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AJNR Am J Neuroradiol published online 29 October 2014
http://www.ajnr.org/content/early/2014/10/29/ajnr.A4153
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ABSTRACT

BACKGROUND AND PURPOSE: Calcifications of the pineal, habenula, choroid plexus, and dura are often physiologic. In the modern CT era with thin-section images and multiplanar reformats, intracranial calcifications have become more conspicuous. We aimed to discover the CT prevalence of pineal region, choroid plexus, and dural calcifications in the first decade of life.

MATERIALS AND METHODS: Five hundred head CTs from different patients (age range, 0–9 years) encountered during a consecutive 6-month period at a single academic children’s hospital were reviewed retrospectively after excluding examinations with artifacts and pineal region masses/hemorrhage. All studies were performed on a 320-detector CT, with 0.5-mm collimation and a 512 × 512 matrix. Five-millimeter reformatted axial, sagittal, and coronal images were analyzed for location and extent of intracranial calcifications.

RESULTS: The mean age was 3.5 ± 5.7 years (range, 0–9 years). There were 285 males (57%) and 215 females (43%). Pineal calcifications were present in 5% (n = 25; age range, 3.2–8.9 years; median, 7 years). Habenular calcifications were found in 10% (n = 50; age range, 2.8–8.8 years; median, 7 years). Twelve percent (n = 58) had choroid plexus calcifications, (age range, 0.1–8.8 years). Dural calcifications were rare, present in 1% (n = 6; age range, 2.9–8.7 years).

CONCLUSIONS: Physiologic intracranial calcifications may be found in the first decade, principally in children older than 5 years. Most epithalamic calcifications are habenular. Pineal and habenular calcifications were never present in children younger than 3 and 2 years, respectively. Choroid plexus calcifications may be present in the very young. Dural calcifications are rare.
320-detector CT scanner (Aquilion ONE; Toshiba Medical Systems, Tokyo, Japan) with 0.5-mm collimation and a 512 × 512 matrix. The epithalamus, choroid plexus, and dura were analyzed for calcifications on 5-mm axial, coronal, and sagittal reformatted images in each case. Epithalamic calcifications included either pineal, habenular, or both. Epithalamic calcifications were also subdivided into pineal or habenular on the basis of their distinct location; focal pineal parenchymal or pineal wall calcifications were classified as “pineal,” while pineal adjacent calcifications were considered “habenular.” The latter are generally localizable anterolateral to the pineal gland along the medial thalamic margins.

Calcification positivity was defined as hyperattenuating material respective to gray matter. In addition to group analysis, we evaluated patient subsets on the basis of age. We recorded the total number of patients in each year of life and calculated the percentages of pineal, habenular, choroid plexus, and dural calcifications in patients of each age (range, 0–9 years). Pineal calcifications were classified as punctate and/or single or larger and/or numerous (>1). All encountered pineal cysts were recorded and manually measured by calipers on a PACS in the maximal diameter; mean pineal cyst sizes were then calculated. Dural and choroid plexus calcifications were grouped by location. Choroid plexus calcifications were classified as either punctate and/or few or large and/or multiple. The potential relationship between choroid plexus and pineal calcifications was explored.

Images were retrospectively reviewed in a blinded manner by 2 fellowship-trained neuroradiologists with American Board of Radiology subspecialty certificates in neuroradiology (M.T.W. and A.F.C.). Both readers had >3 years of clinical experience after board certification. A consensus reading was reached in all cases. Each patient’s electronic medical record was researched for presenting symptomatology, demographic information, and pertinent physical examination findings.

Five hundred patients met the inclusion criteria, with a mean age of 3.5 ± 5.7 years (median, 2.5 years; range, 0.0–9 years). The percentages of patients with intracranial calcifications in each age group are shown in Fig 1. There were 285 males (57%) and 215 females (43%). The most common clinical indications for the examinations were trauma (n = 209), seizures (n = 66), shunt (n = 57), and headache (n = 41). None of the patients had undergone prior therapeutic radiation, and no histories of metabolic disorders, endocrinopathies, or renal diseases could cause calcium-phosphate imbalance and premature or excessive intracranial calcification. One patient had a history of tuberous sclerosis but had no intracranial calcification. None of the patients had documented histories of TORCH infection or neurofibromatosis type 1, entities known to be associated with calcifications in or around the choroid plexus.3

**Data Collection and Statistics**

Data were stored and analyzed in a spreadsheet (Excel 2011; Microsoft, Redmond, Washington) and in SPSS (Version 20.0; IBM, Armonk, New York). Discrete variables were evaluated by using the χ² or Fisher exact test, as appropriate. Continuous variables were analyzed for significance by using the Kruskal-Wallis test with a nonparametric posttest. Correlations were made by using the Pearson product-moment correlation coefficient. P values <.05 were considered significant.

**RESULTS**

Age and intracranial calcification presence were positively correlated (P < .0001) (Fig 1). Most (97%) intracranial calcifications were found in patients older than 5 years. There was no significant difference in sex (P = .41) or race (P = .28) of patients with and without calcifications. A summary of location-specific intracranial calcifications is provided in the Table.

**Epithalamus**

Epithalamic (pineal and/or habenular) calcifications were present in 13% (63/500). Pineal and habenular calcifications were positively correlated (P < .0001). Pineal calcifications were present in 5% (n = 25; age range, 3.2–8.9 years; median, 7 years). One percent (6/500) of patients were younger than 6 years of age (Fig 2). Pineal calcifications were single and/or punctate in 71% (17/24) and larger and/or more numerous in 29% (7/24; range, 7–9 years).

Pineal cysts were found in 3% (n = 15; age range, 0.7–7.5 years; median, 2.6 years). These ranged from 4 to 15 mm (median, 5 mm). Four of these 15 patients had epithalamic calcifications involving the habenula (n = 4) and/or pineal gland (n = 1).

Habenular calcifications were found in 10% (n = 50; age range, 2.8–8.8 years; median, 7 years; Fig 2).

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**FIG 1.** Chart depicting the percentage of patients with calcifications (x-axis) in various regions by year of life (y-axis).
Twelve percent of subjects (n = 58) had choroid plexus calcifications (age range, 0.1–8.8 years; median, 6 years; Fig 3). All choroid plexus calcifications were localized to the lateral ventricular glo-
mus. Most choroid plexus calcifications were punctate and/or few (n = 54). The 4 remaining patients with large and/or numerous choroid plexus calcifications were 8 (n = 3) and 3 (n = 1) years of age. There was a significant association between patients with choroid plexus and pineal calcifications (P = .003).

Dural calcifications were rare, present in only 1% (n = 6; age range, 2.9–8.7 years; median, 6 years; Fig 4). The tentorium was the most common dural site to calcify (n = 4) followed by the falx cerebri (n = 2).

**DISCUSSION**

**Epithalamus**
The epithalamus comprises the pineal gland and habenular com-
plex. We chose to examine the prevalence of epithalamic, pineal,
and habenular calcifications separately for several reasons. First, we think pineal and habenular calcifications can be accurately distinguished in most circumstances with modern CT technology and thin-section images. Second, we suspect that in the past, many habenular calcifications were misclassified as being pineal. Because these structures are functionally unique, we think that they deserve to be analyzed separately.

**Pineal Gland**
The pineal gland is a conical organ situated in the quadrigeminal cistern between the posterior and habenular commissures. Pi-
nocyes, glial cells, neurons, and perivascular phagocytes constitute the bulk of its internal structure. After a period of initial growth, the size of the pineal gland has been shown to remain stable beyond the first 1–2 years of life. Knowledge of normal pineal gland size/volume is relevant in the evaluation of pineal region pathology. Pineal calcifications that would be considered benign in isolation are suspicious for neoplasia if there is concurrent glandular enlargement.

Why does the pineal gland calcify, and where is the calcium localized? Calcified concretions called "corpora arenacea" or "brain sand" found within the pineal gland primarily comprise calcium and magnesium salts. Exophytic membrane debris, a by-product of pineal neuronal and glial polypeptide exocytosis, forms the skeleton of the corpora arenacea around which calcifications are laid. Early calcifications found in younger patients are globular and localize near pinealocytes; whereas in older patients, calcifications tend to be more concentric and are associated with glial cells. In a study of older adults (age range, 62–80 years), Kim et al. found multiple concentrically calcified areas distributed throughout the pineal gland with centrally concentrated larger aggregates. Perivascular, tryptase-secreting mast cells have been found in areas where intrapineal calcium deposits originate, implicating these mast cells in the calcification process. Calcium also tends to localize along cell membranes. The outer pineal gland membrane may calcify, generally in a laminated manner. The laminated pineal calcifications that are found in older individuals may be related to circannual calcium level changes. However, the underlying cause of pineal calcium deposition remains elusive. It could represent a physiologic, maturational, or degenerative process.

In 1982, Zimmerman and Bilaniuk studied a large series of 725 patients (age range, 0–20 years) to define the incidence of pineal calcifications in the first 2 decades of life. In that study, the incidence of physiologic pineal calcifications was 11%; 13% of these were found in the first decade, with the youngest patient being 6.5 years. Because the study used CT parameters that would be considered primitive by today’s standards, including a relatively small matrix (160 × 160 or 320 × 320) and large section thickness (8 mm), the incidence of pineal calcifications was likely underestimated. Other literature contradicts the low range of 6.5 years. One study indicated a 3% frequency of pineal calcifications in patients younger than 6 years by using CT with a 512 × 512 matrix and 8-mm section thickness in nearly all cases. The youngest patient with pineal calcification was 3 years of age, and 1% of patients younger than 6 years had pineal calcifications, identical to our study results. Early pineal glandular calcium deposition has also been shown in the histopathology literature, where microscopic calcifications in the human pineal gland have been documented even before birth.

We found pineal calcifications in 5%. While most of these patients were older than 5 years, a few were younger (youngest, 3 years). The apparent decrease in the percentage of patients with pineal region calcifications between 7 and 9 years of age is likely attributable to fewer patients in older patient subsets (Fig 1). The mantra that no pineal gland calcification should be visible in patients younger than 6 years is no longer relevant in the modern CT age. However, pineal calcifications were single and punctate in all of our patients younger than 7 years of age. Occasional cases of larger and more numerous pineal calcifications were seen only in patients older than 7 years. Therefore, we advise cautious scrutiny in those patients younger than 7 years manifesting large and/or excessive pineal calcifications. Suspicious calcifications (premature, large, or excessive) should prompt a glandular volume analysis. A detailed clinical and biochemical assessment should be undertaken. Follow-up imaging should also be considered to exclude early presentation of a pineal tumor.

**Habenula**

The habenula is a small, bilaterally paired nuclear complex located along the inferior dorsomedial thalami. It is structurally connected to many deep brain structures, including the pineal gland, hypothalamus, midbrain, basal ganglia, and septal area, to name a few. Thus, it has a multitude of functions, mainly serving as a relay and processing center between the limbic system and midbrain through the stria medullaris thalami (afferent) and habenulo-interpeduncular tract (efferent), ultimately influencing dopamine and serotonin elaboration. The habenular complex also participates in the sleep-wake cycle. The retinohypothalamic reflex is the process by which light-induced retinal activation recognized in the hypothalamus signals the pineal gland to decrease melatonin production by way of the sympathetic chain. One function of melatonin is to inhibit the habenular complex. In the absence of melatonin-induced habenular inhibition, dopamine inhibition, normally mediated by the habenula, is eliminated, and dopamine excretion increases. Because the pineal gland and habenula are in proximity and share at least 1 common function, it is understandable that they may calcify at similar times, whether degenerative, physiologic, or otherwise. The substance of these calcifications is similar in makeup to that found in the pineal gland; the calcifications contain both calcium and magnesium salts.

To our knowledge, the potential association between sleep dysfunction and habenular calcification has not been systemically examined. However, there may be a link between habenular calcification and pathophysiology, ultimately disrupting the aversion/reward process. For instance, pineal and habenular calcifications have been noted as frequent findings in patients with schizophrenia. Because patients with schizophrenia often have difficulty learning from errors, habenular dysfunction has been implicated. Habenular calcifications were the most common epithalamic calcification in our cohort, present in 10% and in none younger than 2 years of age.

**Choroid Plexus**

The choroid plexus is an intraventricular organ comprising cuboidal epithelial cells, vasculature, and connective tissue. It has multifaceted functionality, chiefly serving in CSF volume modulation with productive, absorptive, and filtering capability. Its bulk resides in the lateral ventricular atria, called the glominus. During the human life span, the choroid plexus tends to accumulate physiologic calcifications. Because most choroid plexus is concentrated in the ventricular atria, it is here that calcifications are
most frequently evident. Pathologic processes such as prior intraventricular infection, inflammation, and hemorrhage and chronic calcium and phosphate imbalance can cause premature or excessive choroid plexus mineralization. Therefore, knowledge of a normally expected calcification time range would be helpful in distinguishing physiology from pathology.

The prevalence of choroid plexus calcification increases with age.19,20 However, few studies have examined choroid plexus calcifications in children. In a large retrospective CT study from 18,000 consecutive children (0–15 years of age) performed with early CT technology (EMI 1010 CT) by Kendall and Cavanagh,20 choroid calcifications were found in 2% of patients younger than 8 years of age. Another CT study that examined scans obtained with 8-mm section thickness and a 256 × 256 matrix demonstrated the prevalence of choroid plexus calcification in the first decade to be 0.5%.3 More recently, Doyle and Anderson13 found choroid plexus calcifications to be more prevalent, present in 16% of their patients. Indeed, choroid plexus psammoma bodies have been described in children histopathologically.21 Dystrophic calcification associated with the arachnoid cells and collagen fibers is thought to be causative.22

We discovered choroid plexus calcifications in 12% of patients in our cohort. In accordance with findings described by Doyle and Anderson,13 we found a significant positive association of choroid plexus and pineal calcifications. Choroid calcifications were visible in patients across all age ranges, from younger than 1 month to 10 years. A variety of clinical scenarios brought these patients to imaging, mainly trauma and headache. In all cases, the choroid plexus glomus was calcified in isolation, either unilaterally or bilaterally. Typically, calcifications were punctate and few. Therefore, choroid plexus calcifications occurring outside the glomus or in excess should be viewed with suspicion for underlying pathology in patients younger than 9 years of age. Minimal choroid plexus calcium deposition is probably physiologic at any age, including early childhood.

Dura
Thick connective tissue composing the dura envelops the central nervous system deep to the skeleton. Dura partitions the intracranial compartment into various subsections, forms the walls of the major dural venous sinuses, and serves as a conduit for neurovascular connections.

Although extremely rare, the CT prevalence of childhood dural calcifications remains a subject without strong support in the literature. In a series of 970 trauma patients, Ratcliffe et al23 found dural calcifications in >20% of men older than 30 years and women older than 50 years on skull radiography; attenuated calcifications were present in less than 2% of those younger than 40 years. Kendall and Cavanagh20 found dural calcifications in 0.8% of patients 0–15 years of age. Dural calcifications were rare in our study, occurring in 1%. Posttraumatic dural calcifications rarely occur but are not unexpected after a calvarial breach from penetrating trauma and surgical intervention. Indeed, most of our patients with dural calcifications had previous craniotomies (4/6).

The other 2 presented with headache (9 years of age) and head trauma (4 years of age). The tentorium was the most common area to calcify. Extensive dural calcifications in young patients should raise suspicion for pathologic conditions, including basal cell nevus syndrome, meningiomas, sequelae of prior subdural or epidural hemorrhage, and calcium/phosphate imbalance.24

This study has several limitations. Calcium detection is inherently limited by CT section thickness; therefore, intracranial calcifications may have been underestimated. However, we believe this underestimation to be of little clinical importance because thinner section soft-tissue algorithm images are not currently part of most practices. Despite their distinctive locations, pineal and habenular calcifications can sometimes be difficult to separate visually; this problem could have resulted in misclassification in some instances (Fig 2). Pineal cysts are common in the age group in our study and can distort regional anatomy.7 We found pineal cysts in 3% of our patient cohort, ranging from 4 to 15 mm; however, only 4 of these patients had epithalamic calcifications.

CONCLUSIONS
Physiologic intracranial calcifications may be found in the first decade, principally in children older than 3 years. Most epithalamic calcifications are habenular in location. Pinal and habenular calcifications were never present in children younger than 3 and 2 years, respectively. Dural calcifications are rare; they are not present in those younger than 2 years of age. Choroid plexus calcifications may be present in the very young (younger than 1 year).

REFERENCES