Physiopathological Mechanism and Assessment of Fetal Asphyxia

Panagiotis Tsikouras¹, Anastasia Bothou², Zacharoula Koukouli¹, Bachar Manav¹, Constantinos Bouschanetzis¹, Dorelia Deuteraiou¹, Xanthi Anthoulaki¹, Anna Chalkidou¹, George Iatrakis³, Stefanos Zervoudis⁴, George Galazios¹

¹Department of Obstetrics and Gynaecology, Democritus University of Thrace, Komotini, Greece
²Research on Female Reproduction, Democritus University of Thrace, Komotini, Greece
³TEI Technical University of Athens, Athens, Greece
⁴Rea Hospital, Athens, Greece

Email: ptsikour@med.duth.gr

Abstract

Treatment and outcome of childbirth depend on the acidobasic balance of the fetal blood related to the oxygen and carbon dioxide level. Hypoxemia could lead to asphyxia that is why fetal monitoring and biochemical parameters assessment are mandatory. Although there are compensatory mechanisms that temporarily protect the fetus, there are also other factors that interfere with the oxygenation of the fetus and determine the development of the fetus and the newborn. Actually, the level of the oxygen, the carbon dioxide, the acidobasic balance and the pH are the cornerstones of the well-being of the fetus.

Keywords

Hypoxia, Asphyxia, Fetal pH

1. Introduction

Successful treatment and outcome of childbirth depend mainly on the assessment of the good status of the fetus. This could be achieved by using modern biochemical and biophysical methods of monitoring. With these methods we try to determine directly or indirectly the degree of oxygenation of the fetus [1].

2. Perinatal Asphyxia

2.1. Terminology

Oxygen partial pressure means the amount of oxygen dissolved in the plasma which is about 2% of the total quantity of oxygen contained in the blood. By determining the oxygen in the blood, taken from the fetal head at childbirth, the
magnitude of the partial oxygen pressure is calculated. 

Oxygen Saturation is the percentage of oxygen-associated to hemoglobin in the fetus and represents 60% - 87%.

Blood oxygen content refers to oxygen dissolved in the plasma bounded to hemoglobin. By this way, a determined provision of oxygen in the tissues is achieved.

Blood oxygen capacity is the maximum blood capacity to carry oxygen, when hemoglobin is fully saturated.

Volume of oxygen transferred to the tissues based on the quantity of the oxygen content in the blood and on the cardiac function, particularly on the volume of the pulse.

The O2 transfer from the atmosphere begins from the maternal lungs and ends up to fetal tissues. This O2 transfer from the maternal to fetal circulation is done by passive diffusion, given that partial pressure of O2 in fetuses is significantly lower than maternal. Basically, the fetus lives and develops in an environment that is hypoxic regarding the maternal environment. However, despite the normal low partial pressure of O2 (20 - 30 mm Hg), fetal tissues have adequate amount of O2 to cover their needs. This is accomplished through the increased transfer ability of O2 in fetal blood and is due to increased fetal hemoglobin levels [2].

If the gas interchange is disturbed or the O2 supply is suspended, the fetal blood O2 reserves act as safeguards. Therefore, in the same time some more mechanisms that reduce further intensive procedures are activated as well and the fetal oxygenation is protected. Such mechanisms are the cardiac supply changes, the increase of blood capacity in O2, reduced movements and fetal development and generally the modification of all fetal functions. When the O2 supply to the fetus is significantly reduced, the anaerobic metabolism is activated, resulting in increased lactic acid production and decrease in fetal arterial blood O2 saturation. The base excess follows and the fetus develops metabolic acidosis that reduces the affinity of fetal Hb to O2 and causes further SpO2 reduction. The severe and prolonged decrease of O2 supply to brain has direct correlation with brain perfusion [3].

In term fetuses normal blood flow in brain is estimated to be 100 ml/min/100gr of tissue. When this tempo is reduced to levels lower than 50 ml/min/100gr of tissue, then the protein synthesis in brain is significantly affected, whereas in levels less than 15 - 20 ml/min/100gr of tissue, the ATP synthesis is disturbed, resulting in damages in neurons’ electric activity. If this low blood flow persists for a long time or is reduced further to levels less than 10 ml/min/100gr of tissue, then the brain damage becomes permanent.

2.2. Physiopathology—Perinatal Asphyxia

During perinatal period, asphyxia is strictly defined as the combination of oxygen deprivation and acidosis with a decline of organ functions.

Asphyxia is the result of events originally associated with hypoxemia and then
with hypoxia. Generally, the term hypoxia means the significant reduce of O₂ in inhaling air and consequently in tissues. In hypoxia, the O₂ supply is inadequate for the tissues’ oxygen requirements. As a result, the aerobic metabolism is completed by the anaerobic in order to maintain the energy balance. If hypoxia is prolonged, the anaerobic metabolism produces lactic acid and the hydrogen ions are increased. It is understood that oxygen deficiency is accompanied by acidosis and as a result the energy balance cannot be maintained anymore. Organ function reduction follows, with the risk of permanent tissue damages. This situation is characterized as asphyxia [4].

Hypoxemia is expressed by reduced oxygen blood pressure and oxygen content and hypoxia with reduced oxygen supply to tissues. Fetal distress-stress and acidosis can be diagnosed with a relevant accuracy by laboratory methods. However, sometimes the distinction of the above mentioned conditions is not easy. Consequently, they are usually evaluated as a single entity. The earliest indication of fetal distress is the intrapartum decrease of fetal heart rate. Other classic clinical indications are colored amniotic fluid and fetal tachycardia [5].

The grade of fetal distress-hypoxia-acidosis development depends on the gravity of the etiological factor and the fetal condition. The term fetal distress concerns primarily the fetus, because it can develop in fetuses with poor counter-balancing mechanisms after the effect of natural procedures, like normal labor. However, frequent labor pain that causes cardiotocography deteriorations does not lead to fetal distress-acidosis in fetuses with sufficient counter-balancing mechanisms. Therefore, situations that cause severe hypoxia, followed by acidosis, result in cardiotocography late decelerations and in decreased pH levels in blood sampling from the fetal scalp [6] [7] [8].

For many years, perinatal asphyxia has been diagnosed on the basis of Apgar Score clinical criteria. Although a serious hypoxic fetus is expected to have low Apgar Score after delivery, asphyxia is not the only cause. Other possible causes that lead to asphyxia are prematurity, infection, trauma and congenital abnormalities of the fetus [5].

The diagnosis of perinatal asphyxia requires the presence of the following characteristics: low Apgar Score, metabolic acidosis and neonatal complications. However, the common use of the term perinatal asphyxia should be avoided. Obviously, the incidence, the description and the time of acidosis should be determined in order to avoid confusion in the use of this term.

Neonatal encephalopathy is thought to be the result of perinatal asphyxia. For the diagnosis of intrauterine asphyxia, a combination of the following factors: metabolic acidosis, low Apgar Score and neonatal complications are necessary [9].

The placenta is like a gas exchange device for the fetus. Oxygen diffuses from maternal to the fetal blood through the placenta and is transferred to the fetus through the umbilical vein.

Un-oxygenated blood with metabolic waste and carbon dioxide goes back to the placenta through the two umbilical arteries. Therefore, the umbilical vein blood expresses the placental acid-base condition, while the blood of the umbili-
cal arteries reflects the acid-base balance of the fetus.

Fetal oxygen supply is determined by the following factors: 1) the level and type of hemoglobin; 2) the blood content in O₂ (pO₂ and SpO₂); and 3) the blood flow [10].

3. Fetus Oxygenation Status Evaluation

3.1. Pathological Oxygenation of the Fetus

During pregnancy and childbirth there exist some factors to lead to reduction in oxygen supply to the fetus:

1) Uterus: General conditions, such as hypotension, ischemia or compression of the mother’s aorta, may cause in a reduction in blood flow to the uterus.

2) Maternal causes: Heart’s diseases that cause pulse volume reduction and hypotension may cause a reduction in blood flow to the uterus. In addition, hemoglobinopathies, respiratory diseases (e.g. asthma), the immune system diseases (e.g. systemic lupus erythematosus) and poorly regulated gestational diabetes mellitus may be responsible for reduced blood flow in the uterus [11].

3) Umbilical cord: The prolapse, compression and wrapping of the umbilical cord around the fetus can create conditions of reduced blood flow to it, resulting in its under-oxygenation.

4) Placenta: Placental deficiency e.g. in prolonged gestation, results in limiting the functional surface of the placenta and in the significant reduction of gas exchange and the supply of the fetus to oxygen. Although normal fetal pO₂ (20 - 25 mm Hg) is lower than that of an adult, the necessity of extra tissue oxygenation in some cases is covered by the increase of cardiac work and the increase of blood flow in systematic circulation [10]. The contribution of increased ability of oxygen transfer of fetal blood is also important. The necessity of extra oxygenation causes acidosis of glucose (main fetal energy reserve) to CO₂ and H₂O and additionally the production of 38 mol ATP/glucose molecule. The CO₂ is diffused easily through placenta and eliminated off maternal organism. Conditions that either cause decrease in greater blood quantities adduction to embryo-placental unit or decrease in the ability of fetal tissues’ extra oxygenation, result in hypoxia-metabolic acidosis establishment with characteristic fetal reactions dependent mainly on fetal state [12].

5) The periodic delay of the contractions may cause occasional decrease in the utero-placental blood flow, resulting in decreased CO₂ elimination and in respiratory acidosis. The decrease in blood flow that lasts more than usually, as it happens in cases of tetanic uterus contractions, may affect unfavorably the procedure of aerobic metabolism even in healthy fetuses. In such cases, the mechanism of anaerobic glycolysis, with the production and accumulation of pyruvate acid and lactic acid, is activated and metabolic acidosis is developed. Additionally, in cases of hypoxia, lactate dehydrogenase, that stimulates the conversion of pyruvate into lactic acid, is activated [13].

6) Fetal causes: Some infections like toxoplasma or cytomegalovirus could decrease the functional surface of the placenta and induce the sub-oxygenation of
the fetus. Anemia caused by hemolysis in Rh-sensitized pregnant women or by embryo-fetal transfusion in monozygotic twins, reduces the fetal oxygenation [14][15].

3.2. Assessment of Fetal Oxygenation

The last 40 years, the assessment of the fetus status is achieved with the cardiotocographic monitoring [16]. Urgent endometrial conditions regarding the oxygenation and mainly regarding the oxygenation of the fetus are provided by the biochemical analysis of the blood, obtained by puncture of the umbilical cord or by sampling from the skin of the fetus’s head or by oxymetry.

4. Conclusion

The time of onset, duration and intensity of under-oxygenation play a key-role in fetal development and neonatal neurological status. Although there are compensatory mechanisms that temporarily protect the fetus, there are also other factors that interfere with the oxygenation of the fetus and determine the development of the fetus and the newborn. Actually, the level of the oxygen, the carbon dioxide, the acidobasic balance and the pH are the cornerstones of the well-being of the fetus.

References


**Submit or recommend next manuscript to SCIRP and we will provide best service for you:**

Accepting pre-submission inquiries through Email, Facebook, LinkedIn, Twitter, etc.
A wide selection of journals (inclusive of 9 subjects, more than 200 journals)
Providing 24-hour high-quality service
User-friendly online submission system
Fair and swift peer-review system
Efficient typesetting and proofreading procedure
Display of the result of downloads and visits, as well as the number of cited articles
Maximum dissemination of your research work

Submit your manuscript at: [http://papersubmission.scirp.org/](http://papersubmission.scirp.org/)
Or contact ojog@scirp.org