Craniopharyngioma and other cystic epithelial lesions of the sellar region: a review of clinical, imaging, and histopathological relationships

GABRIEL ZADA, M.D.,1 NING LIN, M.D.,1 ERIC OJERHOLM, B.S.E.,1 SHAKTI RAMKISSOON, M.D., PH.D.,2 AND EDWARD R. LAWS, M.D.1

Departments of 1Neurosurgery and 2Pathology, Brigham and Women’s Hospital, Harvard Medical School, Boston, Massachusetts

Objective. Cystic epithelial masses of the sellar and parasellar region may be difficult to differentiate on a clinical, imaging, or even histopathological basis. The authors review the developmental relationships and differentiating features of various epithelial lesions of the sellar region.

Methods. The authors performed a review of the literature to identify previous studies describing the etiological relationships and differentiating features of various cystic sellar lesions, including craniopharyngioma (CP), Rathke cleft cyst, xanthogranuloma, and dermoid and epidermoid cysts.

Results. There is significant evidence in the literature to support a common ectodermal origin of selected sellar and suprasellar cystic lesions, which may account for the overlap of features and transitional states observed in some cases. Research obtained from animal studies and reports of transitional cystic epithelial masses or lesions crossing over from typical to more aggressive pathological subtypes have collectively provided a solid foundation for this theory. Histological features that signify transitional entities beyond simple benign Rathke cleft cysts include squamous metaplasia, stratified squamous epithelium, and ciliated or mucinous goblet cells in squamous-papillary CPs. Several studies have identified key clinical, imaging, and histopathological features that can be used in the differentiation of these lesions.

Conclusions. The pattern of embryological formation of the hypothalamic-pituitary axis plays a major role in its propensity for developing cystic epithelial lesions. Subsequent inflammatory, metaplastic, and neoplastic processes may promote further progression along the pathological continuum, ranging from benign epithelial cysts to aggressive neoplastic cystic CPs. Selected clinical, imaging, and histopathological features can be used collectively to help differentiate these lesions and assign a formal diagnosis, thus accurately guiding further treatment.

(DOI: 10.3171/2010.2.FOCUS09318)

Key Words • craniopharyngioma • Rathke cleft cyst • epithelial cyst • embryology • magnetic resonance imaging

D ifferentiation of cystic lesions of the sellar and parasellar region may pose a diagnostic dilemma to neurosurgeons, radiologists, and pathologists involved in treating patients with these entities. As a result of the pattern of embryological development of the adenohypophysis from its stomodeal origin, as well as subsequent inflammatory, metaplastic, and neoplastic processes that can occur, the potential exists for the formation of a variety of cystic lesions in the sellar and parasellar region. The spectrum of cystic pathology occurring in the sellar region includes CPs, RCCs, colloid cysts, arachnoid cysts, cystic pituitary adenomas, xanthogranulomas, epidermoid cysts, dermoid cysts, and several others. In the majority of cases, a straightforward diagnosis of typical cystic lesions is established with little difficulty based on the lesion’s clinical, imaging, and histopathological characteristics. In some cases, however, a significant degree of overlap in these features occurs and may preclude the assignment of a definitive diagnosis. Establishing an accurate working diagnosis for sellar region pathology is critical in formulating appropriate surgical goals, predicting the likelihood of lesion recurrence, and guiding postoperative adjunctive management.

In 1994, Harrison and colleagues reported 19 cases of cystic epithelial lesions, in which overlapping radiological and histopathological features were evident in almost half, rendering 3 lesions without a formal diagnosis. Their study, in addition to several others, posited that cystic epithelial lesions may comprise varying entities along a continuum of pathology derived from a common ectodermal origin in the primitive craniopharyngeal duct.
thus accounting for the overlapping features observed in selected lesions. At one end of this pathological spectrum are benign RCCs, which are nonneoplastic lesions generally accepted to be derived from the remnant of the Rathke pouch. At the other end are CPs, which are neoplastic, often aggressive lesions thought to arise from squamous epithelial cell rests occurring anywhere along the region of the primitive stomodeum, from the sella to the infundibulum to the tuber cinereum and floor of the third ventricle.

In this summary, a current analysis of the developmental and clinicopathological relationships among CPs, RCCs, and other cystic epithelial derivatives of the sellar region is provided. The embryological development of the hypothalamic-pituitary region and various reasons for its particular predisposition to developing a variety of cystic lesions is discussed. Furthermore, we review the evidence favoring a common ectodermal origin, in contrast to alternative nonectodermal hypotheses, for the origin of cystic epithelial lesions. Finally, we review the typical and atypical features of RCCs and CPs, as well as the clinical, imaging, and histopathological features that have proved to be most useful in differentiating these lesions.

**Craniopharyngiomas: Typical Features**

Craniopharyngiomas can arise anywhere along the vestiges of the stomodeal diverticulum, but they most frequently originate in the region of the infundibulum, where squamous epithelial rests are known to occur. In rare cases, CPs arise in less typical locations along the remnants of the primitive craniopharyngeal duct, including the nasopharynx, sphenoid bone, or as primary intraventricular lesions. Overall, CPs comprise approximately 3% of all intracranial tumors, yet this proportion is notably higher in the pediatric population (10% of all pediatric brain tumors). The estimated incidence of craniopharyngiomas is 0.13 per 100,000 cases per year. Historically, CPs present in a bimodal age distribution with peak ages at the time of presentation of 5–14 years and then 50–74 years. However, these lesions may present in patients of all ages. The clinical presentation of CPs at any age frequently includes headache, vision loss, and hypopituitarism. In children, growth and sexual retardation, obesity, and hydrocephalus are frequently observed as well. Many patients with CPs suffer from chronic obesity, which is thought to develop secondary to hypothalamic dysfunction. Memory loss and cognitive deficits are more common findings in older patients. Diabetes insipidus is seen on presentation in 6–38% of new cases. The 2 major pathological subtypes of CP are the adamantinomatous and squamous-papillary varieties, although mixed-type lesions have been reported. These generalized tumor subtypes vary in age at presentation, tumor location, consistency, imaging characteristics, and histopathological features.

**Typical Imaging Features of CP**

From an imaging standpoint, CPs are typically described as calcified, solid, and/or cystic lesions, typically with a lobular shape and diameter of 20–40 mm. The majority of CPs involve the suprasellar space, with 40–53% of cases exhibiting some intrasellar involvement. Cranioopharyngiomas occasionally extend into the anterior, middle, or posterior fossa and may invade the floor or walls of the third ventricle. Hydrocephalus is observed in up to 38% of cases and is a more common finding in children.

On standard CT scanning, calcification is evident in 60% of tumors and is more common in pediatric cases and the adamantinomatous subtype. The majority of adamantinomatous CPs are mixed solid-cystic or predominantly cystic tumors with a lobulated appearance. On MR imaging, the solid elements are usually iso- or hypointense on T1-weighted images, exhibit inhomogeneous high intensity on T2-weighted images, and heterogeneously enhance following Gd administra-

---

**Fig. 1.** Imaging and histopathological examples of typical adamantinomatous craniopharyngiomas. 
A and B: Sagittal and coronal Gd-enhanced MR images obtained in a patient with a primarily solid suprasellar craniopharyngioma. 
C and D: Coronal Gd-enhanced MR images acquired in a patient with a mixed solid and cystic suprasellar craniopharyngioma. 
E and F: Photomicrographs of typical adamantinomatous craniopharyngioma composed of squamous epithelium arranged in sheets, lobules, and anastomosing trabeculae lined by palisaded columnar epithelium. H & E, original magnification ×100 (E) and 400 (F).
Cystic epithelial lesions of the sellar region

The cystic elements of adamantinomatous CPs typically display a high intensity on T1-weighted images, high or mixed intensity on T2-weighted images, and contrast enhancement of the cyst wall. The squamous-papillary subtype is found in approximately one-third of adult CP cases and rarely shows calcification. The majority of squamous-papillary CPs are predominantly solid or mixed solid-cystic tumors with a spherical shape, and usually exhibit low intensity on T1-weighted images, high intensity on T2-weighted images, and enhancement of the cyst wall after addition of Gd (Fig. 2). The MR imaging appearance of the solid regions is frequently similar to those of the adamantinomatous variety.

Typical Histopathological Features of CP

Histologically, adamantinomatous CPs are thought to arise from squamous embryonic rests and bear similarity to adamantinomas or ameloblastomas of the jaw with the potential for enamel production (Fig. 1). The epithelium is often stratified squamous or adamantinoid type, frequently with evidence of wet keratin nodules. The cystic components are often described as having a characteristic “machine-oil” interior, containing desquamated squamous epithelium and comprised mainly of keratin and cholesterol.

The papillary subtype of CP is known to occur more commonly in adults than children (14–50% of cranio-pharyngiomas in adults compared with only 2% of CP in children). Papillary CPs usually bear similarity to oropharyngeal mucosa and rarely exhibit calcification. The cyst contents are typically yellow and viscous. Histopathological analysis frequently demonstrates squamous epithelium forming pseudopapillae, without discrete nodules of wet keratin or calcium. There has been some debate as to whether adamantinomatous CPs demonstrate a higher potential for recurrence, although recent analyses have reported similar or slightly higher rates than their papillary counterparts. Several investigators have argued that papillary and adamantinomatous CPs may also represent 2 distinct entities that are located at opposite ends of a pathological continuum. It has been reported that various markers, including KL-1 or cytokeratin 7, can be used to distinguish the adamantinomatous and papillary varieties of CP.

Rathke Cleft Cysts: Typical Features

Rathke cleft cysts are benign, cystic remnants of the craniopharyngeal duct that are typically located in the sellar and suprasellar region. They are often discovered incidentally and have been identified in up to 22% of the population according to routine examination of
Typical Imaging Features of RCCs

On MR imaging, RCCs often appear as well-circumscribed, centrally located spherical or ovoid lesions of the sellar region. The majority of these smooth contoured cysts are unilobar with a diameter ranging between 5–40 mm (mean approximately 17 mm) (Fig. 3). They are often identified as having an epicenter located between the anterior and posterior pituitary gland in the region of the pars intermedia. The vast majority of lesions are intrasellar or intra- and suprasellar, with reports of purely suprasellar lesions occurring in a minority of patients. The normal pituitary gland may be displaced in any direction by an RCC, including circumferentially if the cyst arises in and remains encased within the gland.

In the majority of cases, administration of Gd contrast material demonstrates little or no enhancement of the cyst wall or contents on MR imaging. A thin peripheral rim of enhancement has been attributed to inflammation or squamous metaplasia of the cyst wall, or to a circumferential rim of displaced pituitary gland. The MR imaging signal intensity of cyst contents demonstrates high variability on T1- and T2-weighted sequences and has been reported to correlate with the nature of the cystic contents. In the series by Kim et al., the 3 most common signal patterns were a high intensity on both T1- and T2-weighted images, a low intensity on T1-weighted images with a high intensity on T2-weighted images, and a high intensity on T1-weighted images with a low intensity on T2-weighted images. Rathke cleft cysts filled with thin, CSF-like fluid generally exhibit a low intensity on T1-weighted images and a high intensity on T2-weighted images, while cysts with more proteinaceous, mucoid fluid correlate with higher intensity on T1-weighted images. Although most RCCs display a homogeneous signal intensity, up to 40% contain a waxy intracystic nodule composed of protein and cellular debris that typically fails to enhance following contrast administration.

Typical Histopathological Features of RCCs

Histopathologically, RCCs typically demonstrate simple columnar or cuboidal epithelium, often with ciliated or mucinous goblet cells (Fig. 3). Pseudostratified columnar cells are also commonly observed in specimens of RCCs. Squamous metaplasia of RCCs has been noted in 9–39% of patients and is associated with higher rates of cyst recurrence.

Epidermoid Cysts: Typical Features

Epidermoid lesions can occur anywhere in the intracranial cavity. They most commonly arise as extradural lesions or as intracerbral masses in the region of the cerebellopontine angle, but they may also present in the sellar and parasellar region. Epidermoid cysts often arise in a paramedian location, in contrast to typically midline dermoid cysts. Sellar and parasellar epidermoid tumors make up only 0.2–0.7% of major transsphenoidal series. Epidermoid tumors typically present in middle-aged patients with symptoms of mass effect, such as headache and vision loss. Some reports have also described an uncommon clinical presentation mimicking that of pituitary apoplexy. The cyst contents of epidermoid cysts, and many other epithelial sellar region cysts, can be caustic to the surrounding tissue, often resulting in hypophysitis, meningitis, or neurological deficits. Standard MR imaging cannot be reliably used in all cases to definitively establish a diagnosis of epidermoid or dermoid tumors, on account of their nonspecific MR imaging features. Demonstration of restricted diffusion on diffusion weighted imaging, however, has been shown to play a useful role in allowing the differentiation of epidermoid lesions from other types of cystic pathology, in particular arachnoid cysts.

Intraoperatively, epidermoid cysts can often be adhesive lesions not universally amenable to a gross-total resection. The cyst capsule and contents cannot consistently be dissected away from key vascular and nervous structures to achieve an acceptable outcome with minimal morbidity. Histologically, epidermoid cysts are characterized by a squamous epithelium, keratohyaline granule layers, and stratifications of “dry” keratin. Gross-total resection of intradural epidermoid tumors has been reported in 42% of cases, with a long-term recurrence rate of 26%. Resection remains the most effective modality, as no adjuvant measures have been proven to be of significant benefit in the management of these lesions.

Embryologic Development and Origins of Cystic Epithelial Lesions of the Sellar Region

Elucidation of the developmental processes accounting for the formation of the hypothalamic-pituitary system and related cystic lesions occurred as a result of several investigators over the course of several decades. Martin
Cystic epithelial lesions of the sellar region

Rathke\textsuperscript{85} was the first to describe the evagination process of the anterior forlet in 1838. In 1860, Huber von Luschka\textsuperscript{36} was the first to describe the presence of squamous epithelial rests occurring along the axis of the pituitary gland and infundibulum. Although Babinski\textsuperscript{5} and Frohlich\textsuperscript{28} each described epithelial suprasellar tumors at around the turn of the century, Jakob Erdheim\textsuperscript{24} first indicated that these lesions, which we now refer to as craniopharyngiomas, arise from squamous cell rests occurring in the region of the remnant hypophyseal/pharyngeal duct.

Embyrological Development of the Hypothalamic-Pituitary Region

The pituitary gland can be divided into 2 distinct anatomical compartments with different ectodermal origins. The adenohypophysis is ultimately formed by an ectodermal outpouching of the stomodeum (primitive oral cavity) located immediately anterior to the oropharyngeal membrane, known as the Rathke pouch. The neurohypophysis, on the other hand, develops from a downward extension of neuroectodermal tissue originating from the diencephalon, called the infundibulum.\textsuperscript{29,62,88}

The onset of neurulation begins with the primitive streak and node, which first appear on the dorsal aspect of the embryonic disc at approximately the 3rd week of gestational life. Once the neural tube is formed, transverse segmentation occurs at the midbrain, pons, and cervical levels, and a series of evaginations of the neural tube walls eventually produces several important structures in the CNS. The Rathke pouch (also known as the hypophysial diverticulum at this stage) is first noted to appear as an outpouching from the roof of the oral cavity at the 4th week of gestation. During the following weeks, this diverticulum gradually elongates and becomes constricted at its attachment site to the oral epithelium. At 6–8 weeks of life, the pouch loses its connection with the oral cavity and has grown in close contact with the infundibulum dorsally. Between the 3rd and 5th months of gestation, cells in the anterior wall of the Rathke pouch proliferate rapidly to form the pars anterior, whereas cells in the posterior wall do not divide significantly to form abundant glandular tissue. Instead, cells from this less active posterior wall form the pars intermedia, which is frequently not a prominent structure in the adult pituitary gland. Meanwhile, the infundibulum gives rise to the median eminence, the pituitary stalk, and the pars nervosa, or the posterior lobe of the pituitary gland. A small extension of the pars anterior, known as the pars tuberalis, extends superoventrally along the stalk and eventually surrounds it by the 16th week of gestation.

The extensive proliferation of the anterior wall of the hypophysial diverticulum eventually reduces its lumen size to that of a narrow cleft that is not typically recognizable in the adult pituitary. It is likely that CPs, RCCs, and other cystic epithelial lesions originate from remnants of this ectodermal cleft and the associated derivatives of the primitive stomodeum, where squamous cell rests are known to reside. In the past, 2 generalized theories have been proposed to explain the origins of CPs in this region. The “embryogenic theory” states that when the Rathke pouch is detached from the oral epithelium, remnants of ectopic craniopharyngeal duct may be deposited within the sellar region. The craniopharyngeal duct contains ciliated cells, which are derived from parts of the stomodeum and could be the origin of adamantinomatous craniopharyngiomas.\textsuperscript{24,29,80} The “metaplastic theory,” on the other hand, proposes that squamous epithelial cell rests that are found in the adenohypophysis and infundibulum can undergo metaplasia, thus giving rise to the papillary subtype of CP tumors.\textsuperscript{80,84,93} Evidence supporting and refuting the various theories for both ectodermal and non-ectodermal origins of cystic epithelial lesions is further discussed below. It is certainly possible that developmental as well as subsequent metaplastic and neoplastic processes can each play a role in the formation of various lesions along a spectrum of related pathology.

Evidence Lending Support to a Common Ectodermal Origin of Cystic Epithelial Lesions

The evidence lending support to the prevailing theory of a common ectodermal origin for many cystic epithelial lesions of the sellar region is derived from several sources. These sources include animal laboratory investigations, observational reports of transitional lesions containing features of multiple histopathological lesions, and case reports of progression from one lesion type to another, all of which have provided a credible basis for this viewpoint.

In a study by VanGilder and Inukai\textsuperscript{106} in 1973, oral mucosa was transplanted into the brains of 50 baby rats. The transplanted cells differentiated into a spectrum of
histopathological subtypes characterized by stratified squamous, cuboidal, and transitional epithelium as well as cholesterol clefts, calcification, and bone. The authors concluded that similar progenitor cells may differentiate into the spectrum of cystic epithelial lesions observed in the developing human brain. In another study, Iwata et al. performed a microscopic analysis of the hypophysis in rats. The authors found evidence of epithelial cranio-pharyngeal derivatives in approximately 0.16% of rats, and they suggested developmental rather than neoplastic origins of RCCs and related cystic epithelial lesions. Similarly, Schaetti and colleagues examined pituitary specimens in rats and reported finding epithelial cranio-pharyngeal derivatives consisting of cuboidal or columnar epithelium with goblet cells or stratified squamous epithelium. The authors of this study supported a heterogenous origin for many of these various epithelial cystic masses as well.

Several varieties of “transitional” or “crossover” cystic epithelial lesions with nonspecific features have been reported in the literature, collectively providing additional support for a theory of sequential progression in epithelial cystic lesions (Table 1). Although RCCs are considered nonneoplastic cystic lesions, recurrence rates following surgical intervention have been reported to be as high, or higher, than in some series following resection of CPs. More aggressive, or “transitional,” subtypes of RCCs have been reported to comprise a significant proportion of these recurrences and are frequently characterized by less typical histopathological features (Fig. 5). A theory of acute and chronic inflammatory processes, perhaps incited by repeated cyst leakage or microhemorrhage, has been implicated in inciting the process of squamous metaplasia identified in some RCCs. Between 9 and 39% of RCCs demonstrate evidence of squamous metaplasia, which has been independently associated with higher rates of cyst recurrence. The higher recurrence rates observed in RCCs with squamous metaplasia support the idea that these are likely more aggressive lesions that more closely approach the natural history of CP. The presence of stratified squamous epithelial cells in a minority of RCCs, as well as the higher MIB-1 labeling indexes associated with these lesions, also lends support to the theory that at least a subset of CPs may develop from such transitional intermediaries. A theory by Ikeda and Yoshimoto proposed that squamous epithelial cells in aggressive RCCs with higher proliferative indexes eventually outgrow and displace simple epithelium cell types.

More evidence for this hypothesis comes from reports of transitional-type pathology, consisting of histological features falling along different points of this histopathological continuum and occurring within the same lesion. In such cases, the intrasellar portion typically demonstrates features consistent with an RCC, whereas the suprasellar portion usually demonstrates features that are more typical for CP. Furthermore, numerous cases of ciliated epithelial cells or mucin-containing goblet cells occurring in squamous-papillary cranio-faryngiomas have been reported and implicated as providing further support for a common ectodermal origin and transitional entity between RCCs and CPs. As these rare lesions’ epithelial cells undergo transformation and no longer exhibit ciliation, it has been proposed that they develop into the more characteristic stratified squamous cell epithelial cells traditionally observed in the squamous-papillary subtype of CP.

Finally, several cases of ciliated squamous-papillary CPs have been reported to arise directly from preexisting RCCs. In one of these studies, Park et al. reported a case of an RCC with negative β-catenin accumulation that transitioned into a CP with positive β-catenin accumulation. A major limitation of these reports, however, is that for any given case it cannot be proven with certainty that a CP neoplasm actually arose from a preexisting RCC. The possibilities of coexisting lesions or dif-

### TABLE 1: Histological characteristics, incidence, and recurrence rates of various cystic epithelial lesions and their subtypes

<table>
<thead>
<tr>
<th>Lesion Type</th>
<th>Pathological Features</th>
<th>Incidence</th>
<th>Recurrence Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>RCC</td>
<td>simple cuboidal or columnar cells</td>
<td>28–54% of RCCs</td>
<td>3–19%</td>
</tr>
<tr>
<td></td>
<td>pseudostratified columnar</td>
<td>23–49% of RCCs</td>
<td>3–19%</td>
</tr>
<tr>
<td>“transitional”2,7,41,63,68,69,91,108,113</td>
<td>squamous epithelium, including squamous metaplasia; chronic inflammation</td>
<td>9–39% of RCCs</td>
<td>32–39%</td>
</tr>
<tr>
<td>CP</td>
<td>ciliated or mucin goblet cells</td>
<td>infrequent case reports</td>
<td>unknown</td>
</tr>
<tr>
<td>ciliated or goblet papillary30,70,77,79,87</td>
<td>ciliated or mucin goblet cells</td>
<td></td>
<td></td>
</tr>
<tr>
<td>squamous-papillary20,99</td>
<td>stratified squamous epithelium, pseudopapillae</td>
<td>28–33% in adults, 2% in children</td>
<td>0–12%</td>
</tr>
<tr>
<td>adamantinomatous1,26,109</td>
<td>adamantoid epithelium, wet keratin</td>
<td>66–68% in adults, 96–100% in children</td>
<td>13–22%</td>
</tr>
<tr>
<td>xanthogranuloma63,83</td>
<td>xanthogranulomatous component, little epithelium (usually squamous)</td>
<td>34% of suspected CPs</td>
<td>unknown</td>
</tr>
<tr>
<td>epidermoid cyst6,31,112</td>
<td>squamous epithelium, dry keratin, keratohyaline granules</td>
<td>&lt;1% of primary CNS lesions</td>
<td>0–26%</td>
</tr>
</tbody>
</table>

* Superscripted numbers indicate studies discussing the respective lesion.
Cystic epithelial lesions of the sellar region

Differences in biopsy technique and tissue analysis can be alternative explanations for this phenomenon.

Another noteworthy histological category with features resembling those of both CPs and RCCs is the sellar xanthogranuloma. These lesions tend to occur in younger patients (mean 27 years), have a smaller diameter, and remain primarily intrasellar with infrequent calcification. Although they have been reported to comprise a distinct entity, it remains unknown whether they are derived from RCCs or CPs following extensive inflammation and metaplasia, to the point that no epithelium is readily identifiable. Le and coworkers reported that the features of xanthogranuloma were more consistent with RCCs than CPs and demonstrated a high association with squamous metaplasia of these lesions.

Evidence Supporting Nonectodermal Origins of Cystic Epithelial Sellar Lesions

In the past, some investigators have favored theories supporting nonectodermal origins of various cystic epithelial sellar lesions. One alternative explanation is a theory that RCCs and related epithelial sellar and paraseellar cysts, including neureneric cysts and colloid cysts, are derived from an endodermal origin. The presence of histological features such as ciliation, goblet cells, and mucin associated with each of these lesions has provided the main argument in favor of this theory. As Harrison and colleagues described, however, similar histological findings can be identified in a variety of epithelial tumors located throughout the cerebrum. Furthermore, the same authors argued that the contents of the sella are purely ectodermal, with no valid explanation in place for how endodermal derivatives may arise here later during development, including theories of dysraphism. Another theory for the origin of cystic epithelial lesions in this region is that they are derived instead from neuroepithelial sources such as the neural crest, as supported by the finding of amyloid stroma in some examples. A third alternative theory suggests that cystic epithelial lesions are derived from metaplasia of anterior pituitary cells. Some authors have reported RCCs or CPs occurring in conjunction with pituitary adenomas as transitional or collision lesions. However, no evidence for a direct metaplastic origin of these cystic lesions from adenomas, or vice versa, has been proven as an alternative to a purely coincidental hypothesis.

Differentiation of Cystic CP and RCC

Although RCCs and CPs may represent 2 poles of a pathological spectrum, they are for the most part distinct entities, and according to Thapar and Kovacs, have “differences that are much more compelling than are their similarities.” Several previous reports have attempted to identify the clinical, radiological, and histopathological parameters that are most useful in accurately differentiating cystic sellar region pathology. In this section, these key characteristics are reviewed (Table 2).

Clinical Features

The majority of studies report overlapping clinical features for RCCs and CPs in adults, with regard to age distribution and sex. Similarly, presenting clinical features such as headache, endocrine deficits, visual deficits, and diabetes insipidus have not been reliably demonstrated to allow differentiation of CPs from RCCs. The only clinical features that have been reliably correlated with the diagnosis of CP over RCC, according to one study, were a significantly higher incidence of amenorrhea and neuropsychiatric deficits associated with CP.

Imaging Features

In many cases, imaging modalities can be used to more reliably differentiate RCCs from CPs and other cystic lesions. In some cases, this differentiation can be quite challenging, as the majority of studies have been unable to reliably do so based solely on T1- and T2-weighted intensity (Fig. 6). Calcification on CT imaging is often a useful characteristic for differentiat-
ing RCCs from CPs. In previous studies, 42–87% of CPs exhibited calcification, compared with only 0–13% of RCCs. It is important to note, however, that several cases of RCCs have been reported to occur with ossification and no evidence of neoplastic features, and that the presence of calcium is not necessarily pathognomonic for CP.

In 2006, Hofmann and colleagues reported that the imaging parameters that can be used to support a diagnosis of CP over RCC include: greater tumor diameter (> 2 cm), suprasellar location, and presence of calcification. In another study by Choi et al., MR imaging features were reviewed for RCCs, CPs, and cystic pituitary adenomas. Radiological parameters that supported a diagnosis of RCC were an ovoid shape, small cyst volume, and thin or no cyst wall enhancement. Conversely, a radiological diagnosis of CP was supported by features such as superior tumor lobulation, larger tumor volume, compression of the third ventricle, and a reticular enhancement pattern of the solid tumor portion. Kunii et al. used single-shot fast spin echo diffusion weighted MR imaging to differentiate cystic sellar and suprasellar lesions. They reported that RCCs could be identified using this imaging modality because of the lesion’s increased regional apparent diffusion coefficient values, in contrast with those of CPs and hemorrhagic pituitary adenomas. However, this modality was less useful in differentiating RCCs from cystic pituitary adenomas.

### Histopathological Features

The most reliable methods of distinguishing cystic epithelial lesions are clearly based on histopathological

<table>
<thead>
<tr>
<th>TABLE 2: Summary of clinical, imaging and histopathological characteristics that have been demonstrated to benefit in differentiating CPs and RCCs*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Feature</strong></td>
</tr>
<tr>
<td><strong>Clinical</strong></td>
</tr>
<tr>
<td>psychiatric deficits</td>
</tr>
<tr>
<td>amenorrhea</td>
</tr>
<tr>
<td><strong>Imaging</strong></td>
</tr>
<tr>
<td>calcification (CT)</td>
</tr>
<tr>
<td>size (&gt;20 mm)</td>
</tr>
<tr>
<td>location</td>
</tr>
<tr>
<td>suprasellar only</td>
</tr>
<tr>
<td>compressing 3rd ventricle</td>
</tr>
<tr>
<td>ovoid shape</td>
</tr>
<tr>
<td>cyst wall enhancement</td>
</tr>
<tr>
<td>single-shot fast spin echo diffusion weighted imaging</td>
</tr>
<tr>
<td><strong>Histopathology</strong></td>
</tr>
<tr>
<td>calcification</td>
</tr>
<tr>
<td>epithelial lining</td>
</tr>
<tr>
<td>adamantinomatous</td>
</tr>
<tr>
<td>squamous</td>
</tr>
<tr>
<td>simple columnar</td>
</tr>
<tr>
<td>simple cuboidal</td>
</tr>
<tr>
<td>pseudostratified columnar</td>
</tr>
<tr>
<td>ciliated</td>
</tr>
<tr>
<td>mucinous/goblet cells</td>
</tr>
<tr>
<td>keratin nodules</td>
</tr>
<tr>
<td>chronic inflammation</td>
</tr>
<tr>
<td>markers</td>
</tr>
<tr>
<td>CK</td>
</tr>
<tr>
<td>CK 20</td>
</tr>
<tr>
<td>nuclear β−catenin accumulation</td>
</tr>
</tbody>
</table>

* Superscripted numbers indicate studies discussing the respective feature. Abbreviations: NA = not applicable; ++ = quite common; + = more common; +/− = less likely; − = rare or absent.
Cystic epithelial lesions of the sellar region

![Fig. 6. Imaging examples demonstrating the potential difficulty in differentiating atypical CPs and RCCs. A and B: Sagittal and coronal Gd-enhanced MR images obtained in a patient with a recurrent RCC, demonstrating suprasellar, clival, and retrosellar extension. C and D: Sagittal and coronal Gd-enhanced MR images obtained in a patient with a cystic CP, also demonstrating suprasellar and infrasellar extension.](image)

and molecular markers, representing the gold standard for diagnosis. However, even at a microscopic and molecular level the differentiation of these lesions is often not clear cut, which perhaps further elucidates why the clinical and radiological features are often indistinguishable.

In a report by Shin and associates,[37] the histopathological features of RCCs and CPs were reviewed to identify those that correlated significantly with each lesion type. The authors determined that the characteristics correlating significantly with a diagnosis of RCC were simple columnar or cuboidal epithelium, pseudostratified columnar epithelium, and ciliation. Conversely, features supporting a diagnosis of CP included stratified squamous epithelium, calcification, keratin nodules, and chronic inflammation.

Nuclear immunohistochemical staining for β-catenin accumulation has been used as a reliable method of differentiation of some cystic sellar region lesions and has been shown to demonstrate immunoreactivity exclusively in CPs.[38] A similar study demonstrated β-catenin immunoreactivity in 77% of CPs, particularly of the adamantinomatous subtype.[39] Although useful, the downside of this modality is that it cannot be used to reliably differentiate squamous-papillary CPs from transitional RCCs, which is often the more formidable challenge.[30]

The expression patterns of various cytokeratins have also been reported to aid in the differentiation of RCCs from CPs yet with less consistency. In a study reviewing cytokeratin expression in 15 patients with cystic sellar lesions, Xin et al.19 reported that RCCs demonstrate expression of cytokeratins 8 and 20, whereas CP did not. However, a similar study in 2007 by Le and coworkers63 failed to demonstrate as reliable of a pattern, in which cytokeratin 8 reactivity occurred in all cases of RCC and CP.

Conclusions

Varying subtypes of sellar and parasellar epithelial cystic masses may be difficult to differentiate on a clinical, imaging, or even histopathological basis. There is significant evidence to support a common ectodermal origin of such entities, which may account for the overlap of features and transitional states observed in some cases. The pattern of embryological formation of the hypothalamic-pituitary axis plays a major role in its susceptibility to the development of such cystic epithelial lesions. Subsequent inflammatory, metaplastic and neoplastic processes may promote further progression along a pathological continuum ranging from benign epithelial cysts to aggressive neoplastic CPs. Research obtained from animal studies, reports of transitional cystic epithelial masses with nonspecific features, and reports of lesions crossing over from typical to more aggressive pathological subtypes have collectively provided a solid foundation for this idea. Selected clinical, imaging, and histopathological features can be used to aid in differentiating these lesions and assigning a formal diagnosis to guide further treatment.

Acknowledgement

The authors would like to thank Dr. Peter Chiarelli for assistance with journal collection.

Disclosure

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author contributions to the study and manuscript preparation include the following. Conception and design: G Zada, N Lin, ER Laws. Acquisition of data: G Zada, E Ojerholm, S Ramkissoon. Analysis and interpretation of data: G Zada, S Ramkissoon, ER Laws. Drafting the article: G Zada, N Lin, E Ojerholm, S Ramkissoon, ER Laws. Critically revising the article: G Zada, N Lin, E Ojerholm, S Ramkissoon, ER Laws. Reviewed final version of the manuscript and approved it for submission: G Zada, N Lin, E Ojerholm, S Ramkissoon, ER Laws. Study supervision: ER Laws.

References

6. Barrow DL, Spector RH, Takey I, Tindall GT: Symptomatic...


Cystic epithelial lesions of the sellar region

sellar tumors: diagnostic procedures and management. Neurosurg Focus 18 (6A):e6, 2005


Address correspondence to: Gabriel Zada, M.D., Department of Neurosurgery, Brigham and Women’s Hospital, 15 Francis Street, PBB3, Boston, Massachusetts 02115. email: gzada@usc.edu.