Coexistent neck enteric cyst and intraspinal neurenteric cyst: embryopathogenetic implications

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Introduction. Enteric cysts are very rare conditions, occurring mainly in the posterior mediastinum and posterior neck. Their pathomorphology corresponds with that of intraspinal neurenteric cysts. Both formations are derivatives of the posterior foregut. However, their embryopathogenesis has not been elucidated satisfactorily as yet. For those associated with vertebral anomalies, the split notochord theory has been widely accepted. However, this is be hardly conceivable for cases free of these anomalies.

Case report. Here, a patient with concurrent separated enteric and neurenteric cysts and cervical spine dysmorphism is presented.

Discussion. The review of the relevant literature revealed sporadic analogical cases in which a transvertebral communication between the two cysts was present or absent. The latter was associated with a minimal abnormality of the vertebral body.

Conclusion. The authors suggest that normal vertebrae may be formed in patients with enteric cysts, which would make the notochord-split theory plausible also for those free of spinal malformations.

Key words: embryogenesis, cysts, notochord, enteric cyst, neurenteric cyst

INTRODUCTION

Foregut anomalies are sporadic dysembryonal formations. Because of variable anatomical position and histomorphology, they were labeled with different, often confusing terms. In 2009, Sharma et al. proposed a new taxonomy and classification based on a unifying concept of suggested embryopathogenesis, categorizing them as duplication cysts, bronchopulmonary foregut malformations, bronchogenic cysts and foregut and enteric cysts (ECs) (ref.1). The latter (also termed enterogenous cyst, endodermal cyst, gastroenterogenous cyst, gastrocytoma, intestinoma, and archenteric cyst, etc.) are considered to be derivatives of the dorsal portion of the foregut that suggests an origin in the upper part of the gastrointestinal tract (including pharynx, esophagus, stomach, duodenum, pancreas, liver and gall bladder). ECs thus comprise two distinct smooth muscle layers and epithelial lining corresponding morphologically with that of the dorsal foregut itself or the visceria originating from it. ECs may be incorporated in various GIT organs or isolated in different locations with a prevalence in the posterior mediastinum and rarely intraabdominaly or in the posterior neck2-11. Neurenteric cyst (NEC) is considered to be a variant of EC. It differs from it in occasional presence of ependymal or glial elements, and intraspinal or very rarely, intracranial position. The embryopathogenesis of these cysts has not been elucidated satisfactorily as yet and various theories have been postulated1,3. Frequent (40-70%) coexistence with spinal anomalies and persistent attachment to the vertebral column suggests that the split notochord theory is at work in the formation of these anomalies12,13. However, this theory fails in cases devoid of split notochord syndrome consisting of vertebral malformation associated with central nervous system and gastrointestinal tract anomalies.

Here we describe simultaneous occurrence of a posterior neck enteral cyst and an intraspinal neurenteral cyst. We believe that our case along with those found in the relevant literature will contribute to the embryopathogenesis of these malformations.

CASE REPORT

A 29-year old woman presented with a history of headaches and neck stiffness. With the exception of an inconspicuous sinistrocoliosis, the ENT examination was within normal limits. MRI of the neck showed congenital synostosis of vertebrae C2-5, a hemivertebra Th1 and a deformation of vertebrae C6 and 7, which correspond to the MRI findings linked to Klippel-Feil syndrome. Furthermore, two cysts were found (Fig. 1a) with the larger one located in the right posterior neck extending
down to the Th3 level (Fig. 1b). The longitudinal and transversal diameters were 8 by 4 cm. Another small intraspinal cyst lacking any evident connection with the former was also present. The MRI image of the intraspinal cyst met the MRI criteria of neurenteric cysts, moreover in the presence of Klippel-Feil syndrome. The cervical cyst was removed using an external approach. It was found to overlie but not to communicate with the thyroid trachea and esophagus. There were neither fibrous connections nor communication between vertebrae and the cyst and its complete removal as well as postoperative course was uneventful. Because of the absence of any cord compression symptoms, the intraspinal cyst was left without any neurosurgical intervention. Three years later, the patient is doing well, with no clinical signs of recurrence. Histopathologic analysis of the surgical specimen (Fig. 2) showed a cyst with the wall composed of two layers of a smooth muscle. The cyst was covered with gastric mucosa with continuous transition into a flat non-keratinizing squamous epithelium. No cartilage was found.

**DISCUSSION**

Association between ECs and spinal abnormalities, assuming a common mechanism involved in their formation was noticed by some early authors. Later, it was conclusively proven by Bently and Smith who postulated the split notochord theory. This suggests primary split of the notochord as the starting formal event in the development of enteric cysts associated with spinal malformations. The split occurs in a three week embryo and is composed of ectodermal, mesodermal and endodermal
layers. The latter gives rise to the notochordal plate. This is temporarily in contact with dorsal ectoderm and escalates later from the endoderm to form solid notochord that is the primordium of the future axial skeleton. In this pathological scenario, herniation of the endodermal diverticulum through the pre-existing notochord gap and its contact with the ectoderm occur. These events, interfering with normal development of vertebrae, spinal cord and relevant viscera, result in anomalies called posterior enteric remnants. Their eventual nature determined by the extent of the herniation may be subdivided into fistulae, sinuses and posterior enteric diverticula and cysts. The latter two formations develop when solely an intermediate part of the endodermal diverticulum persists. The remaining part may become atrophic, leaving a fibrous band connecting the cyst to the anterior aspect of the vertebral body. The more or less complete persistent gap in the dorsal embryonic tissues results in associated various spinal malformations (butterfly vertebra, anterior or posterior spina bifida, Klippel-Feil syndrome, diastematomyelia, meningocele etc.).

Other mechanisms for development of an endodermal diverticulum may be abnormal escalation of the notochord from the dorsal foregut and persistent neurenteric canal. However, the latter theory is weak. The canal that represents a patent endo-ectodermal passage separating the notochord into two lateral parts is situated in the most caudal aspect of the spine where enteric cysts occur very rarely. This drawback eliminates the Bremer’s theory of accessory neurenteric canal that occurs in a more rostral position.

These theories are based on the formation of endodermal cells lined diverticulum attached to or separating the notochord. Therefore, they are considered to be applicable only for cysts associated with vertebral malformations. For cases free of them, other more or less acceptable explanations were sought.

Shredhar, Piramoon, Holcomb and Dorsey described posterior mediastinal enteric cysts expanding intraspinally through a hole in distorted vertebral body. In our patient, the cysts were associated with spinal anomaly but lacked any apparent transvertebral communication. A similar finding was described by Guillery in a baby who died at the age of 3 months from a congenital heart defect. At the autopsy, however, neither epithelial nor fibrous transvertebral connection between the two cysts was found. The author thus postulated early presence of a cyst between the lateral vertebral anlagen. It later becomes snared into two parts by normal fusion of vertebral masses, with minimal or – as we assume - even no vertebral body malformation and no connection tract left behind. We believe that this hypothesis would explain those cases of isolated enteric cervico-thoracic and intraabdominal cysts published by Andrew, Mahore and Kim, respectively, in which, because of absent spinal dysraphism, the notochord-split theory could seem implausible.

Despite the similar pathomorphology of a dorsal foregut formation, it is not quite clear whether ECs and NECs share identical embryopathogenesis. This hypothesis is supported by the simultaneous presence of both types of cysts. They – as it was in our case and in that described by Guillery – may be entirely separated or communicate through a transvertebral defect. In addition, the occurrence of two unrelated dysembryogenetic events in the same patient is hardly conceivable.

**CONCLUSION**

The presented case of coexistent foregut intraspinal and cervical cysts lacking transvertebral communication suggests that they may develop without spinal malformation. This makes the split notochord theory plausible also for cases with absent spinal dysraphism.

**ABBREVIATIONS**

EC, Enteric cyst; HE, Hematoxylin eosin; NEC, Neurenteric cyst; T2 WI, T2 weighted image.

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**REFERENCES**