CASE REPORT

18F-Fluorodeoxyglucose positron emission tomography/computed tomography findings in a patient with bilateral macronodular adrenal hyperplasia

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ABSTRACT
Adrenocorticotropic hormone-independent macronodular adrenal hyperplasia (AIMAH) is a rare bilateral adrenocorticotropic hormone (ACTH)-independent nodular adrenal hyperplastic disease. Most patients with AIMAH are usually asymptomatic and do not have adrenal hyperfunction. Herein, we reported the case of a 51-year-old female with bilateral macronodular adrenal hyperplasia with mild fluorodeoxyglucose uptake based on PET/CT imaging findings. Her symptoms resolved after surgical resection of the right adrenal gland.

INTRODUCTION
Clinically, hypercortisolemia is usually caused by an unexpected long-term increase in glucocorticoid hormones in tissue circulation. Hypercortisolemia is classified into two main categories: adrenocorticotropic hormone (ACTH)-dependent hypercortisolemia and ACTH-independent hypercortisolemia. ACTH-dependent hypercortisolemia accounts for 80–85% of cases, which 80% were pituitary tumours and 20% were cases of ectopic ACTH syndrome. ACTH-independent hypercortisolemia accounts for 15–20% of all patients and is mainly composed of adrenocortical adenoma, adrenal cortical adenocarcinoma, ACTH-independent macronodular adrenal hyperplasia (AIMAH) and primary pigmented nodular adrenocortical disease. The incidence of AIMAH with hypercortisolemia is less than 2%. We reported a case of bilateral macronodular adrenal hyperplasia in a 51-year-old female with PET/CT findings.

CASE REPORT
A 51-year-old female presented with oedema of the face and both lower extremities since more than 3 months. She had a history of hypertension for 30 years and cerebral haemorrhage 6 months ago, for which she received long-term standardised pharmacotherapy. Physical examination revealed typical Cushing’s syndrome appearance including moon facies and hairy and thin skin throughout the whole body. Serum aldosterone (690.90 pmol L⁻¹) and 24h urinary cortisol (2070.00 nmol L⁻¹) were elevated. Laboratory tests also confirmed mild hypokalaemia and abnormal circadian rhythm of cortisol concentration. No obvious abnormality was observed on pituitary MRI. CT revealed lobulated hypodense masses in both adrenal glands (Figure 1), following which a diagnosis of bilateral adrenal metastases was made. 18F-fluorodeoxyglucose (FDG) PET/CT examination with SIEMENS Biography 16 PET/CT scanner was performed to locate the primary tumour, PET images were acquired with two minutes per bed. The CT-based attenuation-corrected PET images were reconstructed with an iterative true X algorithm (true X 3D, three iterations, 24 subsets) and smoothed with a Gaussian filter with 4 mm FWHM (full width half maximum) (Matrix 168×168), which revealed mild FDG uptake in the adrenal lesions bilaterally (Figure 2). The patient’s immediate family also underwent bilateral adrenal CT examination to rule out familial genetic disease. Her father and sister underwent CT, which did not reveal any abnormality in the adrenal glands. The patient’s younger brother and son also underwent bilateral adrenal CT examination to rule out familial genetic disease. Her father and sister underwent CT, which did not reveal any abnormality in the adrenal glands. The patient’s younger brother and son also underwent CT for the adrenal glands, and revealed bilateral adrenal lesions. She was not willing to undergo genetic testing, and a diagnosis of familial genetic AIMAH with Cushing’s syndrome was made on the basis of the imaging findings and family history. Right laparoscopic adrenalectomy was performed under general anaesthesia, and pathological examination suggested macronodular hyperplasia.
of the right adrenal gland (Figure 3). The patient was discharged 1 week after surgery. Her clinical symptoms disappeared, blood hormone levels returned to normal, and she appeared healthy during the 7-month follow-up.

**DISCUSSION**

The incidence of bilateral hyperplastic adrenal macronodules is very low. The literature is limited to case reports and no longitudinal studies with a large study population. Few reports on AIMAH have described the $^{18}$F-FDG PET/CT imaging findings, which revealed abnormally high uptakes of $^{18}$F-FDG. This condition can easily be misdiagnosed as metastasis or lymphoma. Most metastases have primary malignant lesions in another part of the body and most primary adrenal lymphomas are diffuse large B cell lymphomas with hypermetabolism. However, our case manifested mild FDG uptake on PET/CT image, the maximum standardized uptake value (SUVmax: 3.9) is similar to the value of liver, which is different from the previous literature reported. The possible cause is that AIMAH is pathologically characterized by hyperplasia of cortical nodules and is not a real tumour, so its uptake value is similar to that of normal adrenal gland tissue, which showed a slight FDG uptake. Further studies are required based on large sample for mechanism of varied degree FDG uptake. Ginger-like appearance and retention of adrenal contour are the typical imaging features. The exact aetiology of bilateral adrenal macronodules is unclear and genetic variance in ARMC5 may be the main aetiology. Accordingly, genetic testing is necessary if AIMAH is suspected. AIMAH commonly causes Cushing’s syndrome. Several patients undergo unilateral adrenalectomy of the larger adrenal gland, which can help prevent the need for lifelong steroid replacement. Although unilateral adrenalectomy is safe and effective, subclinical cortisol hyperplasia may
occur after the procedure. Therefore, contralateral adrenalectomy was necessary. Thus, close follow-up is essential for patients who have undergone unilateral adrenalectomy. One-stage bilateral adrenalectomy may be the treatment of choice for elderly patients.10–12

A diagnosis of non-ACTH-dependent hypercortisolism should be considered if both adrenal glands show ginger-like appearance on CT and a slight FDG uptake on PET/CT, without an increase in FDG uptake in other parts of the body. The final diagnosis of AIMAH depends on postoperative histopathology.

LEARNING POINTS
1. The typical characteristics of adrenocorticotropic hormone-independent macronodular adrenal hyperplasia (AIMAH) include ginger-like appearance on CT image, retention of basic adrenal morphology, and mild-to-moderate fluorodeoxyglucose (FDG) uptake. Meanwhile, the other parts of the whole-body PET imaging are negative.

2. 18F-FDG PET/CT imaging could distinguish primary adrenal lymphoma and metastatic adrenal cancer from AIMAH, and offers the evidence for later surgical resection.

PATIENT CONSENT
Under the premise of hiding the patient’s information, the patient agrees that her medical data is published as a medical article exchange.

REFERENCES
1. Newell-Price J, Bertagna X, Grossman AB, Nieman LK. Cushing’s syndrome. Lancet 2006; 367: 1605–17. doi: https://doi.org/10.1016/S0140-6736(06)68699-6
2. Lindholm J, Jeul S, Jørgensen JO, Astrup J, Bjerre P, Feldt-Rasmussen U, et al. Incidence and late prognosis of Cushing’s syndrome: a population-based study. J Clin Endocrinol Metab 2001; 86: 117–23. doi: https://doi.org/10.1210/jcem.86.1.7093
3. Lacroix A. ACTH-independent macronodular adrenal hyperplasia. Best Pract Res Clin Endocrinol Metab 2009; 23: 245–59. doi: https://doi.org/10.1016/j.beem.2008.10.011
4. Verma A, Mohan S, Gupta A. ACTH-independent macronodular adrenal hyperplasia: imaging findings of a rare condition : A case report. Abdom Imaging 2008; 33: 225–9. doi: https://doi.org/10.1007/s00261-007-9236-y
5. Alencar GA, Fragoso MCBV, Yamaga LYI, Lerario AM, Mendonca BB. 18)F- FDG-PET/CT imaging of ACTH-independent macronodular adrenocortical hyperplasia (AIMAH) demonstrating increased (18) F-FDG uptake. J Clin Endocrinol Metab 2011; 96: 3300–1. doi: https://doi.org/10.1210/jc.2011-1397
6. Rockall AG, Babar SA, Sohaib SAA, Isidori AM, Diaz-Cano S, Monson JP, et al. Ct and MR imaging of the adrenal glands in ACTH-independent Cushing syndrome. Radiographics 2004; 24: 433–52. doi: https://doi.org/10.1148/rg.242035092
7. Espiard S, Drouget L, Libé R, Assié G, Perlemeine K, Guignat L, et al. Armc5 mutations in a large cohort of primary macronodular adrenal hyperplasia: clinical and functional consequences. J Clin Endocrinol Metab 2015; 100: E926–35. doi: https://doi.org/10.1210/jc.2014-4204
8. Albigier NM, Regazzo D, Rubin B, Ferrara AM, Rizzati S, Taschin E, et al. A multicenter experience on the prevalence of Armc5 mutations in patients with primary bilateral macronodular adrenal hyperplasia: from genetic characterization to clinical phenotype. Endocrine 2017; 55: 959–68. doi: https://doi.org/10.1007/s12020-016-0956-z
9. Elbelt U, Trovato A, Kloth M, Gentz E, Finke R, Spranger J, et al. Molecular and clinical evidence for an Armc5 tumor syndrome: concurrent inactivating germline and somatic mutations are associated with both primary macronodular adrenal hyperplasia and meningioma. J Clin Endocrinol Metab 2015; 100: E119–28. doi: https://doi.org/10.1210/jc.2014-2648
10. Iacobone M, Albiger N, Scaroni C, Mantero F, Fassina A, Viel G, et al. The role of unilateral adrenalectomy in ACTH-independent macronodular adrenal hyperplasia (AIMAH. World J Surg 2008; 32: 882–9. doi: https://doi.org/10.1007/s00268-007-9408-5
11. Kobayashi T, Miwa T, Kan K, Takeda M, Sakai H, Kanazawa A, et al. Usefulness and limitations of unilateral adrenalectomy for ACTH-independent macronodular adrenal hyperplasia in a patient with poor glycemic control. Intern Med 2012; 51: 1709–13. doi: https://doi.org/10.2169/internalmedicine.51.7041
12. Ito T, Kurita Y, Shinbo H, Otsuka A, Furuse H, Mugiva S, et al. Successful treatment for adrenocorticotropic hormone-independent macronodular adrenal hyperplasia with laparoscopic adrenalectomy: a case series. J Med Case Rep 2012; 6: 312. doi: https://doi.org/10.1186/1752-1947-6-312