

in view of the large functioning mass encasing major vessels. Chemo- and radio-nuclear therapy is not suitable in view of baseline ECOG-4. Decision for best supportive care was made with family members.

#### CONCLUSION

Due to its low prevalence, there is a low index of suspicion when it comes to diagnosing paraganglioma. Adding atypical presentation to this, this may lead to delay diagnosis and treatment. Early diagnosis is utmost important in a case of malignant paraganglioma since the treatment is by surgical removal.

### EP\_A061

#### APOPLEXY IN MICROPROLACTINOMA

<https://doi.org/10.15605/jafes.038.S2.79>

**Zhe Lan Wong, Eileen Tan, Ilham Ismail, Nurbadriah Jasmiad, Chee Keong See, Saiful Shahrizal Shudim**  
*Hospital Sultan Haji Ahmad Shah, Temerloh, Malaysia*

#### INTRODUCTION/BACKGROUND

Pituitary adenoma apoplexy is uncommon and often occurs spontaneously as a result of infarction, haemorrhage or a combination of both. Pituitary apoplexy is potentially life threatening and has long term consequences resulting in permanent hormonal deficiencies. It is often associated with pituitary macroadenoma and occurrence with pituitary microadenoma is rare.

#### CASE

We report a case of micro-prolactinoma with pituitary apoplexy on follow-up pituitary MRI.

A 29-year-old female who presented with secondary oligomenorrhea in December 2021 was subsequently diagnosed with micro-prolactinoma. Cabergoline therapy was initiated promptly. However, her serial prolactin level while on treatment would fluctuate between normal range and up to three times the upper limit. During her initial few months of treatment, she would develop intermittent headache which subsequently subsided. A repeat pituitary MRI was initially planned due to this complaint to exclude pituitary apoplexy, but MRI was postponed due to patient's claustrophobia. The repeat pituitary MRI was only performed in early 2023 revealing the presence of pituitary apoplexy with intratumoral haemorrhage. At the same time, she would also be diagnosed with hypocortisolism and hypothyroidism. Hydrocortisone and thyroxine replacement therapy were initiated and cabergoline therapy was stopped. During her follow-up she did not exhibit overt symptoms of hypocortisolism, hypothyroidism, or visual field defects.

#### CONCLUSION

This case illustrated that pituitary apoplexy could present in pituitary microadenoma and may present with subtle symptoms. Without a high index of suspicion, pituitary apoplexy can be overlooked leading to delay or missing diagnosis. Pituitary apoplexy requires prompt diagnosis with imaging and treatment can improve the clinical outcomes of patients.

### EP\_A062

#### A RARE CASE OF TOTAL LEFT ANTERIOR CIRCULATION INFARCT (TACI) SECONDARY TO THYROTROPINOMA (TSHoma) TREATED WITH OCTREOTIDE MEDICAL THERAPY

<https://doi.org/10.15605/jafes.038.S2.80>

**Yi Jiang Chua, Choon Peng Sun, Zanariah Hussein**  
*Division of Endocrinology, Department of Internal Medicine, Hospital Putrajaya, Putrajaya, Malaysia*

#### INTRODUCTION/BACKGROUND

TSHoma is a rare cause of functioning pituitary adenoma. Patients with TSHoma have a biochemical derangement of elevated free thyroid hormones with unsuppressed TSH. They are usually misdiagnosed and treated for primary hyperthyroidism at the initial diagnosis. Most common symptoms upon presentations are hyperthyroidism, goitre and visual field defects. We reported an asymptomatic middle-aged female who presented acutely with left TACI secondary to atrial fibrillation due to a large functioning TSHoma.

#### CASE

A 53-year-old female, with no known medical diseases, presented with sudden-onset aphasia and right-sided body weakness. She presented to the emergency department with hypertension and tachycardia and ECG showed fast atrial fibrillation. Thyroid function test (TFT) showed that patient had a discordant hyperthyroid result [FT4 51.1 pmol/L (reference range 7.9-14.4) and TSH 4.825 mU/L (reference range 0.34-5.60)]. Repeated TFT on another platform had similar results. MRI of the brain revealed a heterogeneous mass occupying the sellar region with suprasellar extension 1.8 x 2.6 x 3.7 cm (AP x W x CC). TRH stimulation testing confirmed a functioning TSHoma and the patient was started on IM Octreotide LAR with rapid improvement of the TFT and resolution of the atrial fibrillation within 2 months. As this patient was not fit for surgical operation during the acute presentation, she was treated with medical therapy of Octreotide LAR and the patient achieved good improvement in 6 months' time where she was able to function independently. A repeat MRI after 6 months showed a smaller sellar mass (1.6 x 2.3 x 3.5 cm).