Epidemiological Clinical and Profile of Cranio-spinal Dysr aphisms in Madagascar

Bemora Joseph Synèse, Andrianaivo Radotina Tony, Masina Ndalana d’Assise, Ratovondrainy Willy, Rabararijaona Mamiarisoa, and Andriamamonjy Clément

**ABSTRACT**

Cranio-spinal dysraphisms are a set of congenital malformations resulting from a defect in closing the neural tube during embryonic development. The objective of this study was to describe the epidemiological-clinical profile of these malformations in Madagascar. It was a retrospective, descriptive study and metacentric from January 01, 2016 to December 31, 2018; all cases of cranio-spinal dysraphisms seen in two neurosurgery centers in Madagascar. We have collected 32 cases of cranio-spinal dysraphisms. The hospital prevalence was 0.37%. Among these children, the mean age was 8.48 months with an extreme of 1 day to 6 years; there was a feminine predominance (53.12%), with a sex ratio of 0.88. The absence of maternal intake of folic acid, the intake of folic acid outside the recommended periods, the birth order of the children and maternal pathologies during pregnancy were found as etiology. Cranial topography predominated in 56.25% (18 cases) and 68.75% of the children were asymptomatic. An association with hydrocephalus was found in 31.25% of cases. Prevention of known risk factors is essential, antenatal diagnosis is important for early management and improved prognosis. The treatment is surgical.

**Keywords:** Etiology, surgery, Neural Tube Defects, Hydrocephalus.

I. INTRODUCTION

The term cranio-spinal dysraphism corresponds to a set of congenital malformations, resulting from a defect of closure of the neural tube of variable extent, during the fourth week of the development of embryonic life. They are usually divided into cephalic and spinal forms [1]. They are caused by an absence of complete or partial closure of the neural tube between the 22nd and 28th day of gestation [2], [3]. Each year, more than 300,000 children are born with a neural tube defect, 70% of which are in developing countries [4]. The prevalence in developed countries has fallen sharply due to prevention through genetic counseling; the periconceptional intake of folic acid, one of the main predisposing factors found, although its role in the development of cranio-spinal dysraphisms is unknown [5] and the precision of the antenatal diagnosis with legalization of therapeutic abortion [6]. These malformations usually cause significant psychomotor and sphincteric sequelae requiring multidisciplinary management [6]. The objective of this study was to define the epidemiological-clinical profile of these malformations in Madagascar in order to improve their management.

II. METHODOLOGY

We conducted a retrospective, descriptive and multicenter study of children hospitalized and cared for in the Neurosurgery department of the Joseph Ravoahangy Andrianavalona University Hospital Center (CHU-JRA) as well as the Pediatrics department of the Soavina andriana Hospital Center (CENHOSOA) of the 1st January 2016 to December 31, 2018. All children with either cranial and / or spinal dysraphism, with a complete file were studied (clinical information and antecedents well detailed).

The parameters studied were epidemiological data (frequency, prevalence, age, gender, geographical origin and history) and clinical data (discovery of the malformation, topography, appearance of the skin coating, associated malformations and clinical signs). The data that was collected was entered on Microsoft Excel 2019® and analyzed on R® software version 3.5.2 with an IDE (Integrated Development Environment) RStudio® version 1.1.456.
III. RESULTS

During the targeted period, 32 cases of cranio-spinal dysraphisms with a complete file were listed in these two departments. In our series, cranio-spinal dysraphisms represented 0.37% of hospitalizations with a frequency of 10.6 cases per year (6 cases / years for cranial locations and 4.66 cases / years for spinal forms). Among these children, there was a slight female predominance (53.12%) with a ratio of 0.88; the average age was 8.48 months with an extreme of 1 day and 6 years. Depending on the geographical origin, populations living in the highlands had 43.75% craniospinal dysraphism against 34.37% those in coastal areas. Only 56.25% of mothers who had a child with cranio-spinal dysraphism took folic acid during pregnancy. For the birth order of children, it was found that children who were born first are the most affected by the malformation, 43.75% of cases (Fig. 1).

Clinically, all of these malformations were discovered at birth. According to the topography, 17 cases of dysraphisms were cranial, i.e. 53.12% against 46.87% of spinal topography (15 cases), to note that there were no cases of craniolochalisis or rachischisis; and in 8 of the 32 cases, the malformation was ruptured within 48 hours of birth. The location of these various malformations is shown in Table I.

On the evolutionary level, the postoperative complications were dominated by meningitis and SCL leakage with a respective rate of 7.69% and 3.84%. The average length of hospital stay was 12.43 days with extremes of 1 day and 60 days.

IV. DISCUSSION

From an epidemiological point of view, the frequency of cranio-spinal dysraphisms varies from country to country. In our study, this frequency was 10.6 cases per year (6 cases / year for cranial locations and 4.66 cases / year for spinal forms), with a prevalence of 0.37% hospitalization. In Niger, this annual frequency is 14.66 cases [4] for meningoencephaloceles; in Burkina Faso 24.4 cases per year [7] for spinal dysraphisms. The worldwide prevalence of cranio-spinal dysraphisms ranges from 0.3 to 199.4 per 10,000 births. A delay in consultation was observed both in our study and in other African countries [4], [7], in fact the average age at the time of diagnosis was 8.48 months in our study with an extreme of 1 day and 6 years old. This delay could be related to the distance from the neurosurgery centers. According to gender, in our series we noted a slight female predominance found in 53.12% with a sex ratio of 0.88, but statistical analysis had shown that the relationship between gender and cranio-spinal dysraphisms was not significant with a p-value of 0.99. This slight female predominance was known in the literature, it would be due to epigenetic phenomena, the loss of a fairly large proportion of male fetuses with cranio-spinal dysraphism at an early gestational age, differences in susceptibility to environmental factors, … [8], [9]. In our study, 43.75% of children hospitalized for cranio-spinal dysraphism come from the former province of Antananarivo and that 34.37% come from the province and that 21.87% were not specified in the file. The relationship between geographic origins and cranio-spinal dysraphisms was statistically insignificant in our case (p-value at 0.31), however we observe a slight predominance of children from the highlands with cranio-spinal dysraphisms compared to other geographic origins.

This low rate of children from the provinces has also been reported by some authors [10]-[12], which could be explained by the difficulty of accessing care for children from the province. Maternal folic acid deficiency during the periconceptional period is known to be one of the risk factors for cranio-spinal dysraphism. In our case, 43.75% of mothers did not take it during pregnancy. Mothers who did not take folic acid had a high prevalence of cranio-spinal dysraphism; in addition, the difference between folic acid intake and cranio-spinal dysraphisms was statistically significant with a p-value <0.001.

Our result matches the data in the literature [13]-[16]. This folic acid should be taken between 1 month before conception and throughout the first trimester of pregnancy. There is a correlation between the birth order of children and the existence of craniospinal dysraphisms; indeed, the first child is the most at risk of developing this malformation, this possibility has been seen in our study as well as in the literature [6], [18].

Clinically, the diagnosis of cranio-spinal dysraphism should be made prenatal using ultrasound or fetal MRI [19].

### TABLE I: DISTRIBUTION OF CRANIO-SPINAL DYSRAPHISMS ACCORDING TO LOCATION

| Type of malformations | Localisation         | Number (n = 32) | Percent (%) |
|-----------------------|----------------------|-----------------|-------------|
| Cranial dysraphism (17 cases) | Ethmoid-nasal | 2    | 06.25 |
|                       | Frontal              | 2    | 06.25 |
|                       | Parietal             | 5    | 15.62 |
|                       | Occipital-occipital  | 1    | 3.12 |
| Dysraphisme spinal (15 cases) | Occipital-cervical | 1    | 3.12 |
|                       | Cervical             | 1    | 3.12 |
|                       | Cervical-thoracic    | 1    | 3.12 |
|                       | Thoracic             | 2    | 06.25 |
|                       | Thoraco-lumbar       | 1    | 3.12 |
|                       | Lumbar               | 5    | 15.62 |
|                       | Lumbosacral          | 4    | 12.50 |
|                       | Sacred               | 1    | 3.12 |

Associated with this malformation, we found hydrocelephalus (31.25%), an Arnold-Chiari malformation (9.37%), anal imperforation and clubfoot with a respective rate of 3.12%.

Fig. 1. Distribution of children by birth order.
It should be noted that ultrasound is a dependent manipulative technique, it takes some experience to be able to visualize this malformation. In our case, no malformation was diagnosed antenatal. The location of this malformation is variable; for the cranial form, the occipital topography is by far the most frequent [4], [6], [11], found in 18.75% in our series; for spinal forms, the lumbar (15.62%) and lumbosacral (12.50%) localization is the most frequent both in the literature and in our study [6], [7], [10], [21], [23]. For associated malformations, we found 31.25% hydrocephalus; this malformation is variable in the literature, it can be hydrocephalus, Dandy Walker and other polymalformative syndrome [7], [4], [10], [12], [23], [24].

The course of cranio-spinal dysraphism remains variable, the length of hospitalization depends on the course and the existence of a possible complication. The most frequent complication is an infection such as meningitis (7.69%) and SCL leakage (3.84%), found both in our study and in the literature [4], [6], [10], [12], [22], [23].

V. CONCLUSION

Cranio-spinal dysraphisms are among the congenital malformations mainly linked to a deficiency in folic acid. Antenatal diagnosis remains essential, based on the use of ultrasound or a fetal MRI. Faced with this malformation, it is necessary to look for associated malformations, especially hydrocephalus. The course and the prognosis depend on the topography. Complications are frequent; the best management remains prevention by supplementation with folic acid and early surgical treatment of malformations to prevent their rupture.

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Bemora was born on December 20, 1987 in Maroantsetra, he is the second son of a family of three children. He began his medical studies in 2006 after having obtained the baccalauréat with honors. He was admitted to the competitive examination to be a specialist in 2012 and continued his studies in France then became a specialist in Neurosurgery in 2018. He was admitted to the clinical examination in 2019 and currently, he is head of clinic in Neurosurgery at the Faculty of Antananarivo medicine at CHU JRA Madagascar. He has participated in various national and international research and publications.

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