Early Blood Glucose Level Post-Admission Correlates with the Outcomes and Oxidative Stress in Neonatal Hypoxic-Ischemic Encephalopathy

Inn-Chi Lee 1,2,*, Jiann-Jou Yang 2,3 and Ying-Ming Liou 4,5

1 Division of Pediatric Neurology, Department of Pediatrics, Chung Shan Medical University Hospital, Taichung 40201, Taiwan
2 Institute of Medicine, School of Medicine, Chung Shan Medical University, Taichung 40201, Taiwan; jiannjou@csmu.edu.tw
3 Genetics Laboratory and Department of Biomedical Sciences, Chung Shan Medical University, Taichung 40201, Taiwan
4 Department of Life Sciences, National Chung-Hsing University, Taichung 40227, Taiwan; ymlion@dragon.nchu.edu.tw
5 The iEGG and Animal Biotechnology Center and Rong Hsing Research Center for Translational Medicine, National Chung Hsing University, Taichung 40227, Taiwan
* Correspondence: y610@mercury.csmu.edu.tw

Abstract: The antioxidant defense system is involved in the pathogenesis of neonatal hypoxic-ischemic encephalopathy (HIE). To analyze the relationship between first serum blood glucose levels and outcomes in neonatal HIE, seventy-four patients were divided, based on the first glucose level, into group 1 (>0 mg/dL and <60 mg/dL, n = 11), group 2 (≥60 mg/dL and <150 mg/dL, n = 49), and group 3 (≥150 mg/dL, n = 14). Abnormal glucose levels had poor outcomes among three groups in terms of the clinical stage (p = 0.001), brain parenchymal lesion (p = 0.004), and neurodevelopmental outcomes (p = 0.029). Hearing impairment was more common in group 3 than in group 1 (p = 0.062) and group 2 (p = 0.010). The MRI findings of group 3 exhibited more thalamus and basal ganglion lesions than those of group 1 (p = 0.012). The glucose level was significantly correlated with clinical staging (p < 0.001), parenchymal brain lesions (p = 0.044), hearing impairment (p = 0.003), and neurodevelopmental outcomes (p = 0.005) by Pearson’s test. The first blood glucose level in neonatal HIE is an important biomarker for clinical staging, MRI findings, as well as hearing and neurodevelopment outcomes. Hyperglycemic patients had a higher odds ratio for thalamus, basal ganglia, and brain stem lesions than hypoglycemic patients with white matter and focal ischemic injury. Hyperglycemia can be due to prolonged or intermittent hypoxia and can be associated with poor outcomes.

Keywords: newborns; hypoxic-ischemic encephalopathy; biomarker; thalamus; basal ganglion; glucose; hearing; MRI; outcomes; oxidative stress

1. Introduction

Birth asphyxia is a physiological derangement seen in newborn infants due to a prolonged or profound mismatch between oxygen demand and oxygen delivery [1–4]. It can cause mild to severe neurodevelopmental disabilities. Moderate to severe asphyxia can cause irreversible cerebral cell damage, neonatal seizure, and death, leading to a syndrome of hypoxic-ischemic encephalopathy (HIE) that has multi-organ involvement. Rescue hypothermia has been proven effective and has few adverse effects on newborns with HIE [5–7], and used to reduce neurological injury; nevertheless, a 45–55% risk of death or moderate–severe disability remains in treated infants [5,6,8]. Rescued hypothermia therapy has brought pressure on clinicians to make an early and accurate assessment of neonatal HIE and predict the severity of encephalopathy that will ensue [9]. Although hypothermia...
therapy is proven effective in moderate and severe neonatal HIE [5,10,11], it has not been proven to be beneficial for mild HIE [12–14]. Adjunctive tools or biomarkers for the optimal assessment of neonatal HIE are needed for early diagnosis and timely treatment.

There are two distinctive mechanisms involved in HIE. The first is hypoxic injury. The brain is susceptible to hypoxia, particularly in regions, such as the hippocampus, basal ganglion, thalamus, and brain stem [15,16]. This is called selective neuronal necrosis and status marmoratus of the basal ganglia and thalamus [17,18]. The second is ischemic change. This change occurs due to ischemic injury caused by hypotension or focal infarct, including the anterior, middle and posterior cerebral arteries, as well as their branches [19,20]. The pathogenic mechanisms underlying neonatal HIE can be categorized into three phases. The first phase involves primary energy failure due to the hypoxic-ischemic injury, the secondary phase is a consequence of reoxygenation and reperfusion, and the third phase wherein the hypoxic-ischemic injury can worsen and the inflammation can turn into a subacute and chronic condition [21–25]. The antioxidant defense system is involved in the pathogenesis of neonatal HIE, particularly in the aforementioned second and third phases [26–28]. During the second phase, the activity of the antioxidant defense system is exhausted due to oxidative stress, leading to further damage, including lipid peroxidation, protein denaturation, enzyme inactivation, and DNA damage [29–31]. Glucose concentration can affect the oxidant–antioxidant balance system in the second and third phases, and impair the antioxidant defense system. Thus, glucose imbalance, including hyperglycemia or hypoglycemia, is presumed to play an important role in neonatal HIE and is also a potential diagnostic and prognostic biomarker.

The normal glucose concentration in the blood of newborn infants is between 2.5 mmol/L (45 mg/dL) and 7.0 mmol/L (126 mg/dL). Most newborns have a blood glucose concentration in the middle of the normal range, approximately 3.5 mmol/L (63 mg/dL) to 5 mmol/L (90 mg/dL) [32]. Hypoxia can cause enhanced or unchanged glucose levels and even decreased or increased concentrations of blood glucose, serum insulin, and plasma glucagon [33–37]. These changes occurred since hypoxia was found to stimulate insulin secretion from newborn rats but was inhibited in juvenile rats [38]. The fasting blood glucose concentration remained unchanged in response to acute hypoxia (hours) [33], but increased after 3 days of hypoxia [39,40].

A retrospective cohort study of neonates with encephalopathy showed that hypoglycemia was more often associated with watershed hypoxic-ischemic brain injury than basal ganglia hypoxic-ischemic injury [41], which has been supported in several studies [42,43]. In a prospective cohort analysis of neonates with HIE treated with hypothermia, hypoglycemia in neonatal HIE was associated with an increased odds ratio in a watershed or focal–multifocal brain injury [42]. Hypoglycemia has effects on brain injury, resulting from a hypoxia-ischemia mechanism and is associated with worsening corticospinal tract injury, leading to declined motor and cognitive outcomes at 1 year of age [43]. Regarding hyperglycemia, several retrospective studies [44,45] have reported that hyperglycemia and glucose variability are associated with an increased risk of death and major disability. In hypothermia-treated neonates with HIE, early hypoglycemia or hyperglycemia was noted to cause hearing impairment [46]. In a study of 214 treated neonates with HIE, hyperglycemia was associated with death or unfavorable outcomes at 18 months [44]. Hypoglycemia and hyperglycemia after birth are significant factors that correlate with complications; however, there are also notable differences between the outcomes of glucose abnormalities.

Early diagnosis and rapid treatment are critical for the long-term prognosis of neonatal encephalopathies. Identifying the glucose level to correlate with clinical staging and outcomes can help clinicians begin early treatment. In this study, the first glucose level after the first admission was obtained to correlate with clinical staging, hearing outcomes, magnetic resonance imaging (MRI) findings, and neurodevelopmental outcomes for early diagnosis of neonatal HIE.
2. Patients and Methods

2.1. Patients

We retrospectively reviewed the patient charts of neonates diagnosed with HIE based on a clinical history of fetal distress, metabolic acidosis, or positive-pressure ventilation immediately after birth, at Chung Shan Medical University Hospital from 2015 to 2020. The clinical stages of HIE were classified as Sarnat stage I (mild), II (moderate), and III (severe) [5,6]. Blood glucose levels were measured at the time of admission.

Further examinations for HIE, including head ultrasound (HUS), MRI, automated electrocardiography (aEEG), continuous neonatal conventional EEG monitoring, hearing testing (automated auditory brainstem response (aABR)), and auditory brainstem response (ABR) testing, were performed before discharge. For stage I patients, HUS was performed at birth and at 1, 3, 7, and 14 days of age. An MRI was performed if the clinical condition was suspected to be a brain lesion among these patients.

An experienced pediatric neurologist and neonatologist consultant divided the patients into group 1, classified as Sarnat stage I (mild) HIE, and group 2, classified as stage II (moderate) and III (severe). The differences in blood biomarker levels were compared between the two groups (Figure 1).

![Flow chart of the study procedure demonstrated in neonatal hypoxic-ischemic encephalopathy cases and their first glucose level after admission. MRI, magnetic resonance imaging.](image-url)

Figure 1. Flow chart of the study procedure demonstrated in neonatal hypoxic-ischemic encephalopathy cases and their first glucose level after admission. MRI, magnetic resonance imaging.
The HIE patients were then divided into three groups according to the first glucose level after the first admission after birth: group 1 (>0 mg/dL and <60 mg/dL, n = 11), group 2 (≥60 mg/dL and <150 mg/dL, n = 49), and group 3 (≥150 mg/dL, n = 14). The analysis of the three-group outcomes were based on short-term (clinical staging, hearing test, and MRI findings) and long-term neurodevelopmental changes at 1 year.

2.2. MRI Classification

The MRI findings were divided into two groups to study the correlation between the biomarkers and MRI changes. The first group showed no brain lesions in the parenchyma, while the second group showed brain lesions in the parenchyma. In the second group, brain MRI was classified into two subgroups based on the location of the lesions: one of the basal ganglia, thalamus, or brain stem (midbrain, pons, and lower brain stem), and the other involved areas other than the basal ganglion, thalamus, and brain stem.

2.3. Hearing Tests before First Discharge

For patients who failed the aABR test twice during universal newborn screening, ABR testing, otoacoustic emissions, and steady-state evoked potentials were performed [47]. The ABR waveforms were analyzed, and the latency of peak V was defined and adjusted by an experienced pediatric neurologist or otolaryngologist. The degree of hearing loss was classified as normal (<25 and ≤35 dB nHL) and abnormal, including mild (>35 and ≤45 dB nHL), moderate (>45 and ≤65 dB nHL), severe (>65 and ≤90 dB nHL), or profound (>90 dB nHL) [48,49].

2.4. Measurement of Neurodevelopmental Outcome at >1 Year of Age

The third edition of the Bayley Scales of Infant and Toddler Development (Bayley-III) was used to evaluate the neurodevelopmental outcomes at >1 year of age. Cognitive and motor subscales were used to interpret the neurodevelopmental outcomes. The Bayley-III scores were defined as follows: normal if both cognitive and motor subscale scores were ≥85, and abnormal if one of the cognitive and motor subscale scores was <85 [17,18].

2.5. Statistical Analysis

The independent t-test was performed to compare the means of two independent groups for the significant differences between groups, and the categorical variables were analyzed using the chi-square test. The Fisher’s exact test was performed when the sample size was small. The odds ratio (OR) was calculated by dividing the odds of the first group by the odds of the second group. Furthermore, the Mann–Whitney U test was performed if the sample distribution was nonparametric, and the statistical significance was set at a p-value of < 0.05. For correlation analyses, Pearson’s test was performed to measure the strength of the linear association between the two variables. All statistical tests were performed using SPSS (version 14.0; SPSS Institute, Chicago, IL, USA).

Ethical approval for the study was provided by Chung Shan Medical University Hospital’s internal review board (IRB #: CS14003) and was performed in accordance with the relevant guidelines.

2.6. Informed Consent Statement

Since this is a retrospective study, informed consent was not required.

3. Results

3.1. Demographic Data in Newborns with HIEs

After excluding 18 patients due to congenital anomalies (n = 7), preterm with a gestational age less than 36 weeks (n = 10), or with a confirmed genetic defect later on (n = 1), 74 patients with HIE were enrolled. Eleven belonged to group 1, 49 belonged to group 2, and 14 belonged to group 3 (Figure 1). Among the three groups, factors, including birth weight, sex, age, and inborn or outborn method of delivery (cesarean section or
vaginal delivery), were not significant (Table 1). The 1 min and 5 min Apgar scores were not significantly different in the three groups (Table 1). The initial blood glucose level is shown in Figure 2.

Table 1. Seventy-four neonatal hypoxic-ischemic encephalopathy cases were classified into three groups, according to the first serum glucose level taken before 6 h of birth.

| First Glucose Level after Admission | >0 mg/dL and <60 mg/dL (Group 1, n = 11) | ≥60 mg/dL and <150 mg/dL (Group 2, n = 49) | ≥150 mg/dL (Group 3, n = 14) | p between Group 1 and Group 2 | p between Group 2 and Group 3 | p between Group 1 and Group 3 |
|-------------------------------------|------------------------------------------|------------------------------------------|----------------------------|--------------------------------|----------------------------|--------------------------------|
| Mean ± SD (Range)                   | 35.1 ± 19.3 (9.0–58.0)                    | 104.4 ± 23.4 (60.0–145.0)               | 222.0 ± 62.4 (152.0–332.0) | t(58) = −1.13, p = 0.263 | t(58) = 1.77, p = 0.082 | t(61) = −0.628, p = 0.642 |
| Gestational age (weeks)             | 38.3 ± 2.0                                | 38.7 ± 1.3                              | 38.6 ± 1.2                  | t(61) = −1.4, p = 0.164 | t(23) = 1.82, p = 0.084 |
| Birth weight (gm)                   | 3251.2 ± 729.1                            | 2972.2 ± 368.6                          | 2798.3 ± 446.2             | t(23) = −0.47, p = 0.623 |
| Inborn                              | Male (54.5%)                              | 6 (54.5%)                               | 27 (55.1%)                 | χ²(1, n = 60) = 0.01, p = 0.973 |
|                                    | Female (45.5%)                            | 22 (44.9%)                              | 6 (42.9%)                  | χ²(1, n = 63) = 0.02, p = 0.892 |
|                                     | Inborn (54.5%)                            | 19 (38.8%)                              | 5 (35.7%)                  | χ²(1, n = 63) = 0.17, p = 0.683 |
|                                     | Outborn Method of delivery                | 30 (61.2%)                              | 9 (64.3%)                  | χ²(1, n = 25) = 0.24, p = 0.623 |
| Cesarean section                    | 4 (36.4%)                                | 18 (36.7%)                              | 5 (35.7%)                  | χ²(1, n = 60) = 0.01, p = 0.982 |
| Vaginal delivery                    | 7 (63.6%)                                | 31 (63.3%)                              | 9 (64.3%)                  | χ²(1, n = 63) = 0.01, p = 0.944 |
| Apgar score at one minute           | 4.4 ± 2.1                                | 3.7 ± 2.1                               | 3.5 ± 2.9                  | t(58) = 0.74, p = 0.466     |
| Apgar score at five minutes         | 6.3 ± 2.1                                | 5.4 ± 2.4                               | 4.7 ± 2.6                  | t(58) = −0.86, p = 0.397     |

HIE, hypoxic-ischemic encephalopathy and SD, standard deviation.

Figure 2. The initial blood glucose level.
3.2. Clinical Staging and Glucose Level

Among the patients in group 1, 2 (18.2%) had stage I, 7 (63.6%) had stage II, and 2 (18.2%) had stage III HIE. In 49 cases in group 2, 27 (55.1%) had stage I, 14 (28.6%) had stage II, and 8 (16.3%) had stage III HIE disease. Among the 14 cases in groups 3, 1 (7.1%) had stage I, 6 (42.9%) had stage II, and 7 (50.0%) had stage III HIE (Figure 1 and Table 1). The differences were significant among the group with clinical staging ($\chi^2 (4, n = 74) = 16.5$, $p = 0.002$). However, in groups 1 and 3, the differences in clinical staging distribution were not significant ($\chi^2 (2, n = 25) = 2.9$, $p = 0.238$) (Table 2). Glucose levels were significantly correlated with clinical staging ($r (72) = 0.379$, $p < 0.001$).

Table 2. The clinical staging, hearing outcomes, imaging findings, and neurodevelopmental outcomes in the three groups with neonatal hypoxic-ischemic encephalopathy.

| Clinical Staging | Group 1, $n = 11$ | Group 2, $n = 49$ | Group 3, $n = 14$ | $p$ Values Among Group 1, Group 2, and Group 3 | $p$ Values between Group 1 and Group 3 |
|------------------|------------------|------------------|------------------|-----------------------------------------------|--------------------------------------|
| Stage I ($n = 30$) | 2 (18.2%) | 27 (55.1%) | 1 (7.1%) | $\chi^2 (4, n = 74) = 16.5$, $p = 0.002$ | $\chi^2 (2, n = 25) = 2.9$, $p = 0.238$ |
| Stage II ($n = 22$) | 7 (63.6%) | 14 (28.6%) | 6 (42.9%) | | |
| Stage III ($n = 22$) | 2 (18.2%) | 8 (16.3%) | 7 (50.0%) | | |
| Patients without detected lesion in brain parenchyma ($n = 46$) | 4 (36.4%) | 37 (75.5%) | 5 (35.7%) | $\chi^2 (2, n = 74) = 11.0$, $p = 0.004$ | $\chi^2 (1, n = 25) = 0.001$, $p = 0.97$ |
| Patients with detected lesion in brain parenchyma ($n = 28$) | 7 (63.6%) | 12 (24.5%) | 9 (64.3%) | | |
| Abnormal MRI or CT | | | | | |
| Basal ganglion, thalamus, and brain stem ($n = 20$) | 1 (14.3%) | 10 (83.3%) | 9 (100%) | $\chi^2 (2, n = 28) = 12.3$, $p = 0.002$ | $\chi^2 (1, n = 16) = 8.9$, $p = 0.002$ |
| White matter or focal ischemic injury without lesion of basal ganglion, thalamus, and brain stem ($n = 8$) | 6 (85.7%) | 2 (16.7%) | 0 (0%) | | |
| Hearing outcomes | | | | | |
| Patients with hearing impairments ($n = 13$) | 1 (8.1%) | 6 (12.2%) | 6 (42.9%) | $\chi^2 (2, n = 74) = 7.7$, $p = 0.021$ | $\chi^2 (1, n = 25) = 3.5$, $p = 0.062$ |
| Patients without hearing impairments ($n = 61$) | 10 (91.9%) | 43 (87.8%) | 8 (57.1%) | | |
| Neurodevelopmental outcomes at 1 year old | | | | | |
| Unremarkable ($n = 45$) | 5 (45.5%) | 35 (71.4%) | 5 (35.7%) | $\chi^2 (2, n = 74) = 7.11$, $p = 0.029$ | $\chi^2 (1, n = 25) = 0.24$, $p = 0.622$ |
| Abnormal ($n = 29$) | 6 (54.5%) | 14 (28.6%) | 9 (64.3%) | | |

MRI, magnetic resonance imaging; CT, computed tomography. * The findings of the image included 27 stage 1 with brain ultrasounds, and 47 (3 stage I, 30 stage II, and 14 stage III) with brain MRI or CT. Bold fonts indicate significance.

3.3. Correlation of Parenchymal Brain Lesion and Glucose Level

Lesions in brain parenchyma by MRI, computed tomography, or ultrasound were detected in 7 (63.6%) patients in group 1, 12 (24.5%) patients in group 2, and 9 (64.3%) in group 3 (Figure 3). The differences in the brain parenchymal lesions between the 3 groups were significant ($\chi^2 (2, n = 74) = 11.0$, $p = 0.004$). When groups 1 and 3 were compared, the brain parenchymal lesions were not significantly different ($\chi^2 (1, n = 25) = 0.001$, $p = 0.97$) (Table 2). The glucose levels were significantly correlated with the parenchymal brain lesions ($r (72) = 0.238$, $p = 0.044$).
Patients without hearing impairments (n = 61) 10 (91.9%) 43 (87.8%) 5 (45.5%)

Hearing impairment with neonatal HIE were 1 (8.1%) in group 1, 6 (12.2%) in group 2, and 6 (42.9%) in group 3. The hearing impairment across the three groups were significantly different (χ² (2, n = 74) = 7.7, p = 0.021). Hearing impairment was more common in group 3 than in group 1 (χ² (1, n = 25) = 3.5, p = 0.062) and group 2 (χ² (1, n = 63) = 6.6, p = 0.010). Of the 72 patients with HIE, glucose level was significantly correlated with hearing impairment (r(72) = 0.341, p = 0.003).

3.5. Correlation of Neurodevelopmental Outcomes and Glucose Level

The neurodevelopmental outcome at least 1 year of age was correlated with the first glucose level after admission. In group 1, 5 (45.5%) had unremarkable and 6 (54.5%) had abnormal neurodevelopmental outcomes. In group 2, 35 (71.4%) had unremarkable and 14 (28.6%) had abnormal neurodevelopmental outcomes. In group 3, 5 (35.7%) had unremarkable and 9 (64.3%) had abnormal neurodevelopmental outcomes. The neurodevelopmental outcomes in the three groups were significantly different (χ² (2, n = 74) = 7.11, p = 0.029). However, the ratio of abnormal neurodevelopmental outcomes at >1 year of age in groups 1 and 3 was not significant (χ² (1, n = 25) = 0.24, p = 0.622) (Table 2). In the 72 patients with HIE, the glucose levels were significantly correlated with the neurodevelopmental outcomes (r(72) = 0.331, p = 0.005).

3.6. The Differences of Other Blood Biomarkers in the Group 1, Group 2, and Group 3 Patients

We compared groups 1 and 3 and found significant difference in the levels of lactate dehydrogenase (LDH) (3560.4 ± 2851.1 vs. 863.3 ± 510.3; U = 60; p = 0.001); serum aspartate transaminase (SGOT) (504.0 ± 539.0 vs. 113.0 ± 89.9; U = 90; p = 0.019); serum alanine transaminase (SGPT) (162.8 ± 163.2 vs. 33.4 ± 30.3; U = 84; p = 0.009); platelets (164,888.9 ± 58,650.3 vs. 246,714.3 ± 67,271.5 mm³ µL; U = 87; p = 0.007); C-reactive protein (4.3 ± 5.9 vs. 0.04 ± 0.2 mg/L; U = 122; p = 0.025); and creatine kinase (CK) (5080.1 ± 7238.7
vs. 1830.2 ± 2808.8 U/L; U = 106; p = 0.047) (Table 3). The glucose levels were significantly correlated with LDH (r(64) = −0.401, p < 0.001); SGPT (r(62) = −0.354, p = 0.005), SGPT (r(62) = −0.324, p = 0.010); platelet count (r(67) = 0.208, p = 0.086); and CK (r(67) = −0.235, p = 0.066). However, the lactate levels; white blood cell counts; hemoglobin levels, blood urea nitrogen levels; creatinine levels; prothrombin time; activated partial thromboplastin time; and albumin, sodium, potassium, creatine kinase-MB, and troponin levels were not significant (Table 3). The findings highlight that the systemic biomarker levels were higher in group 1 than in group 3.

| Biomarkers | Group 1, n = 11 | Group 2, n = 49 | Group 3, n = 14 | p Values between Group 1 and Group 2 | p Values between Group 2 and Group 3 | p Values between Group 1 and Group 3 |
|------------|----------------|----------------|----------------|---------------------------------|---------------------------------|---------------------------------|
| WBCs       | 23,372.0 ±     | 19,784.2 ±     | 23,688.6 ±     | U = 199, p = 0.935              | U = 214, p = 0.072              | U = 41, p = 0.166               |
| (9100–34,000 mm³ L⁻¹) | 17,485.5      | 7501.3         | 8536.4         |                                 |                                 |                                 |
| Platelet   | 164,888.9 ±    | 238,977.8 ±    | 246,714.3 ±    | U = 87, p = 0.007              | U = 297, p = 0.008             | U = 21, p = 0.176              |
| (84–478 mm³ L⁻¹) | 58,650.3      | 74,946.5       | 67,271.5       |                                 |                                 |                                 |
| Hemoglobin | 17.3 ± 2.6     | 16.8 ± 2.1     | 18.9 ± 12.1    | U = 160, p = 0.318             | U = 256, p = 0.297             | U = 42, p = 0.176              |
| (13.88 ± 1.34 g/dL) |               |                |                |                                 |                                 |                                 |
| SGOT (30–100 U/L) | 504.0 ± 539.0 | 129.8 ± 160.4  | 113.0 ± 89.9   | U = 90, p = 0.019              | U = 276, p = 0.045             | U = 31, p = 0.044              |
| SGPT (6–40 U/L) | 162.8 ± 163.2 | 36.5 ± 58.9    | 33.4 ± 30.3    | U = 84, p = 0.009              | U = 244, p = 0.411             | U = 32, p = 0.046              |
| BUN (3–12 mg/dL) | 11.2 ± 4.1    | 10.7 ± 3.8     | 11.9 ± 5.7     | U = 173, p = 0.691             | U = 264, p = 0.568             | U = 63, p = 0.975              |
| Creatinine | 1.0 ± 0.2      | 0.9 ± 0.2      | 1.1 ± 0.3      | U = 156, p = 0.472             | U = 198, p = 0.858             | U = 54, p = 0.549              |
| (0.03–0.50 mg/dL) |               |                |                |                                 |                                 |                                 |
| Lactate    | 86.9 ± 76.8    | 68.6 ± 46.9    | 92.5 ± 33.7    | U = 213, p = 0.964             | U = 173, p = 0.017             | U = 55, p = 0.403              |
| (4.4 to 14.4 mg/dL) |               |                |                |                                 |                                 |                                 |
| LDH (170–580 U/L) | 3560.4 ± 2851.1 | 875.7 ± 657.8 | 863.3 ± 510.3 | U = 60, p = 0.001             | U = 300, p = 0.009             | U = 22, p = 0.009              |
| PT (13.0 ± 1.43 s) | 19.8 ± 15.8   | 16.4 ± 6.1     | 16.8 ± 3.0     | U = 179, p = 0.726             | U = 235, p = 0.217             | U = 60, p = 0.850              |
| aPTT (42.9 ± 5.80 s) | 62.4 ± 23.7  | 56.0 ± 25.5    | 71.0 ± 23.0    | U = 155, p = 0.352             | U = 173, p = 0.018             | U = 46, p = 0.284              |
| Albumin    | 3.3 ± 0.8      | 3.5 ± 0.4      | 3.7 ± 0.3      | U = 164, p = 0.363             | U = 236, p = 0.294             | U = 44, p = 0.315              |
| (2.5–3.4 g/dL) |               |                |                |                                 |                                 |                                 |
| Na (133–146 mmol/L) | 135.6 ± 1.8  | 136.0 ± 3.6    | 135.1 ± 3.9    | U = 182, p = 0.631             | U = 230, p = 0.127             | U = 46, p = 0.262              |
| K (3.2–5.5 mmol/L) | 3.8 ± 0.6    | 4.1 ± 0.7      | 4.2 ± 0.7      | U = 155, p = 0.269             | U = 294, p = 0.714             | U = 46, p = 0.218              |
| CK (39–308 U/L) | 5080.1 ± 7238.7 | 1986.1 ± 2398.8 | 1830.2 ± 2808.8 | U = 106, p = 0.047 | U = 225, p = 0.040 | U = 29, p = 0.043 |
| CK-MB (0–4.5 ng/mL) | 75.9 ± 70.8  | 68.7 ± 94.5    | 23.7 ± 19.7    | U = 61, p = 0.736             | U = 64, p = 0.084             | U = 11, p = 0.188              |
| Troponin I (0–30 pg/mL) | 915.7 ± 1365.1 | 154.9 ± 420.0 | 97.8 ± 62.3    | U = 77, p = 0.076             | U = 195, p = 0.459             | U = 28, p = 0.236              |
| CRP (1.5–20 mg/L) | 4.3 ± 5.9    | 5.1 ± 19.6     | 0.04 ± 0.2     | U = 122, p = 0.025             | U = 249, p = 0.102             | U = 23, p = 0.002              |

Table 3. Biomarkers exhibited in group 1, group 2, and group 3.

Bold fonts indicate p < 0.05. HIE, hypoxic-ischemic encephalopathy; ST, standard deviation; WBCs, white blood cells; GOT, aspartate transaminase; GPT, alanine transaminase; BUN, blood urea nitrogen; LDH, lactate dehydrogenase; PT, prothrombin time; aPTT, activated partial thromboplastin time; CK, creatine phosphokinase; CK-MB, creatine kinase-Mb; K, potassium; Na, sodium; and CRP, C-reactive protein.

4. Discussion

A significant contribution of this study is the correlation of the first glucose level of neonates with HIE with clinical staging, findings of brain MRI, hearing outcomes, and neurodevelopmental outcomes at 1 year. Hypoglycemia and hyperglycemia are associated
with advanced staging, brain parenchymal lesions, hearing impairment, and abnormal neurodevelopmental outcomes. Hyperglycemia was strongly related to hearing loss and thalamus, basal ganglia, and brain stem lesions; however, hypoglycemia was closely related to white matter lesions. This finding is also compatible with high systemic biomarker levels indicating liver injury (LDH, SGPT, SGPT, and platelets) in the hypoglycemic group. This finding supports the hypothesis that hyperglycemia is caused by prolonged or intermittent hypoxia. In addition, cases with increased blood glucose levels can be more severe than those with hypoglycemia due to the involvement of the thalamus, basal ganglia, and brain stem, especially the 8th nuclei with hearing impairments.

As hypothermia therapy needs to be performed in a timely manner, the use of a simple and convenient method, such as blood glucose measurement, can be useful in the early prediction of the staging of neonatal HIE and the need for the initiation of treatment. Although a combination of other biomarkers, such as lactate and LDH levels, can help predict the severity of HIE, obtaining the glucose level is a rapid and convenient method. This is beneficial for management, as it allows early rescue hypothermia performed 6 h after birth and permits the use of neuroprotective drugs.

Despite similar clinical staging and MRI findings, hearing impairment in patients with hyperglycemia with a first blood glucose level >150 mg/dL is worse than that in patients with hypoglycemia, with a first blood glucose level <60 mg/dL. This finding can be explained by several hypotheses. First, hypoglycemia can further induce fatty acid oxidation and cause ketosis, which can have a protective effect in the brain; hyperglycemia does not have this effect. Second, hypoxia demonstrated significant increases in plasma glucose and insulin [38]; however, intermittent or prolonged hypoxia can increase insulin resistance in genetically obese mice [35] that causes reflex hyperglycemia. Therefore, further studies on the effects of glucose abnormalities to neonatal HIE outcomes are warranted. Third, hypoglycemia can be induced by hyperinsulinemia due to neonatal HIE; however, the compensation mechanism of hyperglycemia in HIE can be exhausted, reflecting the poor condition seen in newborns.

In neonates with encephalopathy, periods of hyperglycemia were common and temporally associated with worse aEEG background scores, reduced sleep–wake cycling, and increased electrographic seizures, including after adjusting for clinical markers of hypoxia-ischemia. Hyperglycemia epochs were also associated with poor aEEG background scores, including after adjusting for hypoxia-ischemia severity. Our data support the hypothesis that the proactive avoidance of hyperglycemia can be a neuroprotective strategy for infants with neonatal encephalopathy [50]. Hypoglycemic or hyperglycemic blood levels can affect MRI findings. In hypoglycemia, the watershed or focal multifocal infarcts were observed on MRI scans [43], whereas hyperglycemia was more associated with the basal ganglia or global injury findings on MRI scans [43]. In 56 neonatal HIE, all of whom died, we studied their neurodevelopmental outcomes and first 24 h glucose level and highlighted 9 patients with first glucose levels over 200 mg/dL.

The findings of the aforementioned studies [28,29] were compatible with our findings that hypoglycemia and hyperglycemia can increase the risk of poor outcomes in neonatal HIE based on MRI findings. However, in our study, we highlighted that hyperglycemia was associated with a high risk of hearing impairment, which is crucial for childhood neurodevelopment. In hypothermia-treated neonates with HIE for 42 babies, 4 (9.5%) had hearing impairments. The development of hearing loss was associated with abnormal blood glucose levels, low Apgar scores, and evidence of multi-organ dysfunction and increased SGPT and SGPT levels [46], which are compatible with our findings. In addition, we also highlighted that the hyperglycemic patients had more thalamic and basal ganglion injuries than those with hypoglycemia before the first 6 h. These findings suggest that hyperglycemia can cause selective neuronal necrosis that causes injury to susceptible brain tissue, including the basal ganglia, thalamus, and brain stem. We hypothesized that the mechanism of neonatal HIE is related to glucose and clinical staging (Figure 4). Hyperglycemia in the reoxygenation and reperfusion stage can lead to further brain injury
due to the consequence of oxidation stress. Hypoglycemia can cause ketogenesis by acting as an alternative cerebral fuel and as antioxidants. This can explain why the hypoglycemia group had better outcomes than the hyperglycemia group in the study. Hyperglycemia caused by insulin resistance can contribute to further brain injury as the consequence of oxidation stress that can be a useful biomarker of poor neurological outcomes and worse neurological consequences [51]. Thus, avoiding hyperglycemia after admission is mandatory in the clinical management of neonatal HIE.

![Diagram of the hypothesized mechanism of hypoglycemia and hyperglycemia in neonatal hypoxic-ischemic encephalopathy](image)

Figure 4. The hypothesized mechanism of hypoglycemia and hyperglycemia in neonatal hypoxic-ischemic encephalopathy.

However, this study has some limitations. We presented a limited number of HIE cases. Our findings can be biased and comprised owing to the fewer cases than needed for reliable results. Therefore, further studies with an increased number of cases are warranted. Furthermore, in the HIE stage I group with favorable outcomes, an aggressive image study is not available from the national insurance agency in Taiwan. However, a series of HUS can support the imaging findings, and a clinical follow-up of up to 1 year can diagnose the patients without significant brain parenchymal lesions.

5. Conclusions

The first blood glucose level is an important biomarker for clinical staging, MRI findings, hearing impairment, and neurodevelopmental outcomes in neonatal HIE. Hyperglycemic patients had higher ORs in the thalamus, basal ganglia, and brain stem lesions than hypoglycemic patients who were often related to white matter and focal ischemic injury. This finding supports the fact that hyperglycemia possibly occurred due to prolonged or intermittent hypoxia and oxidation stress, and led to worse outcomes due to the involvement of the thalamus and basal ganglia. In neonatal HIE, early glucose levels after the first admission can be a rapid and convenient biomarker for a timely diagnosis and early treatment administration.

Author Contributions: I.-C.L. designed the study, performed the data analysis, collected the data, performed statistical analysis, and drafted the manuscript. I.-C.L., J.-J.Y., and Y.-M.L. participated in the data acquisition. I.-C.L. supervised the project. All authors have read and agreed to the published version of the manuscript.

Funding: This work was supported by Chung Shan Medical University and National Chung Hsing University grant NCHU-CSMU-11009 and Taiwan Ministry of Science and Technology. MOST 110-2314-B-040-011.

Institutional Review Board Statement: The studies involving human participants were reviewed and approved by CS2-14003, Chung Shan Medical University Hospital.

Informed Consent Statement: Since this is a retrospective study, informed consent was not required.
**Data Availability Statement:** All of the data is contained within the article.

**Acknowledgments:** The authors thank Yi-Ho Weng and I-Ting Chen for performing the electroencephalography with great patience.

**Conflicts of Interest:** The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; or in the decision to publish the results.

**References**

1. Badurdeen, S.; Roberts, C.; Blank, D.; Miller, S.; Stojanovska, V.; Davis, P.; Hooper, S.; Polglase, G. Haemodynamic Instability and Brain Injury in Neonates Exposed to Hypoxia-Ischaemia. *Brain Sci.* 2019, 9, 49. [CrossRef]
2. Armstrong, K.; Franklin, O.; Sweetman, D.; Molloy, E.J. Cardiovascular dysfunction in infants with neonatal encephalopathy. *Arch. Dis. Child.* 2012, 97, 372–375. [CrossRef]
3. Saugstad, O.D.; Rootwelt, T.; Aalen, O. Resuscitation of asphyxiated newborn infants with room air or oxygen: An international controlled trial: The Resair 2 study. *Pediatrics* 1998, 102, e1. [CrossRef]
4. Deorari, A.K.; Broor, S.; Maitreyi, R.S.; Agarwal, D.; Kumar, H.; Paul, V.K.; Singh, M. Incidence, clinical spectrum, and outcome of intraventricular infections in neonates. *J. Trop. Pediatr.* 2000, 46, 155–159. [CrossRef] [PubMed]
5. Azzopardi, D.V.; Strohm, B.; Edwards, A.D.; Brocklehurst, P.; Gunn, A.J.; Halliday, H.; Juszczak, E.; Levene, M.; Marlow, N.; Porter, E.; et al. Moderate hypothermia to treat perinatal asphyxial encephalopathy. *N. Engl. J. Med.* 2009, 361, 1349–1358. [CrossRef]
6. Edwards, A.D.; Brocklehurst, P.; Gunn, A.J.; Halliday, H.; Juszczak, E.; Levene, M.; Strohm, B.; Thoresen, M.; Whitelaw, A.; Azzopardi, D. Neurological outcomes at 18 months of age after moderate hypothermia for perinatal hypoxic-ischaemic encephalopathy: Synthesis and meta-analysis of trial data. *BMJ* 2010, 340, c363. [CrossRef]
7. Nakamura, T.; Asanuma, H.; Kusuda, S.; Imai, K.; Hosono, S.; Kato, R.; Suzuki, S.; Yokoi, K.; Kubok, M.; Yamada, S.; et al. Multicenter study for brain/body hypothermia for hypoxic-ischemic encephalopathy: Changes in HMGB-1. *Pediatr. Int.* 2017, 59, 1074–1079. [CrossRef]
8. Løvmo, E.; Tyson, J.; Shankaran, S.; McDonald, S.; Ehrenkranz, R.; Fanaroff, A.; Donovan, E.; Goldberg, R.; O’Shea, T.M.; Higgins, R.D.; et al. Elevated temperature after hypoxic-ischemic encephalopathy: Risk factor for adverse outcomes. *Pediatrics* 2008, 122, 491–499. [CrossRef] [PubMed]
9. Murray, D.M. Biomarkers in neonatal hypoxic-ischemic encephalopathy-Review of the literature to date and future directions for research. *Handb. Clin. Neurol.* 2019, 162, 281–293.
10. El-Dib, M.; Parziale, M.P.; Johnson, L.; Benson, C.B.; Grant, P.E.; Robinson, J.; Volpe, J.J.; Inder, T. Encephalopathy in neonates with subgaleal hemorrhage is a key predictor of outcome. *Pediatr. Res.* 2012, 71, 155–162. [CrossRef] [PubMed]
11. Azzopardi, D.V.; Strohm, B.; Edwards, A.D.; Deyt, L.; Halliday, H.L.; Juszczak, E.; Kapellou, O.; Levene, M.; Marlow, N.; Porter, E.; et al. Moderate hypothermia to treat perinatal asphyxial encephalopathy. *N. Engl. J. Med.* 2009, 361, 1349–1358. [CrossRef]
12. Edwards, A.D.; Brocklehurst, P.; Gunn, A.J.; Halliday, H.; Juszczak, E.; Levene, M.; Strohm, B.; Thoresen, M.; Whitelaw, A.; Azzopardi, D. Neurological outcomes at 18 months of age after moderate hypothermia for perinatal hypoxic-ischaemic encephalopathy: Synthesis and meta-analysis of trial data. *BMJ* 2010, 340, c363. [CrossRef]
13. Nakamura, T.; Asanuma, H.; Kusuda, S.; Imai, K.; Hosono, S.; Kato, R.; Suzuki, S.; Yokoi, K.; Kubok, M.; Yamada, S.; et al. Multicenter study for brain/body hypothermia for hypoxic-ischemic encephalopathy: Changes in HMGB-1. *Pediatr. Int.* 2017, 59, 1074–1079. [CrossRef]
14. DuPont, T.L.; Chalak, L.F.; Morriss, M.C.; Burchfield, P.J.; Christie, L.; Sanchez, P.J. Short-term outcomes of neonates with mild hypoxic-ischemic encephalopathy treated with hypothermia. *J. Perinatol.* 2003, 23, 308–315. [CrossRef] [PubMed]
15. Schmidt-Kastner, R. Genomic approach to selective vulnerability of the hippocampus in brain ischemia-hypoxia. *J. Neurosci.* 2008, 28, 402–409. [CrossRef] [PubMed]
16. Schimpf, A.L.; Lacourciere, Y.; Pageaux, G.P.; Morin, A.; Lebel, A.; Ahrens, C.; Elia, S.; Dupuis, P.; Alarcon, R.; Vezina, M.; et al. Moderate hypothermia to treat perinatal asphyxial encephalopathy. *J. Child. Neurol.* 2013, 28, 132–137. [CrossRef]
17. Nakajima, W.; Ishida, A.; Morita, N.; Nakajima, W.; Nakajima, W.; Nakajima, W. Neurobiology of hypoxic-ischemic brain injury. *J. Neurosci.* 2000, 20, 7994–8004. [CrossRef]
18. Nakajima, W.; Ishida, A.; Lange, M.S.; Gabrielson, K.L.; Wilson, M.A.; Martin, L.J.; Blue, M.E.; Johnston, M.V. Apoptosis has a prolonged role in the neurodegeneration after hypoxic ischemia in the newborn rat. *J. Neurosci.* 2000, 20, 7994–8004. [CrossRef]
19. Nakajima, W.; Ishida, A.; Lange, M.S.; Gabrielson, K.L.; Wilson, M.A.; Martin, L.J.; Blue, M.E.; Johnston, M.V. Apoptosis has a prolonged role in the neurodegeneration after hypoxic ischemia in the newborn rat. *J. Neurosci.* 2000, 20, 7994–8004. [CrossRef]
20. Nakajima, W.; Ishida, A.; Lange, M.S.; Gabrielson, K.L.; Wilson, M.A.; Martin, L.J.; Blue, M.E.; Johnston, M.V. Apoptosis has a prolonged role in the neurodegeneration after hypoxic ischemia in the newborn rat. *J. Neurosci.* 2000, 20, 7994–8004. [CrossRef]
21. Nakajima, W.; Ishida, A.; Lange, M.S.; Gabrielson, K.L.; Wilson, M.A.; Martin, L.J.; Blue, M.E.; Johnston, M.V. Apoptosis has a prolonged role in the neurodegeneration after hypoxic ischemia in the newborn rat. *J. Neurosci.* 2000, 20, 7994–8004. [CrossRef]
22. Nakajima, W.; Ishida, A.; Lange, M.S.; Gabrielson, K.L.; Wilson, M.A.; Martin, L.J.; Blue, M.E.; Johnston, M.V. Apoptosis has a prolonged role in the neurodegeneration after hypoxic ischemia in the newborn rat. *J. Neurosci.* 2000, 20, 7994–8004. [CrossRef]
23. Davidson, J.O.; Wassink, G.; van den Heuij, L.G.; Bennet, L.; Gunn, A.J. Therapeutic Hypothermia for Neonatal Hypoxic-Ischemic Encephalopathy—Where to from Here? *Front. Neurol.* 2015, 6, 198. [CrossRef]

24. Vannucci, R.C. Experimental biology of cerebral hypoxia-ischemia: Relation to perinatal brain damage. *Pediatr. Res.* 1990, 27, 317–326. [CrossRef]

25. Jensen, A.; Berger, R. Fetal circulatory responses to oxygen lack. *J. Dev. Physiol.* 1991, 16, 181–207.

26. Kos, K.; Goričar, K.; Solitrovska-Salomon, A.; Dolžan, V.; Rener-Primec, Z. Genetic polymorphisms, Gene-Gene Interactions and Neurologic Sequelae at Two Years Follow-Up in Newborns with Hypoxic-Ischemic Encephalopathy Treated with Hypothermia. *Antioxidants* 2021, 10, 1495. [CrossRef] [PubMed]

27. Esih, K.; Goričar, K.; Dolžan, V.; Rener-Primec, Z. The association between antioxidant enzyme polymorphisms and cerebral palsy after perinatal hypoxic-ischaemic encephalopathy. *Eur. J. Paediatr. Neurol.* 2016, 20, 704–708. [CrossRef] [PubMed]

28. Qin, X.; Cheng, J.; Zhong, Y.; Mahgoub, O.K.; Akter, F.; Fan, Y.; Aldughaim, M.; Xie, Q.; Qin, L.; Gu, L.; et al. Mechanism and Treatment Related to Oxidative Stress in Neonatal Hypoxic-Ischemic Encephalopathy. *Front. Mol. Neurosci.* 2019, 12, 88. [CrossRef]

29. Piwkowska, A.; Rogacka, D.; Audzeyenka, I.; Jankowski, M.; Angielski, S. High glucose concentration affects the oxidant-antioxidant balance in cultured mouse podocytes. *J. Cell. Biochem.* 2011, 112, 1661–1672. [CrossRef]

30. Cascant-Vilaplana, M.M.; Sánchez-Illana, A.; Píñeiro-Ramos, J.D.; Llorens-Salvador, R.; Quintàs, G.; Oger, C.; Galano, J.M.; Vigor, C.; Durand, T.; Kuligowski, J.; et al. Do Levels of Lipid Peroxidation Biomarkers Reflect the Degree of Brain Injury in Newborns? *Antioxid. Redox Signal.* 2021, 35, 1467–1475. [CrossRef]

31. Vlassaks, E.; Nikiforou, M.; Strackx, E.; Hütten, M.; Bekers, O.; Gazzolo, D.; Li Volti, G.; Martinez-Martinez, P.; Kramer, B.W.; Gavilanes, A.W. Acute and chronic immunomodulatory changes in rat liver after fetal and perinatal asphyxia. *J. Dev. Origin. Health Dis.* 2014, 5, 98–108. [CrossRef]

32. Nicholl, R. What is the normal range of blood glucose concentrations in healthy term newborns? *Arch. Dis. Child.* 2003, 88, 238–239. [CrossRef]

33. Brooks, G.A.; Butterfield, G.E.; Wolfe, R.R.; Groves, B.M.; Mazzeo, R.S.; Sutton, J.R.; Wolfel, E.E.; Reeves, J.T. Increased dependence of blood glucose after acclimatization to 4300 m. *J. Appl. Physiol.* 1991, 70, 919–927. [CrossRef] [PubMed]

34. Oltmanns, K.M.; Gehring, H.; Rudolf, S.; Schultes, B.; Rook, S.; Schweiger, U.; Born, J.; Fehm, H.L.; Peters, A. Hypoxia causes glucose intolerance in humans. *Am. J. Respir. Crit. Care Med.* 2004, 169, 1231–1237. [CrossRef] [PubMed]

35. Polotsky, V.Y.; Li, J.; Punjabi, N.M.; Rubin, A.E.; Smith, P.L.; Schwartz, A.R.; O’Donnell, C.P. Intermittent hypoxia increases insulin resistance in genetically obese mice. *J. Physiol.* 2003, 552, 253–264. [CrossRef] [PubMed]

36. Cheng, N.; Cai, W.; Jiang, M.; Wu, S. Effect of hypoxia on blood glucose, hormones, and insulin receptor functions in newborn calves. *Pediatr. Res.* 1997, 41, 852–856. [CrossRef]

37. Chen, X.Q.; Dong, J.; Niu, C.Y.; Fan, J.M.; Du, J.Z. Effects of hypoxia on glucose, insulin, glucagon, and modulation by corticotropin-releasing factor receptor type 1 in the rat. *Endocrinology* 2007, 148, 3271–3278. [CrossRef]

38. Raff, H.; Bruder, E.D.; Jankowski, B.M. The effect of hypoxia on plasma leptin and insulin in newborn and juvenile rats. *Endocrine* 1999, 11, 37–39. [CrossRef]

39. Sawhney, R.C.; Malhotra, A.S.; Singh, T. Gluocregulatory hormones in man at high altitude. *Eur. J. Appl. Physiol. Occup. Physiol.* 1991, 62, 286–291. [CrossRef]

40. Young, P.M.; Sutton, J.R.; Green, H.J.; Reeves, J.T.; Rock, P.B.; Houston, C.S.; Cymerman, A. Operation Everest II: Metabolic and hormonal responses to incremental exercise to exhaustion. *J. Appl. Physiol.* 1992, 73, 2574–2579. [CrossRef]

41. Wang, D.S.; Poskitt, K.; Chau, V.; Miller, S.P.; Roland, E.; Hill, A.; Tam, E.W. Brain injury patterns in hypoglycemia in neonatal encephalopathy. *AJNR Am. J. Neuroradiol.* 2013, 34, 1456–1461. [CrossRef] [PubMed]

42. Basu, S.K.; Ottolini, K.; Govindan, V.; Mashat, S.; Vezina, G.; Wang, Y.; Ridore, M.; Chang, T.; Kaiser, J.R.; Massaro, A.N. Early Glycemic Profile Is Associated with Brain Injury Patterns on Magnetic Resonance Imaging in Hypoxic-Ischemic Encephalopathy. *J. Pediatr.* 2018, 203, 137–143. [CrossRef] [PubMed]

43. Tam, E.W.; Haeusslein, L.A.; Bonifacio, S.L.; Glass, H.C.; Rogers, E.E.; Jeremy, R.J.; Barkovich, A.J.; Ferriero, D.M. Hypoglycemia is associated with increased risk for brain injury and adverse neurodevelopmental outcome in neonates at risk for encephalopathy. *J. Pediatr.* 2012, 161, 88–93. [CrossRef] [PubMed]

44. Basu, S.K.; Kaiser, J.R.; Guffey, D.; Minard, C.G.; Guillet, R.; Gunn, A.J. Hypoglycaemia and hyperglycaemia are associated with unfavourable outcome in infants with hypoxic ischaemic encephalopathy: A post hoc analysis of the CoolCap Study. *Arch. Dis. Child. Fetal Neonatal Ed.* 2016, 101, F149–F155. [CrossRef] [PubMed]

45. Al Shafouri, N.; Narvey, M.; Srinivasan, G.; Vallance, J.; Hansen, G. High glucose variability is associated with poor neurodevelopmental outcomes in neonatal hypoxic ischaemic encephalopathy. *J. Neonatal-Perinat. Med.* 2015, 8, 119–124. [CrossRef]

46. Fitzgerald, M.P.; Reynolds, A.; Garvey, C.M.; Norman, G.; King, M.D.; Hayes, B.C. Hearing impairment and hypoxia ischaemic encephalopathy: Incidence and associated factors. *Eur. J. Paediatr. Neurol.* 2019, 23, 81–86. [CrossRef]

47. Wroblewska-Seniuk, K.E.; Dąbrowski, P.; Szyfter, W.; Mazela, J. Universal newborn hearing screening: Methods and results, obstacles, and benefits. *Pediatr. Res.* 2017, 81, 415–422. [CrossRef]

48. Hsueh, C.Y.; Huang, C.Y.; Yang, C.F.; Chang, C.C.; Lin, W.S.; Cheng, H.L.; Wu, S.L.; Cheng, Y.F.; Niu, D.M. Hearing characteristics of infantile-onset Pompe disease after early enzyme-replacement therapy. *Orphanet J. Rare Dis.* 2021, 16, 348. [CrossRef]
49. Viswanathan, N.; Vidler, M.; Richard, B. Hearing thresholds in newborns with a cleft palate assessed by auditory brain stem response. *Cleft Palate-Craniofac. J.* 2008, 45, 187–192. [CrossRef]

50. Tas, E.; Garibaldi, L.; Muzumdar, R. Glucose Homeostasis in Newborns: An Endocrinology Perspective. *NeoReviews* 2020, 21, e14–e29. [CrossRef]

51. Hoehn, K.L.; Salmon, A.B.; Hohen-Behrens, C.; Turner, N.; Hoy, A.J.; Maghzal, G.J.; Stocker, R.; Van Remmen, H.; Kraegen, E.W.; Cooney, G.J.; et al. Insulin resistance is a cellular antioxidant defense mechanism. *Proc. Natl. Acad. Sci. USA* 2009, 106, 17787–17792. [CrossRef] [PubMed]