Neonatal diseases and oxidative stress in premature infants: an integrative review

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Abstract
Objective: To describe the relationship of oxidative stress and antioxidant biomarkers in cord blood of premature newborns and the prognosis of diseases in the neonatal period.

Sources: This study consists of an integrative review. Searches were conducted in electronic databases Scopus, PubMed, Web of Science, and Medline/Lilacs through the Virtual Library on Health Issues, using the descriptors: "premature infants", "preterm infants", "preterm birth", "preterm", "oxidative stress", "antioxidants", "infant, premature, diseases" and "cord blood". Original articles published between 2016 and 2021 in Portuguese, English, or Spanish, which analyzed oxidative stress and/or antioxidant levels through cord blood of premature newborns and evaluated clinical outcomes, were included.

Summary of the findings: Of the 1,003 studies reviewed, after exclusion of duplicate articles, analysis of titles, abstracts, and full texts, 18 articles were included. 72.2% (n = 13) of analyzed studies reported a positive association between oxidative stress and the development of prematurity-related diseases; 27.7% (n = 5) showed no significant relation. Outcomes that showed a positive association were: intrauterine growth restriction, necrotizing enterocolitis, bronchopulmonary dysplasia, intraventricular hemorrhage, fetal inflammatory response syndrome, early-onset neonatal sepsis, retinopathy of prematurity, morbidity, and mortality.

Conclusion: The analysis of oxidative stress and antioxidants in cord blood of premature newborns may be useful in the prognosis of some pathologies. The consequences of oxidative damage are known to be associated with increased morbidity in the short and long term. Further investi-
Introduction

Prematurity is a major public health problem. Complications from premature birth are the leading cause of death in children under five years of age. In 2017, of the 2.5 million newborns who died from preventable causes, nearly two-thirds were premature.

Oxidative stress has been implicated as a possible pathophysiological condition to contribute to this unfavorable situation. Premature neonates lack well-developed antioxidant and immune defense mechanisms, making them more susceptible to oxidative stress injury. The main reasons for this susceptibility are a hypoxic-hyperoxic challenge, presence of infections, deficiency in antioxidant defense, and high levels of free iron.

The transition from intrauterine to the extrateruterine environment greatly increases free radical production, which is normally downregulated by the antioxidant defense system. Oxidative stress is caused by excessive reactive oxygen species (ROS), which are generated when there is an imbalance in this regulation. In oxidative stress, there is an inability of the antioxidant defense system to repair the ROS damage due to either an excessive formation or impaired inactivation of ROS, or a combination of both.

ROS include free radicals and oxygenated molecules of non-free radicals. Both of them can generate oxidative stress and redox reaction imbalance. Oxygen-free radicals are extremely reactive chemical species; they react with various cellular molecules — such as phospholipids, amino acids, and nucleic acids — and lead to lipid peroxidation, DNA strand breaks, and other damaging processes which culminate in cellular injury.

Oxidative stress-induced damage has an important role in several pathological pathways involved in neonatal diseases. For example, some studies point out that most complications of prematurity, such as bronchopulmonary dysplasia (BPD), retinopathy of prematurity (ROP), necrotizing enterocolitis (NEC), intraventricular hemorrhage (IVH), periventricular leukomalacia (PVL), and white matter lesions seem to be related to oxidative injury.

According to Özsurecko et al., there are still gaps in knowledge about the potential role of oxidative injury in the pathogenesis of neonatal diseases. New studies should be conducted to investigate more extensively diagnostic and prognostic values of various oxidative stress and antioxidant biomarkers in order to reduce oxidative tissue injury in neonates.

Cord blood has been the most widely used matrix when evaluating mechanisms related to prematurity as are allow monitoring of biomarkers without causing any distress to the babies and providing relevant information.

From this perspective, the objective of this study is to describe the relationship between oxidative stress and antioxidant biomarkers in cord blood of premature newborns and the development of diseases in the neonatal period of preterm newborns.

Method

This study consists of an integrative literature review, which is a method that enables the analysis and synthesis of results in a systematized manner, providing support for decision making and improvement of clinical practice. The preparation of this review respected all the pre-established phases for its realization, covering the following steps: formulation of guiding questions; selection and retrieval of articles according to inclusion and exclusion criteria; data collection; evaluation of selected studies; discussion and interpretation of results, and presentation of the review.

Studies that evaluated biomarkers of oxidative stress and/or antioxidant levels in cord blood of preterm newborns and evaluated clinical outcomes were included. Searches were conducted from January to March 2021, and studies published between 2016 and 2021 in Portuguese, English, or Spanish were included. Original peer-reviewed articles and indexed in the electronic databases Scopus (Elsevier), PubMed (via National Library of Medicine), Web of Science (main collection), and Medline/Lilacs through the Virtual Health Library were chosen. Specific descriptors and their synonyms were used, according to the Medical Subject Headings (MeSH) terms and their equivalents in Portuguese, established by the Health Sciences Descriptors (DeCS). The terms were combined using the Boolean operators “AND” and “OR” to compose the search strategy. The following terms were used in searches: “premature infants” OR “preterm infants” OR “preterm birth” OR “preterm” AND “oxidative stress” OR “antioxidants” AND “infant, prematurity, diseases.” The search equation used was: (“premature infants” OR “preterm infants” OR “preterm birth” OR precem) AND (“oxidative stress” OR antioxidants) AND diseases.

To be included in this review, studies should evaluate oxidative stress and/or antioxidant biomarkers in cord blood of preterm infants (i.e. born before 37 weeks of gestational age (GA)) and associate them with a clinical outcome (a clinical condition, disease or epidemiological indicators, such as morbidity or mortality). Quantitative or qualitative data were extracted from included articles. Studies were excluded if they were duplicate publications; review studies; if they evaluated oxidative stress and/or antioxidants in premature newborns but did not associate it with a clinical outcome; studies that did not evaluate oxidative stress and/or antioxidants using cord blood; experimental studies with objective out of the scope of this review.

Two authors independently screened the titles and identified by the searches, and those which met the eligibility criteria were selected for the full-text review. The selected full-text articles were further evaluated by two independent
authors, and the studies were definitively included in the review when they met all the inclusion criteria. Any differences between the two reviewers were resolved through a third independent author. The analysis of the studies was conducted in a systematized way, using a structured instrument containing the following information: title of the article; journal; database; Qualis evaluation; authors; country; language; year; objectives; sample characteristics; analyzed variables; data analysis; results; and conclusions. The studies were critically analyzed regarding their authenticity, methodological quality, the relevance of the information, and representations to ensure the scientific integrity of the review.

Results

One thousand and three articles were identified by the search strategy. Were excluded 968 articles after title/abstract analysis and after excluding duplicates, so 35 articles were fully assessed for eligibility. Two additional studies were included by assessing article references. After full-text analysis, 19 articles were excluded because they did not meet the inclusion criteria. Thus, a total of 18 studies were included in the review. Figure 1 shows the flowchart of the article selection process. Included studies were presented as follows concerning the year of publication: 5 in 2016, 3 in 2017, 2 in 2018, 3 in 2019, 3 in 2020, and 2 publications in 2021. Synthesis of the articles, year of publication, type of study, sample, source of specimen, biomarkers, diseases evaluated, and the main outcomes are described in Table 1. 3-5,8-10,12,13,19-28

Figure 1 Flowchart of the study selection process for the integrative review on oxidative stress in premature newborns.
| Article / Country       | Type of study | Sample | Source of specimen | Oxidative stress and antioxidants biomarkers | Diseases evaluated | Main outcomes                                                                                                                                 |
|------------------------|---------------|--------|--------------------|---------------------------------------------|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------|
| Moore et al., 2016[^1] | Prospective   | 31 PTN | Cord blood and urine | 8-OHdG                                      | CLD, IVH, BPD, ROP  | CLD was associated with lower levels of 8-OHdG. There were no significant differences regarding IVH, BPD, and ROP incidences.                  |
| Perrone et al., 2016[^3] | Cohort        | 120 PTN | Cord blood          | IsoPs, NPBI, AOPP                          | IUGR                | Placental injuries (inflammation or impaired perfusion) were associated with elevated OS levels. PTN with vascular perfusion lesions had higher levels of AOPP, low GA, and the IUGR. |
| Dietze et al., 2016[^4] | Prospective   | 31 PTN | Cord blood and saliva | Cortisol                                    | NEC, IVH, BPD, ROP  | The higher the GA at the beginning of prenatal care, the lower cord blood cortisol and tended to have a higher risk of NEC. In salivary cortisol, NB whose mothers smoked had a higher risk of IVH. |
| Ghany et al., 2016[^5] | Case-control  | 200 NB  | Cord blood and venous blood | MDA; TAC, Catalase, Vitamin A, Vitamin E | NEC, IVH, BPD, ROP  | Positive relationship between reduced antioxidant levels at birth and the risk of neonatal morbidities, including BPD, IVH, NEC.               |
| Bandypadhyay et al., 2017[^6] | Transversal  | 109 NB  | Cord blood          | MDA, 8-OHdG                                 | IUGR                | Term NB and late PTN with IUGR had higher levels of MDA and 8-OHdG compared to infants suitable for GA through meconium-stained amniotic fluid. |
| Norishadkam et al., 2017[^7] | Case-control study | 50 NB  | Cord blood          | MDA, Catalase, SOD, TAC                      | DNA damage          | There was no significant association between MDA, SOD, TAC, catalase, and early DNA damage in cord blood plasma for PTN.                     |
| Ozalkaya et al., 2017[^8] | Prospective   | 51 PTN | Cord blood          | TAC, PON-1, TOS, OSI                        | FIRS, RDS, IVH, BPD, ROP, Sepsis | Higher levels of TOS were associated with NB with PPROM; Higher PON-1 was associated with higher risk of PPROM, FIRS or both. NB with PPROM and FIRS had higher incidence of RDS. NB with FIRS had higher mortality. |
| Bharadwaj et al., 2017[^9] | Cohort        | 143 NB  | Maternal blood and cord blood | Protein carbonyls, MDA, TAC                 | Neurologic desease, IUGR, Sepsis | OS increases in NB and mothers with PE. Decreased maternal TAC is associated with negative neuro-motor results. Maternal TAC during PE is useful to predict poor motor development at the corrected age of one year. |
| Coutinho et al., 2018[^10] | Transversal   | 21 PTN | Cord blood, saliva, and urine | MDA, SOD, GPx, Catalase                     | Sepsis              | GPx in cord blood can be a diagnostic tool for suspicion of early-onset neonatal sepsis in PTNs of mothers with risk factors for sepsis. |
| Arman et al., 2018[^11] | Case-control study | 80 PTN | Cord blood          | TAC, TOS, OSI                              | Cardiac functions   | TAC, TOS, and OSI were significantly higher for NB of mothers with PE. Echocardiographic parameters are not affected by the oxidant state. |
| Article / Country                  | Type of study       | Sample          | Source of specimen | Oxidative stress and antioxidants biomarkers | Diseases evaluated | Main outcomes                                                                                                                                 |
|-----------------------------------|---------------------|-----------------|-------------------|--------------------------------------------|-------------------|-----------------------------------------------------------------------------------------------------------------------------------------------|
| Elkabany et al., 2019            | Egypt               | Prospective     | 40 PTN            | MDA; AOPP 8-OHdG TAC Copper; Zinc           | RDS               | AOPPs and 8-OHdG can be used as serum biomarkers for OS among NBs with RDS to monitor disease progression.                                  |
| Dekker et al., 2019              | Netherlands         | Clinical trial  | 52 PTN            | 8iPGF2α                                    | Grade III IVH     | 8iPGF2α did not differ among groups. There were also no differences regarding intubation, the incidence of grade III IVH, or death before hospital discharge. |
| Silva et al., 2019               | Brazil              | Transversal     | 140 NB            | Vitamin E                                  | IUGR              | IUGR was more frequent in PTB; most of the infants had low vitamin E levels.                                                                     |
| Stefanov et al., 2020            | United States       | Prospective     | 63 NB             | MDA Glutathione                            | Endothelial       | MDA was higher in cord blood than at 24 hours of life, regardless of GA. PTN had higher ET-1 levels in cord blood than 24 hours of life, but overall, ET-1 had no significant association with OS. |
| Liu et al., 2020                 | China               | Case-control    | 816 NB            | MDA SOD ROSs                               | Morbidity and mortality | Higher levels of MDA and ROSs are associated with lower Apgar scores; NICU admission and ventilation, i.e., are significantly associated with higher morbidity and mortality. |
| Pajai e Bezalwar 2020           | India               | Observational   | 45 PTN            | MDA Nitrites Vitamin C Vitamin E           | Morbidity and mortality | PTN shows higher MDA and nitrates levels and reduced levels of vitamins C and E, especially in males, indicating increased morbidity and mortality. |
| Agrawal et al., 2021             | India               | Nested case      | 189 PTN           | MDA Copper Zinc Vitamin A                  | ROP               | MDA and vitamin A in umbilical cord plasma were independent predictive variables of ROP.                                                              |
| Coviello et al., 2021            | Netherlands         | Prospective     | 44 PTN            | F2-isoprostanes                            | White matter injury | Early verification of plasma IsoPs can help discriminate abnormal WMI scores at term-equivalent age; and represents an early biomarker for identifying PTN at risk of brain injury. |

AOPP, advanced oxidative protein products; BPD, bronchopulmonary dysplasia; CLD, chronic lung disease; DNA, deoxyribonucleic acid; ET-1, endothelin-1; FIRS, fetal inflammatory response syndrome; FT, full-term; GA, gestational age; GPx, glutathione peroxidase; IVH, intraventricular hemorrhage; IsoPs, isoprostanes; IUGR, Intratrauterine growth restriction; MDA, malondialdehyde; NB, newborns; NEC, necrotizing enterocolitis; NICU, neonatal intensive care unit; NPBI, non-protein bound iron; OS, oxidative stress; OSI, oxidative stress index; PE= preeclampsia; PON-1, paraoxonase-1; PPROM, preterm premature rupture of membrane; PTN, preterm newborn; RDS, respiratory distress syndrome; ROP, retinopathy of prematurity; ROS, reactive oxygen species; SOD= superoxide dismutase; TAC, total antioxidant capacity; TOS, total oxidant status; WMI, white matter injury; 8-OHdG, 8-hydroxydeoxyguanosine; 8iPGF2α, 8-iso-prostaglandin F2.
The studies that reported a positive association between oxidative stress and/or development of diseases related to prematurity were 72.2% (n = 13); 27.7% (n = 5) showed no significant association. Positive correlations were found in IUGR,8,13,25 NEC,10,20 morbidity and mortality,5,26 BPD, IVH,10 FIRS,21 early-onset neonatal sepsis,22 RDS12 and ROP,27 CLD,19 DNA damage,4 IVH,24 cardiac functions,23 and endothelial dysfunction9 were not associated with oxidative stress levels in included studies.

Table 25,8,10,12,13,20-22,25-27 shows levels of oxidative stress and/or antioxidants correlated with clinical outcomes. It was observed the following associations: 1) MDA and vitamin A were associated with ROP,27 2) AOPP13, MDA, 8-OHdg8 and vitamin E25 were associated with intrauterine growth restriction; 3) Cortisol,20 vitamin A and vitamin E10 were associated with NEC; 4) Vitamins A and E were associated with BPD and IVH;10 5) PON-1 was associated with FIRS;21 6) GPx was associated with sepsis;22 7) AOPP and 8-OHdG were associated with RDS12 and 8) MDA;5,26 and ROS,26 nitrates, vitamin C, and vitamin E5 were associated with morbidity and mortality.

Three of the 13 studies which found an association between oxidative biomarkers and disease development used different samples to analyze data: venous blood drawn a few hours after birth,28 maternal blood,3 and salivary cortisol.20

Table 2 Oxidative stress biomarkers and antioxidants correlated with evaluated conditions.

| Condition          | Biomarker               |
|--------------------|-------------------------|
| ROP                | ↑ MDA27, ↓ Vitamin A27, AOPP13, ↑ MDA8, ↑ 8-OHdG8, ↓ Vitamin E25, ↑ Cortisol20, ↓ Vitamin A10, ↓ Vitamin E10, ↓ Vitamin A10, ↓ Vitamin E10 |
| IUGR               |                         |
| NEC                |                         |
| BPD                |                         |
| IVH                |                         |
| FIRS               | ↓ PON-11, ↓ GPx22, ↑ AOPP12, ↑ 8-OHdC12 |
| Sepsis             |                         |
| RDS                | ↑ MDA5,26, ↑ ROS26, ↑ Nitrates5, ↓ Vitamin C5, ↓ Vitamin E5 |
| Morbidity and mortality |                         |

AOPP, advanced oxidative protein products; BPD, bronchopulmonary dysplasia; FIRS, fetal inflammatory response syndrome; GPx, glutathione peroxidase; IVH, intraventricular hemorrhage; IUGR, intrauterine growth restriction; MDA, malondialdehyde; NEC, necrotizing enterocolitis; PON-1, paraoxonase-1; RDS, respiratory distress syndrome; ROP, retinopathy of prematurity; ROS, reactive oxygen species; 8-OHdG, 8-hydroxydeoxyguanosine.

In general, studies have shown that premature newborns have a higher level of oxidative stress biomarkers compared to term newborns due to several factors. Moreover, worse prognosis (low Apgar score, admission to the neonatal intensive care unit, assisted ventilation and hospital stay time) and increased morbidity and mortality have been related to higher levels of oxidative stress and lower levels of antioxidants with the development of some pathologies.

Discussion

In premature newborns, oxidative stress is a physiological event during the normal transition from the intrauterine to the extrauterine environment. Outside the uterus, free radical production increases significantly and must be counterbalanced by the antioxidant defense system. Healthy full-term newborns can tolerate these drastic changes; however, when intrauterine development is incomplete or abnormal, this tolerance might be affected. The identification of reliable biomarkers to analyze the oxidant-antioxidant system dysregulation is essential to improve neonatal care. Furthermore, evaluation of oxidative stress through cord blood may be useful in determining the prognosis of some pathologies.29

Most of the studies included in this review showed a relationship between increased levels of oxidative stress biomarkers and/or decreased levels of antioxidants in cord blood and higher risk of clinical outcomes, such as neonatal diseases and morbimortality. Among the conditions that showed a greater association with increased oxidative stress and/or reduced antioxidant levels were ROP,27 IUGR,8,13,25 RDS12, sepsis13 and morbidity and mortality.5,26 Falsaperla et al.11 also observed that the imbalance between the newborn’s oxidant and antioxidant factors seems to play an important role in the onset of the main pathologies of the preterm infant, such as BPD, ROP, NEC, IVH, periventricular leukomalacia, and white matter lesions.

One of the selected studies evaluated 189 premature newborns (38 with ROP and 151 without ROP) in a case-control study and showed that MDA and vitamin A concentration in cord blood are independent predictor variables of ROP.27 ROP is an eye disease that affects 7 to 15% of premature infants and is caused by abnormal vascular growth in the retina and may cause significant visual impairment or even blindness.20

To monitor ROP progression among neonates, AOPPs and 8-OHdG might be relevant biomarkers of oxidative stress. In a study conducted by Elkabany et al.,12 these markers were measured in the cord blood of 80 premature newborns at <34 weeks GA (40 newborns with RDS and 40 newborns without RDS), with a positive association. Another marker that might be considered as a diagnostic tool is GPx. One study evaluated 21 preterm infants (30 and 36 weeks) and showed that early-onset neonatal sepsis had a significant correlation with GPx levels in preterm infants from mothers with risk factors for this disease.22

A review by Casavant et al.30 showed that, compared to term infants, premature newborns had lower levels of antioxidants, vitamin A, vitamin E and catalase (an enzyme that neutralizes ROS and is linked to increased rates of NEC and BPD). In a case-control study conducted by Ghany et al.10
with 100 preterm and 100 full-term newborns, levels of vitamin A, vitamin E, catalase, TAC, and MDA were analyzed. The study described a significant relationship between decreased antioxidant levels at birth and the risk of neonatal morbidity, including BPD, NEC, and IVH. In addition, cortisol was another marker that showed a significant association with NEC. NEC is the main cause of morbidity and mortality in premature babies, with a mortality rate of up to 30% and an increased risk of delayed neurological development, especially in pregnancies with inadequate prenatal care.20

Among the reports which evaluated IUGR, Silva et al.25 analyzed the concentration of vitamin E in umbilical cord serum in 140 newborns (64 premature and 76 term) to test a correlation between the biomarker and intrauterine growth. Results showed that IUUGR was more frequent in premature newborns, and most of them had low vitamin E levels. IUGR is a complication of pregnancy, often described when the fetus is estimated to be small for the GA and has an incidence ranging between 3 and 7% of births. In the study by Perrone et al.13 with 129 premature newborns, it was observed that newborns with vascular perfusion lesions had higher levels of AOPP, low GA, and IUGR. Bandypadhyay et al.8 evaluated 109 newborns (27 premature and 82 term ones) and related IUGR to higher levels of AOPP, MDA, and 8-OHdG.

The study by Pajai and Bezalar5 associated higher morbidity and mortality rates with increased levels of MDA and nitrates and decreased levels of vitamins C and E, especially in premature male newborns. Liu et al.20 stated an association between MDA and ROS levels and a higher risk of a low Apgar score, NICU admission, and mechanical ventilation. Ozalkaya et al.21 determined TAC, PON-1, TOS, and OSI levels in 51 preterm infants (<34 weeks). They showed a significant association between PON-1 levels and PPROM and FIRS, alongside a higher incidence of RDS and higher mortality of infants with FIRS.

Alternative markers used in the studies are the antioxidant activity levels. Antioxidant activity measurement may be generally more useful than oxidative stress levels because the results allow a greater understanding of potential mechanisms and therapeutic interventions.14 In this review, antioxidants that showed a significant association between their levels and the development of morbidities were PON-1,21 GPx,22 and vitamins A, C, and E.5,10 It is essential to measure multiple antioxidants and include measures that identify specific ROS or RNS that may be associated with imbalanced antioxidant levels or activity. The inclusion of these additional measures provides a more comprehensive comprehension of the biological processes involving antioxidants.14

Also, data obtained from some studies reveal a positive association between oxidative stress and neurological diseases. However, these analyses were not obtained from cord blood. One of them stated that analysis of plasma IsoPs (24 and 48 h after birth) might represent an early biomarker to identify the risk of brain damage in premature patients. Bharadwaj et al.13 evaluated the neurodevelopment of 71 children and measured oxidative/antioxidant stress levels. They concluded that maternal TAC in PE is useful for predicting impaired motor development at one year of corrected age. Dietze et al.20 observed through salivary cortisol analysis that newborns whose mothers smoked more than ten cigarettes per day were at higher risk for IVH. IVH affects 30 to 60% of preterm infants and is characterized as white matter lesion due to microvascular events that occur in the germ matrix, putting the infant at increased risk for neurodevelopmental delays and additional brain damage.20

On the other hand, several studies did not confirm the association between oxidative stress biomarkers and the development of pathologies, morbimortality or prognosis. Moore et al.19 examined 31 premature newborns to evaluate the association between CLD and 8-OHdG levels. They observed that CLD was associated with lower levels of oxidative stress, which contradicts previous studies. Authors justify these conflicting results by different methodologies used in sample collection and analysis, alongside other additional factors that might have affected 8-OHdG levels. Dekker et al.24 evaluated 52 preterm infants for 8PGF2a levels in order to find a higher risk of developing IVH, but no significant association was found. Norishadkam et al.4 compared 25 premature and 25 term newborns to verify the relationship between oxidative stress and DNA damage and also found no significant association. Stefanov et al.5 evaluated the relationship between glutathione, MDA, and endothelin-1 levels to analyze endothelial dysfunction in 63 infants born between 24 to 42 weeks. Still, there was no significant relationship between the two factors analyzed. Finally, Arman et al.26 evaluated the global oxidant and antioxidant status in newborns of mothers with and without PE; there was no statistical difference between groups. The absence of significant associations in these studies may be justified by clinical factors affecting biomarkers levels, the small sample size in most studies, and the complexity of the neonatal transition period, in which a not fully comprehended mixed-redox state might occur.9

The present study has some limitations. Although the utility of cord blood samples for the assessment of oxidative stress biomarkers and their possible clinical implications is evident, they cannot be used in follow-up analysis after birth. Alternative non-invasive monitoring options that have shown some positive results are newborn saliva and urine samples, which may be used for longitudinal follow-ups.29 Also, different methodologies, biomarkers, and the limited number of participants in each study make results unreliable when extrapolating to the general population of preterm babies.

An accurate assessment of oxidative damage and the possibility of targeted treatments might improve neonatal care.27 Currently, none of these biomarkers are used in clinical practice. However, further researches on the field might help to overcome the technical and economic barriers and enable their routine use. As a future perspective, in the next few years, diagnostic strategies directed to the identification of the risk of oxidative stress-related pathologies might be developed, and guidelines for their prevention and treatment might be updated, reducing the morbidity and mortality of premature newborns.

The authors of the present study conclude that the analysis of oxidative stress and antioxidant levels in cord blood of premature newborns might be useful in assessing the diagnosis and prognosis of some clinically relevant pathologies. Oxidative stress and antioxidant activity are involved in the pathophysiology of the development of several neonatal
diseases, and their consequences are associated with increased short- and long-term morbidity, impaired neurodevelopment, and increased mortality. More information and research in the area are needed to impact these clinical outcomes.

Conflicts of interest
The authors declare no conflicts of interest.

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