treatment-success or treatment-failure by comparing treatment responders and non-responders as is currently the novel approach developed for cancer treatment trials(111,112). PAH should be viewed as a cardiopulmonary system disease, which is based on the paradox of apoptosis, apoptosis-resistant growth of phenotypically abnormal cells, and angiogenesis in the lungs, and apoptosis and capillary loss in the heart. Thus, new therapeutic strategies should consider that drugs originally designed to tackle angiogenesis in PAH lungs could potentially have a profound, negative impact on the failing right ventricle. Integrating new pathobiological concepts will become critical, as we design new medical therapies in order to change the prognosis of the patients with this fatal group of diseases.

**Authorship**

All authors contributed to the conception, design and drafting of the review. All authors approved the final version of this manuscript.

**References**

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Figure legends

Figure 1: Hallmarks of severe angioproliferative pulmonary arterial hypertension.

Genetic determinants, epigenetic factors and other conditions synergize and result in the injury and
apoptosis of pulmonary arteriolar endothelial cells. Endothelial cell apoptosis triggers compensatory proliferation of phenotypically abnormal endothelial cells. Apoptosis may also be the initiating event activating the bone marrow to release precursor and stem cells and eliciting a local immune response of the vascular wall. The end result is the complex vascular lesion, the plexiform lesion. These complex lesions are obliterating the lumen of small arteries, predominantly localized to vessel bifurcations. The mature lesions are composed of endothelial cells (magenta), smooth muscle cells (yellow), apoptotic-cells (purple) and phenotypically altered apoptosis-resistant endothelial cells (red). These cells, together inflammatory cells, populate a microenvironment of exuberant cell growth.
Figure 1
Figure 2: Participation of the immune system in pulmonary vascular remodeling.

The initial arteriolar endothelial cell injury can be due to mechanical stress, toxins or antibodies. The ‘two-hit’ hypothesis considers that defective immune regulation as in HIV/AIDS and scleroderma, participates in the lumen-obliterating cell growth (1) and in the impaired injury resolution (2). The disease model depicted here reflects the complexity of multicellular interactions within and around the vascular wall and the active role of the bone marrow and of inflammatory and immune cells in the pathobiology of PAH. The first of several immune cells, the multi-growth factor-producing mast cell, to be identified in the perivascular space of pulmonary arterioles in lungs from patients with severe pulmonary hypertension, may populate these vessels because of hyperplasia (in situ expansion of mast cell number), or after being recruited from the bone marrow. An impaired activity of regulatory T cells (Treg cells) may facilitate the angioobliteration.
Figure 2

'Two-hit' Disease Initiation
Pulmonary Endothelial Injury + Defect in Immune Regulation

Impaired Injury Resolution

Precursor Stem Cell in Adventitial Niche

Hematopoietic Stem Cell (HSC)

Endothelial Myoepithelioma

Vasa vasorum

Mast Cell

Megakaryocyte

Bone Marrow
Figure 3: The “sick lung circulation”– right heart failure axis.

Increased pulmonary vascular resistance generates a significant afterload for the right ventricle resulting in right ventricular hypertrophy. The structural and functional changes during the development of right ventricular failure can be characterized clinically (2A-F). Right ventricular dysfunction is likely associated with multiple cellular changes (3) such as oxidative stress, apoptosis, inflammation, fibrosis and metabolic remodeling; these factors also contribute to RV dysfunction and failure. The cellular changes are either the consequence of chronic RV pressure overload or the effect of circulating factors released from the “sick lung circulation”.
Figure 3

1. Pulmonary Vascular Remodeling
   - A. Increased Vascular Resistance
   - B. Increased Pulmonary Arterial Pressure/RV afterload

2. Cardiac functional and structural changes
   - A. Increased wall stress
   - B. Increased RV preload
   - C. Right atrium dilatation
   - D. Abnormal RV Contraction
   - E. Paradoxical Septal Movement
   - F. Restrictive LV filling

3. Cellular changes in the RV

Pressure Overload
   - RVF
   - Fibrosis
   - Oxidative Stress
   - Apoptosis
   - Inflammation
   - Apoptosis
   - Mitochondrial dysfunction
   - "Sick Lung Circulation"
Figure 4: Integration of hemodynamic and cellular events that characterize the cardio-pulmonary disease of severe pulmonary arterial hypertension. The pathophysiology of PAH which operates with mechanical concepts is complemented with concepts of cell injury and repair and interactions of complex multicellular systems. The model is built on the principle of abnormal pulmonary blood flow. Regional blood flow abnormalities underlie the pulmonary vascular remodelling, which in turn causes further alterations of the regional blood flow, perpetuating a cycle of cell death and cell proliferation. Vascular rarefaction occurs in the form of pruning of the small lung vessels and in the right ventricle in the form of myocardial capillary rarefaction.
Figure 4

- Hyperdynamic circulation
- Pulmonary Vasoconstriction
- Abnormal Pulmonary Blood Flow
- Endothelial cell injury and apoptosis
- "Wound healing gone awry"
- Proliferation, Apoptosis-resistance
- Angio-obliterative remodeling
- Vascular rarefaction
- Left-to-Right shunt
- Pulmonary Hypertension