Unraveling the Mystery of Rathke’s Cleft Cyst Presenting with Hyponatremia: A Case Report with a Comprehensive Review of Literature

Shayan Huda1,2, Souvik Singha1, Ali Haidous1, Manju Harshan5, Luis Medina Mora3, Maria Devita6, Phillip R. Bukberg3, Bidyut K. Pramanik4, Amy McKeown1, John A. Boockvar1

1 Department of Neurosurgery, Lenox Hill Hospital, Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, New York, New York, United States
2 CUNY School of Medicine, New York City, New York, United States
3 Department of Endocrinology, Diabetes and Metabolism, Lenox Hill Hospital, Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, New York, New York, United States
4 Department of Radiology, Lenox Hill Hospital, Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, New York, New York, United States
5 Department of Pathology, Lenox Hill Hospital, Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, New York, New York, United States
6 Department of Nephrology, Lenox Hill Hospital, Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, New York, New York, United States

Abstract

Background Rathke’s cleft cyst (RCC) is a benign cystic lesion that is commonly discovered incidentally and remains asymptomatic in most cases. However, its association with the syndrome of inappropriate antidiuretic hormone (SIADH) secretion leading to hyponatremia (HN) is rare and has only been sporadically reported in the medical literature. In this article, we present a unique case of RCC manifesting with HN and discuss the diagnostic and management challenges encountered in a neurosurgical context. Additionally, we provide a comprehensive review of existing literature on RCC presenting with HN to enhance our understanding of this rare presentation.

Case Description A 56-year-old woman with acute-onset blurry vision, headaches, and low fluid intake was diagnosed with euvolemic HN secondary to SIADH. Further evaluation revealed an intrasellar cystic lesion consistent with RCC, which was successfully resected through endoscopic transnasal transsphenoidal surgery, resulting in a complete recovery without the need for hormone replacement.

Conclusion The most likely explanation for the HN due to SIADH in this case is the release of accumulated antidiuretic hormone (ADH) due to compression by the cyst and the irritating effect of inflammation at this location. Accurate evaluation and classification of HN are essential for proper diagnosis and management, considering the rarity of RCC presenting with HN. A multidisciplinary approach to treatment can lead to favorable functional outcomes; however, further research is necessary to better comprehend this unique clinical entity and optimize neurosurgical approaches.
Introduction

Rathke’s cleft cyst (RCC) is a benign cystic lesion that arises from remnants of Rathke’s pouch, an embryonic structure involved in the development of the anterior pituitary gland. These cysts are typically found in the sellar and suprasellar regions and are usually small in size and asymptomatic and most commonly identified as incidental findings. Larger lesions may result in symptoms by compressing the optic chiasm, pituitary stalk, and hypothalamus. The most typical symptoms in adults are vision loss, endocrine problems, and headaches, creating diagnostic challenges due to their overlapping clinical presentations with other sellar and suprasellar lesions.

Hyponatremia (HN) is a sellar–suprasellar lesion that can be a result of various factors altering the normal physiology of the hypothalamic–pituitary–adrenal axis. RCC presenting with HN due to the syndrome of inappropriate antidiuretic hormone (SIADH) secretion has been reported only a few times in the published literature (Table 1).

SIADH is characterized by euvolemic HN, typically with low serum osmolality, increased urine osmolality, and high urine sodium concentration. Patients with SIADH often present with nonspecific symptoms, such as nausea, vomiting, headache, confusion, seizures, and muscle cramps, which can range from mild to severe depending on the degree of HN and its rate of development. In SIADH, antidiuretic hormone (ADH) is secreted despite normal physiological conditions, resulting in impaired water excretion and dilutional HN. The pathophysiology of SIADH involves various mechanisms that lead to increased ADH secretion or enhanced renal responsiveness to ADH. Commonly reported etiologies of SIADH include tumors (such as lung cancer, small cell lung carcinoma, and oat cell carcinoma), central nervous system disorders, pulmonary diseases, medications, and certain systemic conditions.

In the context of RCC, SIADH can occur due to mechanical and/or chemical alternation of the hypothalamic–pituitary axis. Surgical resection of the cyst may be necessary in the cases where symptoms are significant or when complications like SIADH arise. Further research and understanding of the pathophysiology, diagnostic approaches, and management strategies for both RCC and SIADH are essential to optimize patient outcomes and improve clinical management in neurosurgery. This report focuses on a rare case of HN caused by SIADH in a patient with RCC, highlighting the challenges faced in treatment and management in the neurosurgical setting. We obtained written informed consent from the patient for the publication of this case report and accompanying images.

Case Report

A 56-year-old woman with no significant past medical history presented with acute-onset blurry vision with headaches, nausea, and low fluid intake for the past few days. Laboratory investigations revealed severe HN with a serum sodium level of 114 mEq/L, serum glucose level of 98 mg/dL, serum blood urea nitrogen (BUN) of 6 mg/dL, and creatinine of 0.50 mg/dL. Other investigations revealed serum osmolality of 247 mOsm/kg, urine sodium of 53 mEq/L, and urine osmolality of 321 mOsm/kg. Her endocrine workup revealed an 8 a.m. serum cortisol level of 12.06 µg/dL, thyroid-stimulating hormone (TSH) of 3.64 µIU/mL, free-T4 (FT4) of 1.10 ng/dL, adrenocorticotropic hormone (ACTH) of 17.4 pg/mL, follicle-stimulating hormone (FSH) of 68.15 IU/L (postmenopausal), luteinizing hormone (LH) of 33.76 IU/L (postmenopausal), prolactin 10.32 ng/mL, and insulin like growth factor 1 (IGF-1) of 116 ng/mL. Her blood pressure was within normal limits, and she was not dehydrated. Based on these, she was diagnosed as euvolemic HN secondary to SIADH. She was managed with hypertonic saline and fluid restriction, without success for 1 week in the hospital.

For further evaluation, she underwent brain magnetic resonance imaging (MRI), which showed an intrasellar cystic lesion, measuring 8 × 7 × 7 mm, which was T1 hypointense and T2 hyperintense, and was nonenhancing on T1-weighted contrast imaging. Based on the findings, our working diagnosis was RCC (Fig. 1). She then underwent endoscopic transnasal transsphenoidal resection of the cyst. She tolerated the procedure well and recovered well from anesthesia. During the postoperative period, she was treated with hydrocortisone and tolvaptan (Arginine Vasopressin-selective V2 receptor antagonist). Her HN gradually improved to 135 mEq/L in the postoperative period. Postoperative MRI revealed no residual disease with operative site changes (Fig. 1).

A histopathological examination of the surgical specimen revealed a cyst lined by ciliated columnar cells adjacent to unremarkable adenohypophyseal tissue. Reticulin stain showed normal lobular architecture of adenohypophysis and the cells are positive for cytokeratin, FSH, LH, ACTH, prolactin, growth hormone, TSH (rare cells), T pit (few cells), SF1, Pit 1, and negative for GATA3 and Ki-67 supporting the diagnosis. The diagnosis was consistent with RCC (Fig. 2).

At discharge, her Eastern Cooperative Oncology Group Performance Status (ECOG PS) was 1 and her Karnofsky performance status (KPS) was 90. At the 3-month follow-up, she is asymptomatic, with ECOG PS of 0 and KPS of 100, with normal serum electrolyte levels without any requirements of hormone replacement.

Discussion

The embryological Rathke’s pouch gives rise to RCCs. The primitive oropharynx develops a rostral outpouching between the third and fourth weeks of gestation, which eventually gives rise to the anterior lobe, pars intermedia, and pars tuberalis of the pituitary gland. The craniopharyngeal canal is formed by this rostral outpouching from the oropharyngeal ectoderm or Rathke’s pouch. The infundibulum originates from the diencephalon around the same time as the neurohypophysis. Around the fifth week of gestation, both structures come into contact with one another to form the adenohypophysis and neurohypophysis, respectively. By the sixth week of gestation, the Rathke pouch shrinks to a tiny cleft. RCCs, which are typically found in the pars...
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<td>Iwai et al</td>
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<td>SIADH (due to large cystic lesion) with secondary adrenal insufficiency (due to compression of the pituitary gland/stalk)</td>
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<td>Hsu et al</td>
<td>18/F</td>
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<td>Ogawa et al</td>
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<td>Retro-orbital pain, nausea, vomiting, and bitemporal hemianopsia</td>
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<td>73/M</td>
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| 51/F        | 33/M    | Sudden severe headaches, visual changes, dizziness, nausea, vomiting, and metallic taste in his mouth | 121                                     | Sellar and suprasellar region; transsphenoidal resection | RCC            | Transient hypocortisolemic state resulting in hyponatremia | Outcome: NA  
Levothyroxine and hydrocortisone |
| 75/M        | 70/M    | Drowsiness                                                                            | 126                                     | Sellar and suprasellar region; transsphenoidal resection | RCC            | Same                                                                                   | Outcome: NA  
Fludrocortisone + dexamethasone |
| 33/F        | 66/F    | Loss of consciousness                                                                  | 126.2                                   | Sellar and suprasellar region; transsphenoidal resection | RCC            | Same                                                                                   | Outcome: NA  
Hydrocortisone |
| Sivakumar et al⁸ | 49/M   | Headache, nausea, appetite loss, and fatigue; 1 mo later, thirst, polydipsia, and polyuria | 112                                     | Intrasellar region; medically managed                | RCC (based on imaging) | Chemical damage to the posterior pituitary caused by leaked RCC content: ADH hypersecretion (SIADH)  
Axons degenerate from the posterior pituitary to the magnocellular cell bodies: CDI | Outcome: NA  
Good outcome  
Medical management  
For SIADH: fluid  
For CDI: dDAVP |

Abbreviations: ADH, antidiuretic hormone; CDI, central diabetes insipidus; dDAVP, desmopressin; RCC, Rathke’s cleft cyst; SIADH, syndrome of inappropriate antidiuretic hormone.
Fig. 1 (A–D) Preoperative magnetic resonance imaging (MRI) reveals 8 × 7 × 7 mm round focus of T1 hypo-, T2, and fluid-attenuated inversion recovery (FLAIR) hyperintense and T1 contrast nonenhancing lesion in the intrasellar region and at the dorsal aspect of the pituitary gland, just left of the midline without any suprasellar extension. The pituitary stalk is in the midline with normal thickness (2–3 mm). (E, F) Postoperative MRI reveals a T1 hyperintense signal at the operative site and (G) postoperative computed tomography (CT) demonstrates a left-sided cystic cavity without any evidence of residual lesion.

Fig. 2 (A) Histological examination reveals a portion of the cyst lined by ciliated columnar cells (hematoxylin and eosin [H&E] stain at 40X), juxtaposed to unremarkable adenohypophysial tissue and containing detached thick eosinophilic material. (B) Reticulin staining (20X) confirms the normal lobular architecture of the adenohypophysis. (C) Cells have positive immunoreactivity for cytokeratin (20X). (D) Negative glial fibrillary acidic protein (GFAP) (20X).
intermedia, are formed as a result of the persistence and expansion of these clefts. Various theories have been described in the published literature to explain the HN associated with RCC (Table 1). Hypocortisolemia as a Result of Chronic Inflammation of the Pituitary Gland Causing Hyponatremia Cortisol promotes sodium reabsorption and potassium excretion in the collecting tubules, which helps maintain plasma volume and serum sodium levels. In cases of hypocortisolemia, there can be a reduction in sodium reabsorption, leading to secondary HN. This hypothesis is further supported by the observation that patients recovered from HN with corticoid treatment alone with low baseline serum cortisol levels, without requiring additional sodium supplementation. The development of HN can be initiated by recurrent intracystic hemorrhage and cyst rupture, leading to inflammation of the cyst wall. This inflammatory process, along with histological repair, results in changes from a single cuboidal epithelium of RCC to stratified epithelia and significant interstitial fibrosis causing pituitary atrophy with reduced reserve. The deficiency in the mineral cortex effect of cortisol can result in HN. Nevertheless, the normal preoperative serum cortisol level eliminates this possibility in our case.

Salt Wasting as a Cause of Hyponatremia Certain intracranial organic lesions can promote sodium diuresis, known as cerebral salt wasting. This condition involves disturbances in the hypothalamic function and an increase in the concentrations of atrial natriuretic peptide (ANP) and brain natriuretic peptide (BNP). Considering that these patients usually present with hypovolemia and signs of dehydration (hypovolemic HN), it is highly improbable that this possibility is responsible for HN in our case.

SIADH as a Cause of Hyponatremia ADH is produced by magnocellular neurosecretory cells located in the hypothalamic supraoptic and paraventricular nuclei. Subsequently, it travels through axons that traverse the infundibular stalk and reaches the posterior pituitary, where ADH is stored in vesicles until its controlled release into the bloodstream. One of the explanations of this plausible causation is that RCC compresses the posterior pituitary, which can lead to SIADH due to the release of accumulated ADH into the bloodstream. Additionally, as the inflammation progressed along the magnocellular axons, SIADH persisted due to the release of AVP from the degenerating axons. Considering the euvolemic status and clinical presentation, this theory (HN secondary to SIADH) appears to be the most plausible explanation for HN in our case.

To address the RCC and its potential contribution to the patient's SIADH, a transsphenoidal resection of the cyst was performed. This neurosurgical intervention aimed to alleviate the effect of the lesion on the hypothalamic–pituitary axis and the possible disruption of normal ADH regulation. Following the resection, the patient experienced an immediate improvement in symptoms, and her HN also got corrected. And in subsequent follow-ups, she did not require any hormone replacement therapy to maintain normal electrolyte levels. The relationship between RCC and SIADH warrants further investigation to comprehensively understand the intricate pathophysiological interplay.

Conclusion The management of SIADH in the neurosurgical setting is highly challenging, demanding a delicate balance between fluid restriction and the prevention of complications arising from HN. Collaborative efforts between neurosurgery, endocrinology, and other specialized disciplines are paramount to optimize patient outcomes and ensure appropriate management. This intriguing case report underscores the rare association between RCC and SIADH. The unusual presentation of SIADH in this context highlights the need for vigilant monitoring and meticulous management of HN in neurosurgical patients. Prompt recognition of SIADH in the setting of RCC, along with interdisciplinary collaboration and timely intervention, holds the key to achieving the best possible outcomes in these complex cases. Further research and investigations are imperative to enhance our understanding of this unique clinical entity and optimize treatment approaches in the realm of neurosurgery.
Conflict of Interest
None declared.

References