Neuropathology of the Guillain-Mollaret Triangle (Dentato-Rubro-Olivary Network) in Sudden Unexplained Perinatal Death and SIDS

Anna Maria Lavezzi*, Melissa Corna, Luigi Matturri and Franco Santoro

Department of Surgical, Reconstructive and Diagnostic Sciences - “Lino Rossi” Research Center for the study and prevention of unexpected perinatal death and SIDS, University of Milan, Italy

Abstract: The present study was undertaken to evaluate the possible alterations of the triangle of Guillain and Mollaret (G-Mt), a neuronal brainstem/cerebellum network (from the dentate nucleus to red nucleus and inferior olivary nucleus) already known for its involvement in the pathogenesis of the palatal myoclonus, in sudden unexplained perinatal and infant death. In 44 cases of perinatal and infant death victims, aged from 26 gestational weeks to 10 months of life, we investigated, besides the histological morphology of the three nuclei, the c-fos and apoptotic expression, as well as the possible effects elicited by maternal cigarette smoking.

A significant increase of lesions (hypoplasia and/or increased c-fos and apoptotic neuronal immunopositivity) of the three nuclei was found in unexplained death victims, compared with age-matched controls. These alterations were related to maternal cigarette smoking habit.

We postulated that the G-Mt is an important network involved in the pathogenesis of a wide spectrum of pathological manifestations and that maternal smoking during pregnancy can exert an adverse influence on this complex, even leading to sudden death in vulnerable periods of perinatal or infant development.

Keywords: Dentate nucleus, Inferior olivary nucleus, Red nucleus, Palatal myoclonus, Maternal cigarette smoking, SIDS, Unexplained perinatal death.

INTRODUCTION

Early studies on palatal myoclonus ascribed the cause of this pathologic manifestation, characterised by rhythmic involuntary contractions of the palatal musculature, to neuronal lesions affecting the dentato-rubro-olivary pathway or Guillain–Mollaret triangle (G-Mt) [1-7]. This is a functional circuit connecting the dentate nucleus (DN) of the cerebellum of one side with the red nucleus (RN) and the inferior olivary nucleus (ION) on the other side, via the superior cerebellar peduncle, the central tegmental tract and the inferior cerebellar peduncle, respectively (Fig. 1).

We previously highlighted the vulnerability of the ION and of the DN to hypoxia, prevalently expressed by increased neuronal apoptosis and c-fos expression, in unexplained perinatal and infant death victims [8]. Since even the RN may play a part in the autonomic responses which occur in hypoxic conditions [9,10], we aimed to evaluate in the present study the possible involvement of the G-Mt in sudden fetal and infant death, like the alterations in nuclei and/or structures of the brainstem and cerebellum checking vital functions, by us observed in these pathologies [11-16].

In particular, we investigated in a large group of perinatal and infant death victims, aged from 26 gestational weeks to 10 months of life, besides the morphological structure of the ION, RN and DN in histological sections of medulla oblon-
toxylin-eosin and Klüver-Barrera stains, and two additional sections were subjected to immunohistochemistry of the apoptosis and c-fos expression. The remaining sections were saved for further investigations.

The examination of the inferior olivary nucleus, the red nucleus, and the dentate nucleus, the target of this study, was performed in all the cases at the same levels of the medulla oblongata (obex), mesencephalon (superior colliculus) and at the midline of either cerebellar hemispheres. These histological sections also allowed to easily analyze further important structures (dorsal motor vagus, tractus solitarius, hypoglossus, ambiguous, pre-Bötzinger and arcuate nuclei in the medulla oblongata; Kölliker-Fuse nucleus and substantia nigra in the mesencephalon; fastigial, globose, emboliform nuclei and cortex layers in the cerebellum).

In 28 cases, 10 fetuses, 4 newborns and 14 infants, after the in-depth examination, no cause of death was found. These cases were defined Sudden Intrauterine Unexplained Death (SIUD), Sudden Neonatal Unexplained Death (SNUD), and Sudden Infant Death Syndrome (SIDS), respectively.

In the remaining 16 cases, 5 fetuses, 3 newborns and 8 infants, a precise cause of death was formulated at autopsy, with no evidence of neuropathological lesions; these constituted the control cases. Specific diagnoses among the fetal and neonatal deaths included: chorioamnionitis (n=2) and congenital heart disease (n=6). Specific diagnoses among the infant deaths were: congenital heart disease (n=4), severe bronchopneumonia (n=3) and pericarditis (n=1).

While taking the medical history, the mother was asked for information about any smoking habit before, during and after pregnancy. Smoking habit was assigned to two categories (smokers vs. nonsmokers). Overall, 22 of the 44 mothers (50%) were smokers of more than 3 cigarettes a day already before the onset of the pregnancy, and 22 were non smokers. Eighteen women in the smoker group were mothers of a victim of sudden unexplained death.

**Immunohistochemical Methods**

In order to analyze the immuno-expression of c-fos, we used specific primary antibody after applying the avidin-biotin-peroxidase technique, in conformity with conventional immunohistochemical procedures. Immunohistochemical visualization of apoptotic cells was obtained by the TUNEL method (TdT-mediated dUTP-biotin nick end labeling). A detailed description of the immunohistochemical procedures that we applied, including the immunopositivity evaluation, is available in our previous works [8,12,14].

Both morphological and immunohistochemical analyses were performed by at least two independent and blinded observers. All the results were compared to evaluate the interobserver reproducibility. In cases of discordance among the investigators, the case was reviewed and discussed until a unanimous result was obtained.

**Statistical Analysis**

Data were analyzed using the SPSS statistical software package (version 11.0; SPSS Inc., Chicago, IL, USA). Fisher’s exact test was applied for comparisons. P values <0.05 were considered statistically significant.

**RESULTS**

**Morphological Examination of the G-Mt**

Histological examination performed on brainstems and cerebella of the 44 victims demonstrated, in most of the cases, particularly in the subjects who died of known cause, a normal morphology, consistently with the gestational age in fetal deaths, of the inferior olivary, dentate and red nuclei.

Nevertheless, developmental cytoarchitectural abnormalities, represented by hypoplasia of the three nuclei (ION, DN and RN), were found in 7 of the 14 SIDS cases (50%), in 2 of the SNUD and in 2 of the 10 SIUD victims (22%).

The diagnosis of DN hypoplasia was expressed when this nucleus showed small-size, simplified structure, with poor convolutions, and decreased neuronal density compared with the DN morphology of age-matched controls.

The hypoplasia of the ION was defined by lack of both medial and dorsal olivary accessory nuclei and reduced undulation of the principal olivary nucleus (Fig. 2).

The hypoplasia of the RN was diagnosed in absence of the large neurons of the caudal magnocellular portion (Fig. 3).
Fig. (2). Inferior olivary nucleus.
A) Normal structure in a control case (female, aged 2 months). Arrows point to accessory subnuclei (dorsal and medial, respectively).
B) Hypoplasia in a case of SIDS (male, aged 4 months) with lack of both the medial and dorsal accessory nuclei.
Klüver-Barrera stain. Magnification: 4x.

Fig. (3). Red nucleus.
A) Normal structure in a control case (male, aged 4 months). Below, higher magnification of the encircled area in the upper image with large neurons of the caudal magnocellular portion.
B) Hypoplasia in a case of SIDS (male, aged 3 months). Below, higher magnification of the encircled area in the upper image with lack of large neurons that are typical of the caudal magnocellular portion.
Klüver-Barrera stain. Magnification: in the upper images 4x; below 10x.

These morphological alterations were not seen in victims of the control group (statistical significance of sudden death group vs. control group: p< 0.01).

Immunohistochemical Examination of the G-Mt

*Immunohistochemistry* - In 10 sudden death victims (5 fetuses, 1 newborn and 4 infants), more than 80% of the
neurons of the three nuclei were intensely c-fos labeled. In the same nuclei of the other cases c-fos expression was negative or scarcely positive (<25% of positive cells) (Fig. 4A).

**Apoptosis Immunohistochemistry**

We observed high percentages (ranging from 70% to 90%) of apoptotic neurons (TUNEL-positive cells with highly condensed chromatin, various nuclear fragmentation stages and apoptotic bodies) in the three analyzed nuclei of 5 sudden perinatal deaths, 7 SIDS victims and two cases in the infant control group, both of whom died of pneumonia (Fig. 4B). Neuronal apoptotic values, ranging from 0 to 20%, were found among the other brainstem and cerebellum nuclei.

High c-fos and apoptotic expressions were present even in hypoplastic DN, RN and ION observed in 8 of the 11 unexplained deaths.

Table 1 summarizes the G-Mt alterations observed in perinatal and infant death victims of the study. Statistical evaluation suggested that sudden unexplained death was significantly associated with G-Mt neuropathological alterations.

In the great part of SIUD, SNUD and SIDS victims with developmental abnormalities of the G-Mt, we also observed morphological alterations of different brainstem and cerebellum structures (precisely, hypoplasia/agenesis of the arcuate nucleus, pre-Bötzinger nucleus, raphé obscurus nucleus, parafacial nucleus and cerebellar cortex delayed maturation). The most frequent association was between G-Mt alterations and hypodevelopment of the medullary arcuate nucleus in SIDS victims, hypoplasia of the pontine parafacial nucleus in sudden fetal deaths.

Finally, we correlated all the findings with the mother’s smoking habit. A significant relation was evident between maternal smoking, sudden unexplained deaths and alterations of the G-Mt. In fact, 12 of the 18 unexplained death victims with smoking mother showed neuropathological findings (hypoplasia and/or increased c-fos and apoptotic expression) of the G-Mt (p<0.01).

**DISCUSSION**

The important role of the G-Mt alterations in the pathogenesis of the palatal myoclonus, a rare movement disorder consisting of rhythmic involuntary jerky movements of the soft palatal musculature, has been outlined by several authors, even if the pathomechanism generating these rhythmic contractions remained unexplained [1-7].

In this study we demonstrated that lesions that disrupt this pathway are not exclusive of the palatal tremor. The results documented in fact significant increase of biological and cytoarchitectural alterations of all the three nuclei of the G-Mt in sudden unexplained fetal and infant death victims, compared with age-matched controls.

In particular, we observed major defects represented by hypoplasia of the DN, ION and RN, above all in SIDS victims, and subtle alterations represented by high neuronal c-fos and apoptotic expression.

### Table 1. Alterations of the G-Mt in Perinatal and Infant Deaths

| Neuronal Pathological Alterations | Sudden Unexplained Death (SIUD + SNUD + SIDS) n = 28 | Control Perinatal and Infant Death n = 16 | p Value |
|-----------------------------------|-------------------------------------------------|----------------------------------------|---------|
| hypoplasia                        | 11                                              | -                                      | 0.035   |
| c-fos positivity*                | 10                                              | -                                      | 0.048   |
| Apoptosis positivity*            | 12                                              | 2                                      | 0.040   |

SIUD = sudden intrauterine unexplained death; SNUD = sudden neonatal unexplained death; SIDS = sudden infant death syndrome.

* = >25% of immunopositive neurons.
These developmental lesions can affect the reactivity and the autonomic compensatory responses of the G-M nuclei to low oxygen conditions. Besides, Tai and Truong [25] demonstrated that cerebral hypoxic insults can also be one of the main causes of myoclonus in animal models, by using Fos protein as a marker of neuronal activity.

The proto-oncogene c-fos in fact is the first gene activated by noxious signals, particularly in presence of hypoxia [17,19]. Therefore, immunohistochemical labeling of the fos-corresponding protein in activated neurons has been recognized as an important component of the neuronal response to brain injuries. Experimental works have demonstrated in particular that exposure to hypoxia markedly increases fos-immunoreactivity within the brainstem and cerebellar deep nuclei [26-28]. Likewise, several Authors have provided evidence, on the basis of TUNEL-staining studies in rodents, that serious DNA fragmentation associated with apoptosis is observable in response to hypoxic injury in the central nervous system neurons, particularly in the brainstem and cerebellum [29,30].

Thus we detain that lesions in the G-Mt may determine a variety of pathological manifestations which, even if quite different, recognize as common etiology the hypoxia. All these pathologies can be related to a defective muscular contraction. Soft palate, and sometimes eye, face and throat muscles are involved in palatal myoclonus [2,4,31]; the upper airway muscles in sudden unexplained death.

The nuclei of the G-Mt, above all the DN, are in fact critical structures for the control of the somatic muscle contraction, including the respiratory muscles [32,33]. The brainstem and cerebellar nuclei damage following low oxygen delivery may contribute to loss of coordination of upper airway muscle activity, thus preventing eupneic breathing and leading to fatal outcome.

It can be reasonably understandable that alterations of the G-M nuclei may give rise to respiratory inhibition after birth, and consequently to sudden neonatal and infant death. Nevertheless, it is difficult to explain how breathing alterations can lead to death during intrauterine life.

We know that periodic fetal breathing regulates the release of tracheal fluid in the lung, and consequently alveolar expansion, to favor lung development [34,35]. Nevertheless, defective intrauterine respiratory movements would not be sufficient to justify fetal loss.

One possibility that we strongly support, is that the G-Mt in fetal life, likewise to other brainstem and cerebellum structures [11-16], participates in the modulation of all the vital functions. Consequently, disruption of the ascending and descending pathways between mesencephalon/cerebellum/medulla can produce severe dysregulation of the autonomic nervous system, triggering the death mechanism before the birth.

Inhalation of smoke is the possible risk factor for hypoxic events that may contribute to G-Mt alterations in perinatal period. In cases of maternal smoking during pregnancy, carbon monoxide, a gaseous combustion product of nicotine, may readily cross the placenta, where it binds to hemoglobin, by passive diffusion. Consequently, carboxyhemoglobin inhibits oxygen release into the fetal tissues causing hypoxia, especially in the most susceptible organs, including the brain. Nicotine can thus give rise to damage in neuronal networks regarded as selectively vulnerable [36-38].

In agreement with these considerations is the observation in this study of a significant relation between maternal smoking and biomorphological alterations of the G-Mt in sudden unexplained perinatal and infant death.

We conclude that the G-Mt constitutes an important network involved in the pathogenesis of a wide spectrum of pathological manifestations. The consequences of developmental alterations of the G-Mt can be minor such as in patal-tremor, less soft in cerebellar atrophy, resulting from olivo-dentate fiber disruption [39], or very serious leading to unexpected death in vulnerable periods of the perinatal life.

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Hypoplasia of the Guillain-Mollaret Triangle

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