Coexistence of Suprasellar Lesion and Pulmonary Congestion: Two Rare Cases and Literature Review

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Abstract

Background: The characteristics of sellar/suprasellar lesion are endocrine disturbance, obesity and sleepiness disorder. However coexistence suprasellar lesion with pulmonary congestion is extremely rarely reported.

Case presentation: Two cases in the present report included typical traits of sellar/suprasellar tumor, besides they were identified preoperative pulmonary congestion by chest computer tomography. Abnormal hormone level accompanying with obesity induced and accelerated preoperative pulmonary congestion. Because coexistence of sellar/suprasellar lesion with pulmonary congestion is very rare, we confirmed the ultra-rare cases by literature reviewing.

Conclusion: Implementation of general anesthesia to patients with sellar/suprasellar lesion is a challenge to every anesthesiologist due to their complex pathophysiological changes. Occurrence of pulmonary congestion increases extra burden to anesthesiologists. Because ignorance of the accompanying pulmonary congestion might produce irretrievable disaster, the importance of preoperative evaluation and makeup of anesthesia plan should be stressed. Additional, the diagnosis of the second case was Erdheim-Chester disease, a very rare disease in children. This was the forth youngest case of this disease reported by searching PubMed and PMC.

Introduction

The overwhelmingly lesion in sellar/suprasellar region is pituitary adenomas, however this region can be affected by a variety of non-pituitary lesions, including craniopharyngiomas, cystic nonneoplastic lesions, inflammatory lesions, and so on [1,2]. Given that various diseases occurring in this region, as well as high operative risks, therefore arriving at the right preoperative diagnosis is important for taking appropriate therapeutic decisions, for instance, selecting surgical approach and necessity, ameliorating surgical complications [3]. Implementation of anesthesia to patients suffering sellar/suprasellar lesions is a challenge to every anesthesiologist, due to these lesions frequently accompanying with various neurologic and endocrine dysfunction, depending on properties of lesion, and mass effect on adjacent structures [4]. For instance, space-occupying mass may squeeze pituitary stalk, such compression is able to lead to development of endocrine disorders [5]. Meanwhile, sellar/suprasellar tumor might lead to obesity by different pathophyslogic mechanisms and produce various clinical symptoms [6]. The development of obesity deteriorates patient quality of life, but also raises airway-management problem to anesthesiologists. Here we reported two cases of supersellar lesion, which had to undergo stereotactic biopsy for preoperative diagnosis. Except common symptoms, such as hormone deficiency, sleep disorder and obesity, both of them appeared obvious preoperative pulmonary congestion. The coexistence of supersellar lesion and pulmonary congestion is very rare. Detailed PubMed and PMC search for sellar/suprasellar lesion associated with pulmonary congestion yielded very few case, we demonstrated its rarity by literature review. Different anesthesia implemented to the two cases produced different clinical results. Awareness is needed of new pathological entities in sellar/suprasellar lesions. We discussed the potential causes for coexistence of sellar/suprasellar lesion and pulmonary, which was useful to better and comprehensive preoperative evaluation.
**Case report**

**Case 1**

A 5-year-old girl (Table 1) was admitted to our hospital presenting polydipsia and polyuria. Her symptoms were presented three years ago. Except that, she had no relevant medical history. Brain Magnetic resonance image (MRI) scan detected a mass lesion in suprasellar area with T1 hypointensity and T2 hyperintensity and homogeneous enhancement. Recently, her parents complained with respect to her daytime somnolence and increase of sleep. Limb seizures appeared occasionally without conscious disturbance. Physical examination was remarkable for her obesity and very short neck, with obvious apnea when sleeping. Coarse breath sounds were found on both sides of chest. Reexamined MRI showed the mass was enlarged (Figure 1). Chest computer tomography (CT) detected increased bronchovascular shadows and pulmonary congestion on bilateral pulmonary fields (Figure 2A). Results of laboratory test prompted central hypothyroidism, but also identified cortisol deficiency, coexisting with hyperosmolar status and hyperlipidemia (Table 2). No adjuvant hormone supplement was administered.

![Preoperative brain MRI of the suprasellar lesion. T1 sagittal, T1 axial and T1 coronal sections showed a well-defined hypo-intensive suprasellar lesion, 31*28*22mm. The lesion was homogeneous enhancement.](image1)

![Preoperative axial cross-section images](image2)

**Figure 1**: Preoperative brain MRI of the suprasellar lesion. T1 sagittal, T1 axial and T1 coronal sections showed a well-defined hypo-intensive suprasellar lesion, 31*28*22mm. The lesion was homogeneous enhancement.

**Figure 2**: Preoperative axial cross-section images

A. Showed signs of pulmonary congestion with dilated pulmonary veins and bronchial cuffing, as well as thickened septal lines.

B. Intraoperative lung window revealed sever bilateral pulmonary edema with ground-glass opacity, concomitance with pulmonary consolidation and pulmonary effusion in bilateral lower lobes

C. Postoperative lung CT on admission ICU 48h demonstrated most lung field clear except consolidation of bilateral lower lobes.
Table 1: The demographic of two cases.

|                | Case 1       | Case 2       |
|----------------|--------------|--------------|
| Age            | 5Y3M         | 5Y5M         |
| Gender         | Female       | Male         |
| Body weight (Kg)| 32           | 35           |
| Length (cm)    | 95           | 113          |
| BMI (Kg/m²)    | 35.46        | 27.41        |

BMI: Body Mass Index.

Table 2: The laboratory test parameter of two cases.

|                  | Case 1       | Case 2       |
|------------------|--------------|--------------|
| ACTH (<46pg/ml)  | <5           | 11.70        |
| Cortisol (5~25ug/ml) | 1.41       | <1           |
| TSH (0.35~4.94ulu/ml) | 2.696       | 5.013        |
| TotalT4 (4.87~11.72ug/dl) | 3.27       | 4.67         |
| Free T4 (0.7~1.48ng/dl) | 0.56        | 0.67         |
| Na+ (135~145mmol/L) | 154.8       | 162          |
| Cl- (95~110mmol/L) | 119.8       | 128          |
| TSH (0.35~4.94ulu/ml) | 2.696       | 5.013        |
| TotalT4 (4.87~11.72ug/dl) | 3.27       | 4.67         |
| Free T4 (0.7~1.48ng/dl) | 0.56        | 0.67         |
| Na+ (135~145mmol/L) | 154.8       | 162          |
| Cl- (95~110mmol/L) | 119.8       | 128          |
| AST (3~35U/L)    | 39.9         | 58.8         |
| α-HBDH (72~182U/L) | 299         | 266          |
| LDH (120~250U/L) | 464          | 396          |
| ALP (3~120U/L)   | 499          | 409          |

ACTH: Adrenocorticotropic Hormone.
TSH: Thyroid Stimulating Hormone.
TG: Triglyceride.
AST: Alanine Aminotransferase.
α-HBDH: Hydroxybutyrate Dehydrogenase.
LDH: Lactate Dehydrogenase.
ALP: Alkaline Phosphatase.

The patient was treated with levothyroxine and corticosteroid; homeostasis was monitored and adjusted daily. Four days after hospitalization, she underwent stereotactic biopsy with general anesthesia. However severe negative pressure pulmonary edema occurred during the process of induction (Figure 2B). Treatment of positive pressure ventilation was administrated immediately, accompanying with hormone supplement. After 48 hours intensive care, she recovered to preoperative status. Chest CT revealed bilateral lung fields were clear except consolidation on bilateral lower lobes (Figure 2C). Even the most of lung fields were clearer than preoperative CT images. However, her parents refused the second surgery for biopsy.

Case 2

A 5-year-old boy (Table 2) complained polydipsia and polyuria about one year, without relevant medical history. Brain MRI identified a suprasellar mass with T1 isointensity and T2 hyperintensity, meanwhile it was heterogeneous enhancement. His body weight increased more than 15kg in transient period, and somnolence with snoring was gradually obvious in recent couple of months. Physical examination demonstrated obesity and somnolence without other obvious symptoms. Reviewing brain MRI demonstrated the lesion was larger than the previous (Figure 3). Chest CT showed bilateral pleural opacity and increased bronchovascular shadows (Figure 4). Laboratory tests identified hypothyroidism and hypocortisolemia, as well as hyperosmolar status and hyperlipidemia (Table 2). After admission, he subsequently commenced levothyroxine and hydrocortisone supplement therapy, and homeostasis was maintained. Because of the lesson from the first case, he underwent stereotactic biopsy with local anesthesia. The operation was successful and pathological diagnosis based on microscope and immunohistochemical findings was Erdheim-Chester disease (ECD) (Figure 5).
Figure 3: Preoperative magnetic resonance imaging of brain T1-weighted MRI
A. sagittal view,
B. axial view and
C. coronal view revealed multilobulated complex mass in the suprasellar region measuring 23*17*15mm. The lesion was iso-intensive and heterogeneous enhancement.

Figure 4: Lung window of CT taken on admission revealed bilateral opacity and interlobular septal thickening and peribronchovascular interstitial thickening.

Figure 5: The results of immunohistochemistry of suprasellar lesion. GFAP (-), Olig-2 (-), S-100 (+), Vimentin (+), LCA (+), CD68 (+), CD3 (-), CD1a (-), CD163 (+), ATRX (+), IDH1R123H (+), Ki-67 (+5-15%), P53 (+), CD117 (-), CD163 (+), ATRX (+), IDH1R123H (+), Ki-67 (+5-15%), P53 (+), CD117 (-), Oct3/4 (-), SALL4 (-), PLAP (-), H3K27M (-), MAC387 (-), BRAFV600E (+), BRAFV600 positive contrast (+). The color photo is only available online.

Discussion and Literature Review

The most common clinical symptoms of sellar/suprasellar lesions appeared in the two cases, such as central hormones insufficiency, obesity, and sleep disorder. Besides that, they possessed obvious preoperative pulmonary congestion. Negligence of it made more severe pulmonary edema and consolidation in case 1, thus more attention had to be paid on the second case with appropriate anesthesia strategy. Follow-up confirmed the second case received prompt treatment due to the successful stereotactic biopsy. What etiologies did cause the preoperative pulmonary congestion? Pediatric sellar/suprasellar tumor share many clinical similarities...
due to their anatomic location adjacent to critical brain structures [7]. It is well knows, lesions penetrating in sellar/suprasellar area frequently appear clinical manifestation of anterior pituitary gland insufficiency caused by compression of the pituitary stalk, such as diabetes insipidus, obesity, circadian rhythm disorders, and so on [8]. Both of them were in state of hypothyroidism. For lung development, lung defense and surfactant synthesis, thyroid hormone is very important. Alteration of thyroid hormone level might affect lung function and central respiratory drive [9]. In 2015, Al-Sofiani reported a female case of non-cardiogenic pulmonary edema with history of hypothyroidism. The patient was not only with normal ejection fraction, but also with sleep apnea. So her statue was similar to the above two cases. Different from cardiogenic pulmonary edema, non-cardiac pulmonary is based on change of alveolar capillary permeability. Hypothyroidism was believed to induce non-cardiogenic pulmonary edema indirectly [10,11]. Because hypothyroidism might increase pulmonary capillary permeability and induce plasma proteins leakage [12], severe hypothyroidism might increase pleural effusion, which was associated with restrictive pulmonary function and respiratory failure [13].

Along with hypothyroidism, hypocortisolism also occurred in the two cases. Patients with hypocortisolism might be suspected to be more vulnerable to lung leakage syndrome [14], because cortisol may decrease capillary permeability and ameliorate lung endothelial barrier dysfunction. Both mechanisms are strongly associated with mortality of septic shock [15,16]. As to other diseases based on pulmonary endothelial injury and/or capillary permeability in cement, for instance ARDS, especially nonresolving ARDS, glucocorticosteroids might be an effective rescue option [17]. Although some authors reported not less than 20% patients with ARDS were administrated systemic steroids [18], its role in early ARDS remains controversial [19]. We reported the two cases with onset of hormone supplement treatment as soon as possible. In case 1, corticosteroid did promote resolution of respiratory failure, confirmed by chest CT follow-up.

There are other traits of sellar/suprasellar lesion in the two cases, such as obesity, sleep disorder. The body mass index (BMI) of the first case was 35 kg/m², meanwhile, body weight of the second case increased 15kg in short time, so as to his BMI was 27 kg/m². Sleep disorder was another remarkable sign, case 1 showed severe obstructive sleep apnea (OSA), and parents of case 2 complained his somnolence and snoring. Whatever obesity and sleep disorder, such signs are results of hypothalamic neuroendocrine disturbance, which is commonly related with space-occupying-effect and/or penetration of sellar/suprasellar lesion [6,8,20]. Vice versa, obesity with OSA not only disturbs concentration and rhythm of hormones, but also impairs peripheral sensitivity to hormones [21]. This is a vicious circle. So we suspected there were some associations between obesity with sleep disturbance and pulmonary congestion, even pulmonary edema. At least it could indirectly accelerate pulmonary pathological alteration by induction of hormone disturbance. Obesity is a status of increase of systemic and pulmonary circulations. In obese patient, elevation of pulmonary capillary blood volume causes subclinical interstitial edema due to abnormal alveolar membrane diffusion. In 2016, Oppenheimer and colleague observed a group of obese patients, who were with variable degrees of pulmonary vascular congestion, but no clinical cardiopulmonary dysfunction manifestation. Their results demonstrated alveolar membrane structure and pulmonary vascular permeability might change in these obese patients. Because airway compression and inflammation caused by obesity per se, these patients were vulnerable to pulmonary edema [22].

The characteristics of sleep disorder of both cases were OSA and snoring respectively. These signs are connected with upper airway obstruction. Because generation of negative intrathoracic press is more remarkable during obstructive episodes, pulmonary edema has been observed as a presenting symptom of OSA [23]. Breathing work is increased because of futile efforts against to closed upper airway. Excessive inspiration concomitant with repeated airway collapse produces exaggerated negative intrathoracic pressure fluctuation, which might raise left ventricular afterload provoking pulmonary congestion [24]. Additional reason we presented the two cases owing to the second case with ECD, confirmed by pathological diagnosis. ECD is a proliferative disorder of non-Langerhans histocytes with extremely low rate in children. About 550 cases have been reported in the literature to date, and pediatric ECD is ultra-rare [25]. There were only no more than 7 cases of pediatric ECD reported in PubMed and PMC literature till 2014 [26,27]. Recently another 7 cases of pediatric ECD were reported again from 2014. In these cases, intracranial ECD were more extremely rare. The second case in this report was just 5-year-old boy and the forth youngest ECD in PubMed and PMC literature until now, additional young cases included a 17-month-old, a 2-year-old and a 3-year-old respectively [28-30]. However we focused on his clinical problem related to anesthesia, but not the pathophysiological of ECD. Limitation

In the two cases, diagnosis of hormones insufficiency, the location of lesion, and pulmonary congestion was very definite due to radiological evidences and laboratory parameters. However the state of sleep disorder was assessed without objective index, besides of indirectly cues and some risks, such as obesity, apnea, excessive sleep and snoring. The ‘golden standard’ of OSA is apnea-hypopnea index of polysomnography [31], because our institution is a specialized neurosurgical hospital, overnight polysomnography test cannot be carried there. In addition, the extended time between diagnosis of brain lesion and occurrence of sleep disorder made parents of these two cases neglect connection between abnormal sleep and brain tumor. So the two cases had never received sleep referral. In fact, patients of sellar/suprasellar tumor with obesity are more likely at greater risk of sleep disorder breathing, however these symptoms related to sleep disturbance could not be identified early until diagnosis of tumor on sellar/suprasellar region [32].
Conclusion

Because of space-occupying effect and/or penetrating pituitary, sellar/suprasellar lesions often produce central hormonal deficiencies. Hormone disturbances put patients of sellar/suprasellar mass in high risk of obesity, which increases the possibility of developing sleep-disorder breathing or OSA. Compared to the mean of age and gender, the risk of OSA might raise 12% follow every increase in BMI of 1 kg/m². These disorders increase difficulties of general anesthesia, especially airway management. Besides, they work together to induce and accelerate pulmonary pathological changes, thus pulmonary congestion occurs. Coexistence sellar/suprasellar lesion with pulmonary congestion is very rare. Awareness is need of new pathological entities in sellar/suprasellar lesions, therefore more attention should be pay on preoperative evaluation and makeup of anesthesia plan and otherwise there might be disaster.

Informed consent

Informed parental consent was obtained from the individual included in the study.

Acknowledgment

Not available.

Ethical Statement

We informed the ethics committee of Sanbo Brain Hospital, Capital Medical University, and got its approval (Permit No.SBNK-2019-010-01).

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