Possible treatment of fetal distress without caesarean delivery

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Abstract

Aims: Hypoxic fetal compromise is cured by early caesarean delivery at present, while the surgery causes maternal morbidity by uterine scar. Vaginal delivery of normal fetus after hypoxia treatment in utero will cure the fetus reducing maternal morbidity.

Methods: Although various methods were proposed, maternal lateral posture should not be overlooked.

Results and Conclusion: Although there will be many topics, most will be tried in the future, while fibrin deposit was diagnosed by high gray-level histogram width (GLHW) in fetal growth restriction, which is treated by fibrin solution with maternal heparin treatment.

Introduction

As fetal arterial blood oxygen pressure (PaO₂) was half of adult, and umbilical arterial PO₂ was 50 or lower mmHg [1], fetus tends to be hypoxic, due to the oxygen supply from maternal arterial blood through placental villi. Fetal hypoxia damaged central neuronal cells [2]. Placenta villi prepares active transfer function of nutritive materials for fetal growth, e.g. glucose concentration of fetal blood is higher than maternal one, while oxygen and carbon dioxide gasses are transferred by passive transfer function. Therefore, the active transfer function is damaged forming fetal growth restriction (FGR), preserving simple passive gas transfer [3], therefore fetal hypoxia appears in FGR after the reduction of estimated fetal weight, and fetal heart rate (FHR) acceleration disappears against fetal movement in early stage of hypoxia, where FHR variability is preserved [4]. As the variability developed by the reaction of fetal brain to minor fetal movements, severe hypoxia appears some weeks after the loss of acceleration, when the fetal heart rate (FHR) shows the loss of variability [8] and severe FHR changes appear after the loss of acceleration [4]. Although cases of severe fetal asphyxia received caesarean delivery, the outcome was ominous, developing severe neonatal asphyxia or death, while FGR cases of normal acceleration was born with normal vaginal delivery [4]. The hypoxic fetal brain damage should be prevented by early caesarean delivery, the outcome was ominous, developing severe neonatal asphyxia or death, while FGR cases of normal acceleration was born with normal vaginal delivery [4].

Methods and results

Maternal oxygen inhalation

O₂ gas inhalation for 2 L per min was usually performed in cases of fetal distress, where umbilical arterial PO₂ elevated, but insufficient to disappear fetal hypoxia [1].

Treatment of fibrin deposit in placental intervillous space

Utsu & others [6] found high echogenicity and high gray level histogram width (GLHW) tissue characterization in a FGR case, who had positive cardiolypin antigen, in the second trimester, detecting fibrin deposit in intervillous space by high ultrasound echogenicity and high GLHW, treated with 5,000 U heparin every day in 17 to 31 weeks of pregnancy, where estimated fetal weight increased to normal level, no hypoxic FHR sign appeared, and normal neonate was achieved, instead the fetus died in previous pregnancy of the same mother [6]. Therefore, a FGR case would be treated with fibrin solving therapy, if fibrin deposit is diagnosed by fetal growth restriction (FGR), high ultrasound echogenicity and high GLHW tissue characterization. The fibrin deposit would be treated even in the loss of FHR acceleration in FGR, to prevent severe hypoxia, where the recovery of acceleration should be confirmed. If not, C-delivery will be indicated. The FHR in the loss of FHR variability is such severe fetal brain damage as the anencephalic fetus, so that C-delivery should be performed before the loss of variability in fetal monitoring [9] (Figure 1).

Disappearance of late decelerations after the change to lateral posture from supine one

Since late deceleration (LD) appear by the compression of iliac arteries with contracted pregnant uterus, stopping placental circulation, the LD disappears after changing to lateral posture from supine one [12, 13]. Thus, maternal posture should be changed to lateral one, when there is LD in the FHR. Another cause of LD must be studied, if the posture change is ineffective. Supine hypotension develops pathologic deceleration, also lateral posture should be tried.

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c. Congenital diseases treated by stem cells and genetic procedures

Congenital diseases develop fetal hypoxia where stem cell therapy will treat tissue defect and neurological damage.

d. Treatment of genetic diseases

Genetic diseases will be treated by chromosomal engineering [14] and gene editing [15] in the future, where associated hypoxia will be removed.

e. Preterm labor treatment

Possible developing of preterm labor contraction is as follows; A positive feedback loop of labor composed of uterine contraction - the nerve from the uterus to brain – hypothalamus – hypophysis - oxytocin secretion and uterine contraction develops the oscillation of labor contractions [18]. Thus, preterm labor is suppressed, if a feedback component is rejected. However, no fetal hypoxia was discussed in preterm labor.

Results

Fetal body weight increased to normal level, no hypoxic FHR sign was recorded, and normal newborn was achieved after fibrin deposit solution therapy with heparin administration to the mother in a case of FGR, which associated placental fibrin deposit, while the fetus, who received no fibrin deposit solution, died in previous pregnancy of the same mother [4].

Maternal posture change was effective to improve variable deceleration reducing umbilical cord compression, treating maternal supine hypotension and preventing FHR deceleration caused by the compression of umbilical cord. Late decelerations, developed by the compression of iliac arteries by contracted pregnant uterus at supine posture (Poseiro effect), promptly disappeared after maternal taking lateral posture [12]. FHR deceleration developed by supine hypotension caused by the pressure of inferior vena cava dissapeared by maternal lateral posture too, along with the recovery of blood pressure. The donor twin damage in the TRAP sequence disappeared after the coagulation of nourishing acardiac anomaly with high intensity focused ultrasound [5].

Future fetal distress treatments

a. Pharmaceutical treatment

The application of antiglutamate [7], free radical scavenger and others have been reported to cure fetal distress, while their definite effect has not been reported to remove fetal hypoxic changes.

b. Treatment of hypoxia due to uterine artery constriction in preeclampsia

The constriction of uterine artery, radial and spiral arteries reduce maternal arterial blood supply to the placenta resulting fetal hypoxia, then finally placental infarction, and fetal death (Figure 2) due to the sympathicotonic preeclampsia [11]. Therefore, anti-sympathetic treatment will increase placental maternal blood supply, to prevent fetal hypoxia in the future.

Figure 1. Microscopic fibrin deposit in intervillous space of a FGR placenta. H-E, 200X (Utas M).

Figure 2. Placental infarction in a preeclampsia complicated by fetal death. (Maeda 1965)
Discussion

Caesarean delivery rate performed by obstetric indication was about 8% and by fetal hypoxia was 3% of all births in 1981 Japan, namely, actual C-delivery number in one million births will be about 30,000 cases in a year Japan. The effort to reduce C-delivery under the diagnosis of NRFS (fetal distress) will be useful to lower the maternal morbidity.

Since fetal hypoxia is primarily caused by decreased placental villi oxygen transfer from maternal blood, maternal arterial blood flow reducing inter-villous fibrin deposit should be treated. Also, the increase of maternal blood flow in the placenta removing sympathicotonic constriction of uterine artery will improve fetal oxygenation in preeclampsia. Maternal hypoxia or hypotension caused by maternal disorders, massive hemorrhage and supine posture will be improved by maternal treatment or posture change.

Secondly, various fetal disorders, congenital diseases and fetal cardiac failure will develop fetal hypotension reducing fetal placental circulation and oxygen intake. Therefore, the treatment of fetal disorders will improve fetal hypoxia. Noninvasive high-intensity focused ultrasound ablated nourishing vessel of a TRAP sequence would improve the donor twin [5]. Various fetal diseases will be treated improving fetal disorders and hypoxia. Such genetic disorders as trisomies would be treated by chromosomal technology [12] and gene editing [13] improving fetal hypoxia in fetal disorders in the future.

In preterm labor, periventricular echo density (PVE), which is hyperechogenicity around the ventricle, preceded neonatal PVL and CP, when the PVE lasted until preterm birth, but no brain damage was noted in the full term delivery [19]. Therefore, normal infant is expected without hypoxia if preterm labor is suppressed by tocolysis until full term delivery.

Conclusion

Although the treatment of fetal hypoxia usually depends on C-delivery at present, obstetricians would endeavor to find such nonsurgical therapy as the solution of fibrin deposit in placental intervillous space, or antisympathetic therapy in preeclampsia. Various maternal, fetal or genetic diseases will be treated in the future to achieve normal baby. Preterm labor will also be reduced preventing CP.

References