

Case Report

A Rare Case Report of Acrania

Arman DM¹, Mukherjee SK², Ekramullah SM⁴, Uddin Z⁴, Rahman M⁵

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Manuscript Preparation- Dr. Sudipta Kumer Mukherjee

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Abstract :

Acrania is a rare congenital anomaly and characterized by partial or complete absence of the calvarium. Although acrania associated with anencephaly is a well recognized entity but isolated acrania is a rare anomaly. Ultrasound allows early diagnosis of this anomaly. The fetus was found to have a completely formed brain, base of the skull and facial structures but lacking a cranium. Authors present a rare case of acrania.

Key words : Acrania, brain, cranium

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Introduction :

Acrania is a rare congenital anomaly characterized by partial or complete absence of the cranium and by existence of abnormal development of the brain tissue¹. The term of acrania and anencephaly are often confused since every case of anencephaly has calvarian bone defects². Approximately 40 cases have been reported in the English literature since the first description by Mannes et al. (1982). The pathogenesis of acrania is unknown, but it is suggested that acrania is a congenital anomaly resulting from failure of the

mesenchyma to migrate under the ectoderm overlying the brain tissue over the cerebral hemispheres³.

The acrania - exencephaly – anencephaly sequence together with spina bifida are the two most common neural tube defects worldwide with a prevalence of 1.86 per 1,000 live births.⁴ The acrania is not actually an isolated neural tube alteration, it belongs to a sequence called acrania exencephaly anencephaly, since the lack of bones that make up the cranial vault will cause a protrusion of the cerebral parenchyma (exencephaly) and with the sudden movements of the

1 Dr. D. M. Arman, Assistant Professor. Paediatric Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh.

2 Dr. Sudipta Kumer Mukherjee, Associate Prof. Paediatric Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh.

3 Prof. Dr. Sheikh Muhammad Ekramullah, Professor and Head, Paediatric Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh.

4 Dr. Zia Uddin, Assistant registrar. Paediatric Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh.

5 Dr. Moshir Rahman, medical officer. Paediatric Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh.

Address of Correspondence: Dr. D. M. Arman, Assistant Professor. Paediatric Neurosurgery, National Institute of Neurosciences and Hospital, Dhaka, Bangladesh. cell + 08801745771780. email- armandmdr@yahoo.com

fetus along with the chemical irritation of the amniotic fluid to the unprotected brain structure causes degeneration and destruction of it causing in the absence of brain mass (anencephaly)^{3,5}. Due to this pattern of progression, anencephaly is considered relatively more common than exencephaly⁶.

It is described as a postneurulation defect that, after the closure of the cranial neuropore, there is an alteration in the migration of the membranous portion of the neurocranium². Among the risk factors for this defect, low intake of foods rich in folates⁷ is important. Access to food enriched with folates, which are public norm in some countries, are the protective factor to avoid malformations⁸, in the same way the genetic origin has been demonstrated as a risk factor for acrania, as is the case of the polymorphism of the methylenetetrahydrofolate reductase found in 2% of Afro-descendants and more than 35% in Chinese and Mexican^{9,10}.

We present a case of acrania in Dhaka, Bangladesh.

Case report :

A 22 years old non-diabetic, non hypertensive, non-alcoholic mother with normal obstetric examination gave birth a male baby with acrania in full term. This baby was her 2nd issue. There is no family history of congenital anomalies. There was no history of intake of teratogenic drugs and other relevant past illness. During ultrasonographic scan a single live fetus with cephalic presentation was found. Ultrasonologist did not mention any congenital anomaly.

Live fetus born through normal vaginal delivery without any complication, having well formed brain without skull bone covering (acrania). Brain tissue was covered with thin membrane. Facial feature were normal. Nasal bone, lips were seen normally formed. New born baby was brought to our hospital after 3 days of delivery on 20.01.2021 from Chouddagram, Cumilla. We found new born baby without cranium, baby was otherwise normal. Dressing with Sofratulle and gauze was done. Death of neonate was occur on the same night at home.



Fig.-1: a,b,c,d: Baby with acrania

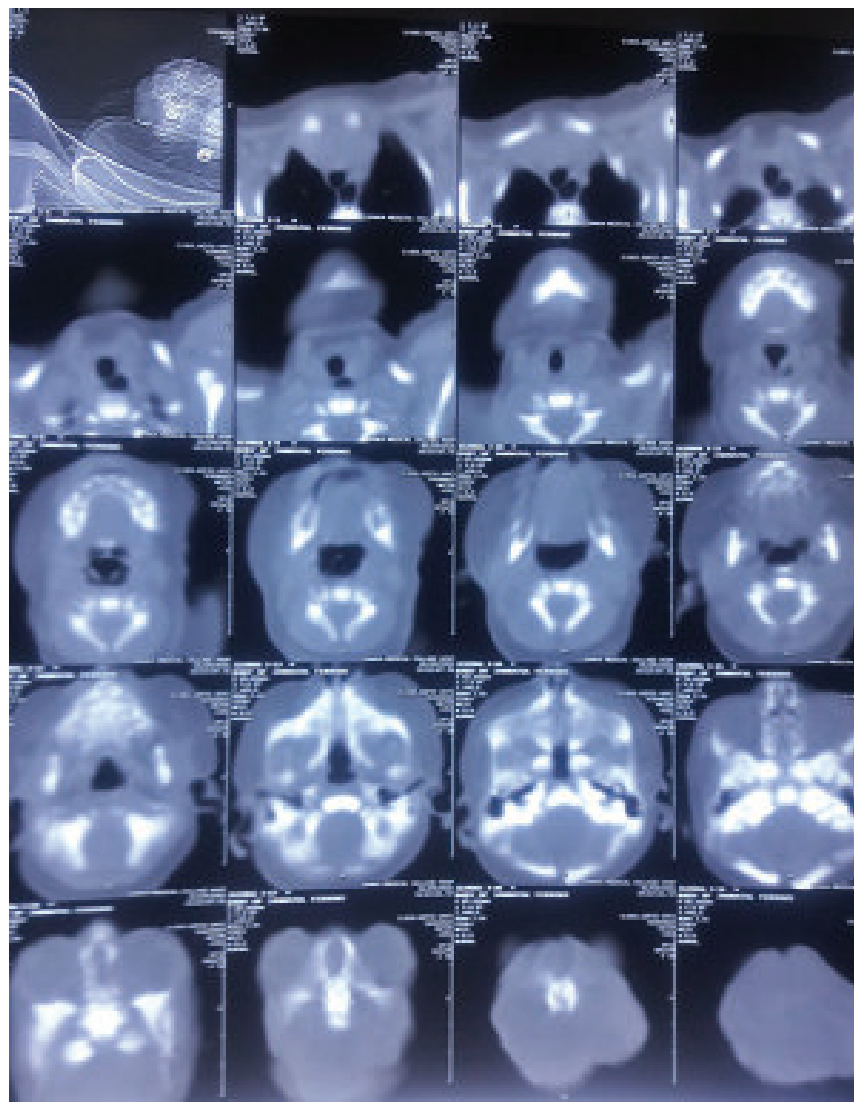


Fig.-2: CT scan of baby with acrania

Discussion:

Acrania is a congenital abnormality characterized by the complete or partial absence of skull bones surrounding the fetal brain with complete, but abnormal development of brain tissue¹¹. Although acrania associated with anencephaly is a well recognized entity with an incidence of about 10:10,000 births, isolated acrania is a rare anomaly. The fetal cranium is not fully calcified before 10–11 weeks; Fetal acrania can be diagnosed from 11 weeks onward¹². At 11–14 weeks gestation, the majority of cranial ossification is in the lateral aspects of the frontal bones and lower parietal bones, and no vault ossification is visible in the midline on a perfect midsagittal image. Hence, misdiagnosis may occur if only midsagittal views of the fetus are obtained. It is important to look

specifically for frontal bone ossification in the axial and coronal planes^{13,14}. For prenatal diagnosis, ultrasound remains the primary modality.¹⁵

Neural tube defects have a prevalence of 1.86 per 1000 live births⁵, among the two most common pathologies are spina bifida and acrania. The latter is not actually an isolated neural tube alteration, it belongs to a sequence called acrania exencephaly anencephaly, since the lack of bones that make up the cranial vault will cause a protrusion of the cerebral parenchyma (exencephaly) and with sudden movements of the fetus and the chemical irritation of the amniotic fluid to the unprotected brain structure causes degeneration and destruction of the brain and causing its absence (anencephaly)³. There is a confusion between what

is acrania and acalvaria, the first is the absence of the scalp and partial or complete cranial vault that inevitably causes anencephaly, while the second is the total or partial absence of the skull bones but with an intact scalp and it will not cause anencephaly¹⁶. Therefore, acrania that is incompatible with life, acalvaria could have a survival expectancy.

It is described as a postneurulation defect that, after the closure of the cranial neuropore, there is an alteration in the migration of the membranous portion of the neurocranium². The fetal neurocranium has two sections, the chondrocranium that forms the base bones and the membranous flat bones that form the cranial vault, acrania is an alteration in said formation and occurs at 4 weeks of gestation when there is a defect in the closure cranial neuropore¹⁷. The diagnosis is made in the 12-week ultrasound where the bones of the cranial vault are not visualized, which are characterized by being a hyperechoic ring surrounding the brain parenchyma¹⁷.

Since the 1960s, the relationship of folate deficiency with neural tube defects was studied, it was determined that folate was essential for the transfer of a carbon unit for the transformation of homocysteine to methionine, DNA methylation and others. Cellular reactions are an essential part for rapid tissue growth and cell replication^{18,19,20}, it is for this reason that many countries are obliged to supply folic acid to all pregnant women. Acrania has a high mortality rate of almost 100%, due to incompatibility with life, it is decided to terminate the pregnancy.

Conclusions :

Acrania is a rare congenital anomaly. Most cases are incompatible with life. Antenatal identification allows the clinician to make appropriate and timely management decisions.

References :

- Mannes EJ, Crelin ES, Hobbins JS, Viscomi GN, Alcebo L. Sonographic demonstration of fetal acrania. *Am J Radiol*;1982. 139: 181-182.
- Harris CP, Townsend JJ, Carey JC. Acalvaria: a unique congenital anomaly. *Am J Med Genet*;1993.46: 694-699.
- Weissman A, Diukman R, Auslender R. Fetal acrania: five new cases and review of the literature. *J Clin Ultrasound*;1997. 25: 5 1 1-5 14.
- Blencowe H, Kancharla V, Moorthie S, Darlison MW, Modell B. Estimates of global and regional prevalence of neural tube defects for 2015: a systematic analysis. *Ann. N.Y. Acad. Sci*;2018.1414: 31– 46.
- Samaniego Haro VJ. Sequence Acrania Exencephaly Anencephaly Report of a Case in the San Vicente De Paul Hospital in Ibarra Ecuador. *Austin Gynecol Case Rep.*;202. 5(1): 1024
- J A. J. Nawale, S. A. Merchant, S. S. R. Koteyar, and P. Masand. "Exencephaly: a rare case diagnosed on antenatal ultrasound: Bombay Hospital Journal;2000. vol. 42, pp. 520–521.
- Elwood JH, Nevin NC. Anencephalus and spina bifida in Belfast (1964–1968). *Ulster Med J.*;1973.42: 213–222.
- Bower C, Antoine HD, Stanley FJ. Neural tube defects in Australia: trends in encephaloceles and other neural tube defects before and after promotion of folic acid supplementation and voluntary food fortification. *Birth Defects Res:A Clin Mol Teratol*; 2009.85: 269–273
- Botto LD, Yang Q. 5, 10-Methylenetetrahydrofolate reductase gene variants and congenital anomalies: a HuGE review. *Am J Epidemiol*;2000. 151: 862– 877.
- Tsang BL, Devine OJ, Cordero AM, Marchetta CM, Mulinare J, Mersereau P, Guo J, Ping Qi Y, Berry R J, Jorge Rosenthal, Krista S Crider, Hamner HC. Assessing the association between the Methylenetetrahydrofolate Reductase (MTHFR) 677C>T polymorphism and blood folate concentrations: a systematic review and meta-analysis of trials and observational studies. *Am. J. Clin. Nutr*;2015. 101: 1286–1294.
- a H, Sezik M, Özkaya O, Aydin AR. Fetal acrania at term: *Perinatal Journal*.; 2004.12(2):96–98.
- Cincore V, Ninios AP, Pavlik J, Dong C. Prenatal Diagnosis of Acrania with amniotic band syndrome. *Obstetrics and Gynecology*; 2003.102(5):1176– 1178. [PubMed]
- Fong KW, Toi A, Salem S, Hornberger LK, Chitayat D, Keating SJ, McAuliffe F, Johnson JA. Detection of Fetal Structural abnormalities with US during early Pregnancy. *RadioGraphics*; 2004.24:157–174.[PubMed]
- Johnson SP, Sebire NJ, Snijders RJM, Tunkel S, Nicolaides KH. Ultrasound screening for anencephaly at 10–14 weeks of gestation. *Ultrasound: Obstet Gynecol*; 1997.9:14–16. [PubMed]
- Radhey Sankhala, Richa Jhuria, Sangeeta Saxena, Dharmraj Meena, Devendra Khatana. A Rare Case Report –Acrania : 2016. volume : 5 Issue : 2
- Bianca S, Ingegnesi C, Auditore S, Reale A, Galasso MG, Bartoloni G. Arancio A, Ettore G. prenatal and postnatal findings of acrania. *Arch Gynecol Obstet.*;2005.271: 256-258
- Peer D, Moroder W, Delucca A. Prenatal diagnosis of the pentalogy of Cantrell combined with exencephaly and amniotic band syndrome. *Ultraschall Med*; 1993.14: 94-95.
- Lanska DJ. Historical aspects of the major neurological vitamin deficiency disorders: the water-soluble B vitamins. *Handb. Clin. Neurol.*;2010. 95: 445– 476.
- Czeizel AE. Nutritional supplementation and prevention of congenital abnormalities. *Curr Opin: Obstet Gynecol*; 1995. 7: 88–94.
- McPartlin J, Halligan A, Scott JM, Darling M, Weir DG. Accelerated folate breakdown in pregnancy: *Lancet*; 1993. 341: 148–149