Primary acalvaria: a case report*

Acalvaria primária: relato de caso

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Abstract Acalvaria is a rare congenital malformation of unknown pathogenesis characterized by the absence of the flat bones of the cranial vault, dura mater and associated muscles, while the central nervous system is usually preserved. The most accepted physiopathogenic theory suggests the presence of a postneurulation defect with normal placement the embryonic ectoderm. The present report describes neonatal imaging findings of primary acalvaria.

Keywords: Primary acalvaria; Skull defect; Congenital malformation.

Resumo Acalvaria é uma malformação congênita rara de patogênese desconhecida, na qual os ossos da abóbada craniana, a dura-máter e a musculatura associada estão ausentes, mas o sistema nervoso central costuma estar preservado. A teoria fisiopatogênica mais aceita sugere um defeito pós-neurulação, com disposição normal do ectoderma embrionário. O objetivo deste relato é descrever os achados de imagem neonatais da acalvaria primária.

Unitermos: Acalvaria primária; Defeito do crânio; Malformação congênita.

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INTRODUCTION

Acalvaria is a rare congenital malformation of unknown pathogenesis characterized by the absence of the flat bones of the cranial vault, dura mater and associated muscles, while the central nervous system is usually preserved although in some cases it may be abnormally developed. This abnormality has been described as a fatal condition and is scarcely described in the literature.

The present case report describes imaging findings of primary acalvaria characterized by the absence of the flat bones of the cranial vault.

CASE REPORT

A 15-year-old primigravida at 38 weeks’ gestation was referred to the authors’ institution for specialized prenatal follow-up. Previous ultrasonography performed at the 31st gestational week had identified fetal unilateral hydrocephalus. There was no maternal history of teratogenic medication, recent infection, diabetes mellitus, hypertension or exposure to drugs during the current pregnancy. Cesarean section was indicated. A female neonate was born at term weighting 2815 grams.

At clinical examination, the neonate’s face was apparently normal, with presence of skin on the frontal region. Cerebral hemispheres prominence and a large defect of the cranial vault were the most obvious findings. The abnormality was characterized by the partial absence of the scalp and the flat bones of the cranial vault, with a thin membranous layer overlying the brain tissue from which one could observe leakage of cephalorachidian fluid compatible with the pia mater (Figure 1). Frontal and parietal bones could not be palpated. The

Figure 1. Clinical examination demonstrates partial absence of the scalp and of the flat bones of the cranial vault. Thin membranous layer overlying the cerebral hemispheres from which leakage of liquor compatible with the pia mater can be observed (arrow). Note the presence of areas with less resistance with exposure of the brain tissue.
newborn was referred to the neonatal intensive-care unit where the investigation was pursued.

Immediately after birth, skull radiography demonstrated absence of frontal, parietal and temporal bones, and presence of facial and occipital bones (Figure 2). The findings were confirmed by computed tomography that also demonstrated unilateral ventricular dilatation (Figure 3). Abdominal ultrasonography and echocardiography did not demonstrate any abnormality.

The brain remained covered by the pia mater for five days. As a result of the growth process, at the sixth day of life, the brain tissue started breaking-up the thin membrane, becoming completely exposed at the tenth day of life. The reconstruction progressed with a local infectious process and partial suture dehiscence.

After two months of life, signs of hypoxic ischemic neurological deterioration could already be observed. After four months of life, transfontanel ultrasonography demonstrated the presence of cystic areas in the brain resulting from hypoxic-ischemic vascular lesion (Figure 4). The neonate progressed to death at the fifth month of life because of respiratory and infectious complications.

DISCUSSION

Primary acalvaria is a rare congenital malformation characterized by either complete or partial absence of the flat bones of the cranial vault, dura mater and associated muscles, with normal skull base and facial bones. Usually, the defect is covered by skin. The intracranial contents are generally complete, although some abnormalities may be associated (1–7).

The etiology and pathogenesis of such abnormality still remain unknown. During the embryonic development, after the anterior neuropore closure around the fourth week migration of the membranous neurocranium occurs under the ectoderm. The condition is said secondary in cases where it is a result of amniotic band syndrome, neural tube defects or use of angiotensin-converting enzyme inhibitors during pregnancy. It is named hypocalvaria in the presence of hypoplastic cranial bones (6).

Association with other malformations such as holoprosencephaly, hydrocephalus and micropolygyria, besides other facial, cardiac anomalies, omphalocles, among others, may be observed (1,6). The rare reports found in the literature suggest a predestination for female fetuses with normal karyotype (1), in agreement with the description of the present case.

Presence of skin covering the cranial vault and partial scalp absence were observed. The presence of ectoderm-derived skin and scalp and the absence of cranial vault, dura mater and associated muscle as a result of mesenchymal migration characterize primary acalvaria.

In this case, association with an extensive type of aplasia cutis congenita cannot be ruled out. This disorder is characterized by the absence of a portion of skin; most of times (84%) manifesting as a defect on the scalp. The skull is affected in 14% to 30% of cases. Extensive and deep lesions are extremely rare, with a high mortality rate, and may involve the periosteum, the skull and the dura mater (8).

REFERENCES