Introduction

Acute respiratory distress syndrome (ARDS) is one of the most common respiratory acute and critical diseases in newborns. It refers to the acute inflammatory reaction of the lung caused by various pathogenic factors inside and outside the lung. It is characterized by progressive dyspnea, intractable hypoxia and decreased lung compliance [1]. With the vigorous development of neonatal intensive care unit, the survival rate of newborns, especially premature infants, has increased year by year, and the incidence of ARDS has also increased significantly, but the mortality rate is still high [2]. In view of the fact that the treatment of neonatal ARDS is only limited to the comprehensive treatment based on the corresponding respiratory support therapy, it is the focus of every neonatal pediatrician to understand the pathogenesis and diagnostic criteria of ARDS and actively take reasonable treatment measures.

Evolution of Ards Definition

Since Ashbaugh et al [3] first put forward the concept of ARDS in the 1960s, pediatricians have realized that children’s ARDS is different from adult ARDS. The 1994 American-European Consensus Conference (AECC) defined Acute Lung Injury (ALI) as $\frac{P_{aO_2}}{F_{iO_2}} \leq 300\text{mmHg}$, ARDS as $\frac{P_{aO_2}}{F_{iO_2}} \leq 200\text{mmHg}$, and later referred to “Acute” as the double meaning of “acute” rather than “adult or acute”, in order to accurately reflect the fact that this syndrome can occur in both adults and children, the diagnostic criteria of ARDS have been widely used in adults and children [4]. In 2012, the European Critical Care Association presided over the revision of the new diagnostic criteria for ARDS in Berlin, Germany. The Berlin definition addresses many of the limitations of the definition of AECC and believes that ARDS is a unique pathophysiological process. It can be described by time, imaging changes and severity, and ARDS can be divided into mild, moderate and severe according to $\frac{P_{aO_2}}{F_{iO_2}}$ ratio [5]. This was followed by a multicenter clinical study conducted by the European Society of Pediatric and Neonatal Intensive Care (ESPNIC) to verify the effectiveness of the Berlin standard in children aged 1 to 18 months [6]. According to the AECC and the Berlin definition, Barreira et al. [7] conducted a multicenter study of children aged from 1 month to 15 years in PICU. The results showed that the mortality rate and ventilator use days of children with severe ARDS were significantly higher than those with mild to moderate ARDS. It is considered that compared with the definition of AECC, the Berlin definition is more effective in predicting mortality and leaving ventilator and can better distinguish the severity of ARDS in children.
Although the definitions of AECC and Berlin are the perfection and expansion of the diagnostic criteria of ARDS, neither of them clearly puts forward the diagnostic criteria of ARDS in children and newborns, and has some limitations in the application of ARDS in children, such as the determination of Pao2 and the calculation of Pao2/Fio2 ratio. The two definitions did not take into account the etiology, risk factors and pathophysiologic differences between adults and children with ARDS. In view of the fact that there is still no specific definition and corresponding clinical trials of ARDS in children, the Paediatric Acute Lung Injury Consensus Conference (PALICC) proposed the definition and classification of Pediatric Acute Respiratory Distress Syndrome (pARDS) for the first time. To clarify the pathogenic factors, etiology and pathophysiology, put forward suggestions for treatment, and determine the focus of research. The PALICC standard states that pARDS includes children of all ages from newborn to adolescence and is unique in pathophysiology etiology and high-risk factors [9]. PALICC standard not only reduces the mortality of ARDS in children and adults, but also provides a strong guarantee for improving the quality of life of patients [10].

Table 1: Diagnostic criteria of neonatal Acute Respiratory Distress Syndrome (ARDS).

| Timing | AECC criteria | Berlin criteria | PALICC criteria | Montreux criteria |
|--------|---------------|-----------------|-----------------|-------------------|
| Age    | None          | None            | Excrete patients with peri-natal related lung disease | None            |
| Chest Imaging | Bilateral infiltrates observed on frontal chest radiograph | Bilateral opacities—not fully explained by effusions, lobar/lung collapse, or nodules | Chest imaging findings of new infiltrate(s) consistent with acute pulmonary parenchymal disease | Diffuse, bilateral, and irregular opacities or infiltrates, or complete opacification of the lungs, which are not fully explained by local effusions, atelectasis, RDS, TTN, or congenital anomalies |
| Causes of pulmonary edema | ARDS; Pao2 / Fio2 ≤200 mmHg (regardless of PEEP) | Respiratory failure not fully explained by cardiac failure or fluid overload | Respiratory failure not fully explained by cardiac failure or fluid overload | Absence of congenital heart disease explaining the oedema (this includes ductus arteriosus with pulmonary overflow if no acute pulmonary haemorrhage exists). Echocardiography is needed to verify the origin of oedema |
| Oxygenation | ARDS: Pao2 / Fio2 ≤200 mmHg with PEEP or CPAP | Mild | Severe | Invasive mechanical ventilation |
| | Mild | Moderate | Severe | Mild | Moderate | Severe |

PEEP: positive end-expiratory pressure; Pao2: arterial oxygen tension; Fio2: inspiratory oxygen fraction; SpO2: arterial oxygen saturation measured by pulse oximetry; CT: computed tomography OI = oxygenation index, OSI = oxygen saturation index. OI = (Fio2 x mean airway pressure x 100) / Pao2. OSI = (Fio2 x mean airway pressure x 100) / SpO2. TTN = transient tachypnoea of the neon. |
Although the PALICC standard has achieved a major breakthrough in the age of the ARDS standard, it specifically excludes Perinatal specific diseases such as acute hypoxia in newborns and secondary severe lung injuries such as meconium aspiration syndrome and congenital disseminated pneumonia. The clinical characteristics of neonatal ARDS have not been pointed out [11]. On the basis of the existing research on ARDS, the international multi-center and multi-disciplinary assistance group established the diagnostic criteria of neonatal ARDS for the first time in 2017, that is, the Montreux criteria [12]. At present, the Montreux criteria is the first ARDS diagnostic standard for newborns in the world. It is consistent with the diagnostic criteria of ARDS in children and adults as far as possible. It not only emphasizes the diagnostic criteria, exclusion criteria and imaging examinations that need to be met. The applicable age of neonatal ARDS, the standard of oxygenation disorder and the threshold of oxygenation index to distinguish mild, moderate and severe were also clarified. Montreux criteria is the expansion and perfection of ARDS consensus guide for children (PALICC standard). It plays a positive role in the diagnosis and treatment of critical newborns (Table 1).

Pathogenesis of Ards

Although the pathogenesis of ARDS has not been fully understood, current studies have shown that systemic inflammation is a key link in the occurrence and development of ARDS, severe inflammation leads to changes in vascular permeability, resulting in acute pulmonary edema [13-15]. After primary injuries such as infection, trauma and chemical factors affect alveolar epithelial cells and vascular endothelial cells, the boundary between alveolar epithelial cells and vascular endothelial cells is destroyed, and pulmonary capillary barrier dysfunction leads to increased permeability. Promote inflammatory cells to enter the alveolar cavity and pulmonary capillaries, protein-rich liquid quickly enter the lung tissue to cause acute pulmonary edema [16-17]. The results showed that the activity of type II secretory phospholipase A2 (sPLA2) was increased in both adults and children with ARDS. SPLA2 could promote inflammation and decompose surfactant phospholipids directly by hydrolyzing Dipalm Oil-Phosphate Diol-Cholinesterase (DDPC). At the same time, oxidative hydrolysis activated by inflammatory reaction can increase the degradation of PS, both of which can cause secondary deficiency of PS, resulting in hyaline membrane formation and alveolar collapse [18]. Therefore, pulmonary inflammation caused by inflammatory diseases, which leads to abnormal PS, is an important reason for the occurrence of ARDS.

Neonatal period is a unique stage which is different from adults and children. It not only has high mortality, but also has particularity in the induction and pathological characteristics of ARDS. Neonatal ARDS can be triggered by direct injury of the lung parenchyma (that is, direct or primary ARDS), such as Meconium Aspiration Syndrome (MAS), pneumonia, or extrapulmonary processes (that is, indirect or secondary ARDS), such as septicemia, necrotizing enterocolitis, Perinatal asphyxia and so on. In addition, some studies have found that the incidence of neonatal ARDS is also related to preterm delivery, caesarean section, diabetic mother, placenta previa and acidosis. It can be seen that the pathogenesis of neonatal ARDS is complex. How to treat ARDS from the aspect of pathophysiology is worthy of more in-depth study (Figure 1).

**Figure 1: Pathophysiological mechanism of neonatal Acute Respiratory Distress Syndrome (ARDS)**

Crs=respiratory system compliance. Rrs=respiratory system resistance.
Prevention and Treatment of ARDS

At present, there are no effective treatment measures for neonatal ARDS, the main treatment methods include respiratory support, PS replacement, extracorporeal membrane oxygenation therapy, nutritional support and liquid management. How to treat ARDS is the key issue of current research.

Treatment of Primary Diseases

A variety of internal and external factors such as meconium aspiration syndrome, pneumonia, perinatal asphyxia, septicemia, neonatal necrotizing enterocolitis and so on can cause neonatal ARDS. Therefore, it is the primary task of prevention and treatment of ARDS to take corresponding treatment measures for the primary disease in the early stage and curb the systemic uncontrolled inflammatory reaction induced by it.

Respiratory Support

Mechanical Ventilation (MV) is an important treatment for ARDS, but inappropriate mechanical ventilation can cause lung injury in children [19-20]. The lung pathology of children with ARDS is characterized by inflammation, partial atelectasis and normal alveoli at the same time. Therefore, in this case, if the traditional mechanical ventilation mode of continuous application of climax volume and inappropriate end-tidal pressure is used, at the same time of opening the collapsed alveoli, the normal alveoli were overexpanded, resulting in normal alveolar injury, that is, ventilator-induced lung injury (ventilator induced lung injury;VILI) [21-22]. Air pressure injury, volume injury, lung injury caused by alveolar overexpansion and atelectasis after low lung volume ventilation were all related to VILI. In order to avoid pulmonary overexpansion and pulmonary wilting injury, it is recommended to use Pulmonary Protective Ventilation Strategy (PLVS), that is, low tidal volume and appropriate PEEP to reduce lung volume injury and lung collapse injury as much as possible [23-24]. A large sample randomized controlled trial in the United States confirmed that low tidal volume and limited platform pressure can reduce the mortality of ARDS [25].

Although PLVS is a major breakthrough in the treatment of ARDS, the latest report points out that the alveoli are still overinflated when PLVS is used, and patients are still at risk of developing VILI. The lower the tidal volume and platform pressure, the more beneficial to the patients. Therefore, some scholars have proposed the concept of Ultra-Protective Lung Ventilation Strategy (UPLVS), that is, it is suggested that the tidal volume should be changed from 6ml/kg to 4ml/kg, and the platform pressure should be changed from 30mmHg to 25mmHg [26-27]. Although lower tidal volume and lower platform pressure in UPLVS can improve the rate of ARDS patients, it can increase the risk of respiratory acidosis, high respiratory rate ventilation and high PEEP ventilation [28]. Both PLVS and UPLVS use lower tidal volume to reduce pulmonary overexpansion but ignore the simultaneous alveolar collapse in children with ARDS. In order to solve this problem, some scholars suggest that intermittent administration of pressure higher than conventional breath pressure during mechanical ventilation and maintaining it for a period of time can not only restore collapsed alveoli, but also prevent repeated alveolar dilatation and collapse, that is, Lung Recruitment Maneuver (LRM) [29-31]. However, at present, this strategy is rarely used in newborns, and large sample multicenter studies are rare, so the method and effectiveness of LRM in neonatal ARDS remains to be further confirmed by clinicians.

In view of the fact that invasive ventilation can lead to ventilator-associated pneumonia and other complications, some scholars have proposed that it is a positive pressure ventilation support technique without endotracheal intubation, that is, Non-Invasive Positive Pressure Ventilation (NIPPV) [32]. At present, a number of studies have confirmed the effectiveness and safety of NIPPV as a first-line treatment to avoid endotracheal intubation in patients with ALI/ARDS [33-36]. The consensus meeting on Acute Lung injury in Children recommended that children with ARDS should use NIPPV, and PEEP noninvasive ventilation at an early stage of the disease [8]. Recent studies have shown that NIPPV can also be used for the transitional treatment of ventilator and extubation in children with ARDS, so further exploring the prospect of NIPPV in the treatment of ARDS has been the focus of pediatricians.

High Frequency Oscillatory Ventilation (HFOV) is a new mechanical ventilation mode developed in recent years. HFOV uses hyperphysiological ventilation frequency concussion to maintain bi-directional pressure, and the ventilation frequency is 4 times higher than the normal frequency. Gas exchange at low tidal volume and constant airway pressure can not only improve oxygenation and efficient clearance of $CO_2$, but also avoid complications such as lung injury caused by overexpansion of the lungs. It is the most advanced high frequency ventilation technology at present [37-40]. Many studies at home and abroad have shown that although HFOV cannot reduce the mortality of children with ARDS, compared with Conventional Mechanical Ventilation (CMV), HFOV can significantly improve oxygenation [41-42]. The latest guidelines recommend that in the absence of clinical evidence of decreased chest wall compliance, HFOV is considered an alternative ventilation mode in patients with moderate and severe ARDS whose respiratory plateau pressure exceeds $28cmH_2O$. It is suggested that the oxygenation, $CO_2$ response and hemodynamic variables should be continuously monitored in HFOV to explore the potential of lung retension by gradually increasing and reducing continuous dilatation pressure, so as to obtain the best lung volume. However, High-Frequency Jet Ventilation (HFJV) and High-Frequency Percussive Ventilation (HFPV) are not recommended in children with ARDS [8].

Prone position ventilation is a commonly used non-mechanical ventilation method for neonatal ARDS. One of the common pathological changes in children with ARDS is the decrease of lung volume, which is mainly characterized by less functional residual...
Pulmonary Surfactant (PS) is a phospholipid formed and stored by type II alveolar epithelial cells. It can not only reduce alveolar surface tension to prevent alveolar collapse at the end of breath, but also increase pulmonary compliance. At the same time, it also has anti-inflammatory and antibacterial effects [57-59]. Because most newborns have primary or secondary PS deficiency in ARDS, the mortality of neonatal ARDS has been reduced by 50% since the emergence of exogenous PS replacement therapy [60]. The results of Wang LP et al. [61] showed that early application of PS combined with mechanical ventilation could significantly improve pulmonary oxygenation and compliance and inhibit inflammation in children with ARDS. A multicenter randomized controlled trial confirmed that intratracheal infusion of PS not only improved oxygenation in patients with ARDS, but also significantly reduced mortality [4]. And studies have shown that improvements in surfactant occur only in patients with direct lung injury (pneumonia, inhalation or near drowning). To verify this conclusion, Willson et al. [62] conducted a multicenter randomized controlled study in 6 different countries in children with ALI/ARDS caused by direct lung injury. The results showed that intratracheal infusion of PS did not improve oxygenation. Therefore, exogenous PS is not recommended for children with ALI/ARDS. Therefore, the optimal use time, dosage and curative effect of PS in neonatal ARDS still need to be further explored.

Nutritional Support and Liquid Management

For children with ARDS, nutrition supply is not only the provision of energy, appropriate nutrition supply can prevent heat consumption, correct malnutrition, shorten the time of mechanical ventilation, regulate immune function and improve oxygenation in order to improve the prognosis of patients with ARDS [63-64]. Therefore, nutritional support as an important means of ARDS treatment has been widely used in clinical practice. A multicenter study of 500 children with PICU showed that children who ate more than 66 percent of the prescribed calories had a significantly lower mortality rate than those who received less than 33 percent of the prescribed calories [65]. Further studies by Wong JJ et al. [66] have shown that consuming enough protein can improve clinical outcomes better than eating enough calories. Neonatal nutritional support includes enteral nutrition and parenteral nutrition. A small amount of enteral nutrition combined with most parenteral nutrition is beneficial to the establishment of intestinal flora, maintain intestinal function and reduce intestinal complications. A randomized controlled trial shows that early rational use of enteral nutrition support can not only improve the clinical efficacy of ARDS patients, reduce the incidence of infection, help to control blood glucose, improve lung function. It can also shorten the time of mechanical ventilation and ICU hospitalization and reduce the hospitalization cost [67].

The main pathological features of neonatal ARDS are non-cardiogenic pulmonary edema, which can affect respiratory function through the following aspects: decrease of pulmonary compliance to increase respiratory work, increase of intrapulmonary shunt resulting in hypoxia, and aggravation of pulmonary hypertension. Combined mechanical ventilation can promote the occurrence of pulmonary inflammation and reduce the secretion of PS, thus aggravating the damage of pulmonary capillary barrier. Therefore, through liquid management to reduce the production of pulmonary edema fluid and promote the discharge of edema fluid is an important link in the treatment of ARDS. The goal of fluid therapy in critically ill children with ARDS is to ensure adequate end organ perfusion. A number of studies have confirmed that early fluid overload can aggravate the clinical outcome of children with ARDS [68-71]. The results of a Meta-analysis showed that strict fluid management strategy in adults and children with ARDS could
reduce the days of ventilator use and shorten the length of hospital stay [72]. At present, liquid overload management mainly includes non-invasive strategies such as diuretics and fluid restriction and invasive methods such as Continuous Renal Replacement Therapy (CRRT). Studies have shown that patients who start using CRRT earlier may have better clinical results [73]. On the premise of ensuring hemodynamic stability and perfusion of tissues and organs, restricted fluid management is helpful to improve the oxygenation and lung injury in patients with ARDS, but whether it can reduce the mortality of patients with ARDS remains to be further confirmed.

Other Treatment

Ambroxol hydrochloride not only has the characteristics of promoting mucous excretion and dissolving secretions, but also has antioxidant and anti-inflammatory effects, and can promote the production of pulmonary surfactant [74]. A Meta-analysis showed that \( P_aO_2/FIO_2 \), \( P_aO_2 \), and \( Sao2 \) increased only 7 days after high dose ambroxol (\( > 15mg/kg \) or \( 1000mg/d \)) treatment in patients with ALLARDS, which may be related to the antioxidant and anti-inflammatory properties of ambroxol [75]. Inhaled nitric oxide (iNO) in the treatment of ARDS is one of the most widely studied interventions in the past two decades. Although iNO treatment can improve oxygenation, there is still no study to prove whether there is a decrease in mortality [76]. A study of 161 children with ARDS showed that inhaling carbon monoxide could reduce the duration of mechanical ventilation, but there was no significant difference in mortality between the two groups [77]. Therefore, at present, iNO is not recommended for routine use in patients with ARDS, but it can be used in the rescue treatment of patients with severe intractable hypoxia [78]. Other treatments, such as liquid ventilation, volume target ventilation, Mesenchymal Stem Cell (MSC) and so on, still need further research and clinical trials to confirm [79-80].

Conclusion

Although great progress has been made in antenatal prevention and postnatal treatment of neonatal ARDS in recent years, there is still no single treatment that can significantly improve the survival of newborns. In view of the high mortality of neonatal ARDS and its specificity in etiology, pathophysiology and so on, the large sample multicenter prospective study on neonatal ARDS diagnosis and treatment is worth looking forward to.

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