Protein Losing Enteropathy after Fontan Surgery – Clinical and Diagnostical Aspects

Attila Tárnok József Bocsi Dominik Lenz Jan Janousek
Department of Pediatric Cardiology, Cardiac Center Leipzig GmbH, University of Leipzig, Germany

Key Words
Protein losing enteropathy · Univentricular heart · TCPC · Hypoproteinemia · Elevated fecal alpha1 antitrypsin · Immunosuppression · T lymphocytes

Summary
Protein losing enteropathy (PLE) is the massive enteric loss of serum protein. PLE may appear in several diseases associated with intestinal mucous membrane damages with or without infection. PLE is mostly associated with total cavopulmonary connection (TCPC) or Fontan-type circulation in patients with a functionally univentricular heart. TCPC is performed at an age of about 2 years or older and has a high survival rate of >90%. Time of PLE onset after TCPC is variable, the exact etiology is still unclear. Increased central venous pressure due to the absence of a subpulmonary ventricle may be a main reason for PLE, affecting 2–15% of the patients with a survival rate of 40% at 5 years and 20% at 10 years. Also immunological reasons for PLE are suspected. Major clinical signs are edemas, ascites, pleural effusion, diarrhea, malnutrition, fatigue, weight loss, and reduced physical development. The most impaired laboratory signs are elevated fecal alpha1-antitrypsin and alpha1-antitrypsin clearance, hypoproteinemia (hypoalbuminemia, hypo-gamma-globulinemia), lymphopenia (selective T helper cell loss) and secondary lymphangiectasia. Therapy considerations should have the aim to decrease central venous pressure to improve hemodynamics. Medical treatment consists of substitution of e.g. albumin, gamma-globulin, glucocorticoid, heparin or calcium, but still >60% of the patients remain symptomatic.

Schlüsselwörter
Eiweißverlustsyndrom · Univentrikuläres Herz · TCPC · Hypoproteinämie · Erhöhtes fäkales Alpha-1-Antitrypsin · Immunsuppression · T-Lymphozyten

Zusammenfassung
Protein Losing Enteropathy

Protein losing enteropathy (PLE), a massive enteric loss of serum protein, is usually characterized by a combination of different symptoms which do not have to be present at the same time. PLE appears in several diseases, e.g. in autoimmune diseases such as systemic lupus erythematosus (SLE) or celiac disease, and is often associated with intestinal mucous membrane damages with or without infection.

The medical history, physical examination and some laboratory parameters (table 1) are also specific for diseases associated with Fontan surgery (total cavopulmonary connection, TCPC). PLE is rare complication in patients who underwent Fontan surgery which is used to palliate complex congenital heart defects with an univentricular physiology [1]. Depending on the study, 9 [2] to 15% [3] of the survivors exhibit a substantial enteric protein loss. Time span between operation and the occurrence of PLE varies significantly from months to years after surgery. The mortality of patients with a manifest PLE is 46–59% after 5 years and >80% after 10 years [3, 4]. Up to now, the etiology of PLE is unknown.

Etiology / Risk Factors

The mechanism of action of PLE is poorly understood. The current theory of post-Fontan PLE is that chronic venous congestion causes the lymphatics to decompress into low-pressure cavities such as the pleural cavity or abdomen. Loss of proteins and lymphocytes, and an inflammatory response leads to chronic malnutrition resulting in a global immune dysfunction [5].

Systematic investigations correlating the risk factors for PLE with clinical parameters suggest elevated systemic venous pressure as a major contributory factor, which is the inherent hemodynamic component of the Fontan circulation [3]. However, there is no strict hemodynamic difference between those patients who develop PLE and those who do not [3]. Several studies suggest at least a participation of the immune system in the pathogenesis of PLE [6–8].

Recent observations indicate that onset of PLE is the result of multiple insults. It has been suggested that viral or bacterial infection, mostly of the gastric tract, can serve as trigger of PLE onset [7–10]. Jejunal biopsies, taken during episodes with PLE, revealed an increased IFN-γ concentration [11], most likely as a response to the viral infection and elevated levels of the pro-inflammatory cytokine TNF-α [12]. Both cytokines are known to impair the integrity of the intestinal epithelial barrier [13]. Similar to other primary diseases associated with PLE (e.g. SLE or celiac disease), episodes of post-Fontan PLE are characterized by a loss of heparan sulfate proteoglycan (HSPG), particularly from the basolateral surface of intestinal epithelial cells. However, neither HSPG expression in the lamina propria nor expression of other matrix components is influenced. In addition, the intestinal architecture remains intact. The reasons why HSPG is lost during episodes of PLE are still unknown [14].

Genetic factors could play a role in susceptibility to PLE development. Patients with univentricular heart differ from a healthy cohort with respect to the blood coagulation system [15] or the immune system [8] already prior to Fontan palliation. This finding may indicate genetic abnormalities. Reduced heparan sulfate accumulation in enterocytes contributes to PLE in a congenital disorder of glycosylation [16]. Also mesenteric hypoperfusion due to the increased mesenteric vascular resistance could serve as a possible trigger for the PLE development [17]. The results of Chaloupecky et al. [18] support the hypothesis that the abnormalities in the balance of coagulation factors observed in patients after Fontan operation are related to the coagulation factor production in the liver. The passive lymph loss caused by high central venous pressure does not explain the selective loss of CD4+ lymphocytes. Thus, it was suggested that the disturbance of the immune system could affect the structural integrity and patency of the intestinal wall, thereby triggering PLE [19]. The most likely explanation is that genetic factors and Fontan-induced venous hypertension predispose for PLE which appears upon a series of sequential or simultaneous environmental insults.

Clinical Description

Clinical signs for PLE are edemas, ascites, pleural effusion, diarrhea, malnutrition, fatigue, and weight loss as well as reduced physical development in children. The laboratory parameters required for the differential diagnosis of PLE are summarized in table 1 [data from 7, 8, 12, 19, 20]. Based on these very heterogeneous laboratory findings, PLE after Fontan procedure must be differentiated from - other diseases with hypoproteinemia, - other primary intestinal mucous membrane damages or primary lymphangiectasia,

Table 1. Required laboratory parameters for diagnosis of PLE

- Elevated fecal α1-antitrypsin secretion (normal range < 100 µg/g)
- Increased α1-antitrypsin serum level (normal range 0.88–1.74 g/l)
- Decreased total serum protein level (normal range 66–87 g/l)
- Decreased serum albumin level (normal range 34–48 g/l)
- Decreased serum γ-globulin and increased α2- and β-globulin level (normal ranges: γ-globulin 4.5–12.1 g/l, α2-globulin 0.9–2.9 g/l, β-globulin 5.4–9.3 g/l)
- No trypsinogen or enteroalkinase deficiency
- Lymphopenia < 1,500 cells/µl (normal range 1,500–3,000 cells/µl)
- Possible acute inflammation (mostly at the onset of PLE increased IL-6, IL-8, TNF-α serum levels, neutrophilia)
- T cell loss, T helper (T4) cell count massively reduced (down to < 100 cells/µl) (normal range 800–1,500 cells/µl)
- Protein level in urine is normal (<0.15 g/24 h)
– congenital enzyme defects (e.g. trypsinogen or enterokinase deficiency), malabsorption, nutrition sensitivity or allergic reactions (gluten, milk components),
– nephrotic syndrome (low serum concentrations of γ-globulin coinciding with elevated α1-, α2- and β-globulin levels are present in both PLE and nephrotic syndrome, but in contrast to nephrotic syndrome patients PLE patients generally do not exhibit proteinuria),
– acquired (e.g. AIDS) or congenital immunodeficiency because of decreased T helper (T4) cell counts (up to <100/µl) and γ-globulin levels.

The diagnosis of PLE is unspecific, even if there is always an elevated fecal concentration of α1-antitrypsin indicating a highly elevated α1-antitrypsin clearance. Nevertheless, since α1-antitrypsin is an acute-phase protein, its increased occurrence in the stool did not allow for an unequivocal doubtless diagnosis of PLE, as this symptom is also associated with other conditions such as enteritis of different origin. In conclusion, it must be emphasized that PLE occurs rarely and in combination with other nonspecific symptoms. The clinical signs become significant most often after the patient underwent Fontan palliation. PLE with other etiologies such as SLE, pericarditis or after superior vena cava-right pulmonary artery shunt (Glenn) is extremely rare. Many patients suffer from infections various harmless origins while developing PLE [7]. For this reason patients are rarely examined by a cardiologist at the very beginning of PLE occurrence. Often patients already had edemas caused by hypoproteinemia before they are admitted to a cardiac intensive care unit.

PLE is often associated with intestinal lymphangiectasia [3, 4, 21]. Although not yet routinely used for diagnosis, the composition of the peripheral blood of patients with PLE shows a pathognomonic change, i.e. highly selective lymphocyte loss (T4 (CD4+) cell count < 100 cells/µl) [7, 19, 20]. This selective cell loss which is not yet completely understood leads to a dramatic decrease of the T4/T8 cell ratio similar to that observed in patients with HIV [22, 23]. But in contrast to HIV patients opportunistic infections are rare in PLE patients [20]. The lymphocyte loss leads to the prevalence of unique T lymphocyte subtypes (CD4– CD8–) in the blood circulation [6, 24]. This phenotype of T lymphocytes is known from the autoimmune lymphoproliferative syndrome (ALPS) [25]. It is noteworthy, that already children with Fontan circulation, but without PLE, have an altered immune phenotype [24] compared to age-matched healthy children [26]. However, also the immune phenotype of children with other congenital heart diseases may differ from that of healthy age-related subjects [27].

Therefore a key tool is immunodiagnostics using either flow cytometry [28] or microscopic (or slide-based) cytometry [29]. Both technologies [30] yield quantitative data on the phenotypes and functions of leukocytes and lymphocytes [23, 31, 32] and are increasingly important tools in hemotherapy and transfusion medicine [33, 34]. Furthermore, cytometry can be used to determine the concentration of multiple soluble blood compounds by multiplexed bead arrays using very small sample volumes [35–37].

Management including Treatment

The prognosis of patients suffering from PLE is poor. Since the pathogenesis is not yet fully understood, there is a broad range of treatments suggested, mostly in single case reports. Therapy considerations should have the aim to decrease central venous pressure to improve hemodynamics. Relief of potential obstructions in the Fontan pathway, conversion of a classical atropulmonary anastomosis with loss of kinetic blood energy due to severe atrial dilatation to a total cavopulmonary connection or creation of a fenestration between the systemic venous and pulmonary venous atrium are possible surgical options.

The current medication consists of the substitution of albumin and/or γ-globulin [38] in combination with diuretics but has often only a temporary effect. In few patients PLE resolves without any obvious correlation to therapy. Case reports indicate that treatment with steroids, heparin, somatostatin analogues or high-dose spironolactone can lead to an improvement [11, 12, 21, 39]. Thus, anti-inflammatory therapy can at least temporarily reduce TNF-α levels, and substitution of heparin can diminish the protein loss by stabilizing the cell membrane of the intestinal mucosa [39, 40]. Cure of PLE in sarcoidosis by calcium substitution or anti-TNF agents indicating allergic or other immunological etiologies have been demonstrated. Furthermore, special diets (high protein, low fat with medium-chain triglycerides and essential fatty acids) are suggested. None of these treatments seem to be successful for all patients, reflecting the multifactorial origin of PLE. Correspondingly, a detailed European study showed that, in spite of surgical intervention or medication, still >60% of the patients remain symptomatic [3] and that the 10-year survival rate of patients suffering from PLE is only 10–20%. For some, but not all, patients with PLE the ultimate option is a heart transplantation [41].

Prospects

Intensive studies about the reasons of PLE are needed to develop therapies in the future. In our opinion, the monitoring of the changes of the immune phenotype could be used to predict the PLE risk [8]. The difficulties of analyzing the changes in the immune phenotype could be overcome by using cytomeric blood analysis [33]. The use of multiple analytes for a rapid analysis of a multitude of cells and for the recognition of specific cell types [29, 42] in combination with mathematical analysis [9, 28, 43] could help to assess the risk of patients and to individualize the therapy [44, 45].
References