Case Report

A rare case of long-term survival from metachronous bilateral adrenal metastasis of lung adenocarcinoma after combined surgical removal and immunochemotherapy

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Abbreviations & Acronyms
18F-FDG = 18F-fluorodeoxyglucose
AM = adrenal metastasis
Bev = bevacizumab
CBDCA = carboplatin
CEA = carcinoembryonic antigen
CT = computed tomography
Doc = docetaxel
FDG-PET/CT = 18F-FDG uptake on positron emission tomography/computed tomography
HE = hematoxylin–eosin
NSCLC = non-small cell lung cancer
PEM = pembrolizumab
Ram = ramucirumab

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Introduction: The prognosis of adrenal metastasis from non-small cell lung cancer is very poor. A recent report described the efficacy of the surgical removal of adrenal metastasis when solitary. However, metachronous bilateral adrenal metastasis is extremely rare, and a treatment strategy has not been established.

Case presentation: Herein, we describe a 52-year-old Asian male who presented with a right adrenal metastasis of non-small cell lung cancer 8 months after immunochemo-therapy and surgical resection of the lung. He underwent combined systemic treatment and a laparoscopic right adrenalectomy; however, after 9 months, a metachronous left adrenal metastasis emerged. A subsequent laparoscopic left adrenalectomy and systemic treatment led to long-term progression-free survival.

Conclusion: The appropriate surgical indication and combined systemic treatment of a metachronous bilateral adrenal metastasis in non-small cell lung cancer may extend the prognosis.

Key words: adrenalectomy, long-term survival, lung adenocarcinoma, metachronous bilateral adrenal metastasis.

Keynote message

We treated a case of bilateral metachronous adrenal metastasis of lung adenocarcinoma that successfully achieved long-term survival by a combination of adrenalectomy and immunochemo-therapy. This indicates the prognosis can be improved by an appropriate judgment with regard to surgical indication and systemic treatment.

Introduction

NSCLC frequently metastasizes to the adrenal gland. Most AM are multiple and the prognosis is poor. However, improved imaging modalities allows the detection of solitary AM in early disease at initial diagnosis or follow-up.1–3 Increasing cases of the surgical removal of AM have gradually been reported, and some patients with isolated AM achieve long-term survival after adrenalectomy.3–9,11–18 However, there is a lack of established evidence comparing adrenalectomy with non-surgical treatment, and clinical prognostic factor for disease relapse after surgical treatment are poorly understood. In addition, metachronous bilateral AM is extremely rare.7 We herein describe long-term survival in a case of metachronous bilateral AM of NSCLC after a combination of surgical removal and systemic immunochemotherapy.

Case report

A 52-year-old Asian man was referred to our department due to a left adrenal mass on CT following treatment for poorly differentiated adenocarcinoma in the left lung. He underwent a surgical resection of the left upper lobe of the lung. The pathological stage was T3aN0M0 (stage IIB), and four cycles of adjuvant chemotherapy (cisplatin, vincristine) were undertaken.
Abnormalities in vital signs or significant findings suspicious of Cushing syndrome were not observed on an initial visit. Except for an elevated CEA level of 23.9 mg/dL, a serum biomarker of lung adenocarcinoma, which became normalized soon after lung surgery, no significant findings were observed in the blood examination, including for adrenal hormones. Contrast-enhanced CT revealed a 15 × 12-mm–sized substantial mass in the left adrenal cortex with 18F-FDG uptake on PET–CT (Fig. 1a,b). Clinically, the mass was assumed to be an AM of lung adenocarcinoma, therefore, after informed consent, the patient underwent two courses of chemotherapy (CBDCA, PEM). However, the tumor did not completely regress, therefore, a laparoscopic left adrenalectomy (Fig. 2e) was performed. Pathological findings revealed atypical cells with anisokaryosis and pleomorphism infiltrating and forming nests in the adrenal cortex (Fig. 2f) as found in the primary lung adenocarcinoma (Fig. 2a–d). The adrenal tumor was weakly positive for TTF-1 (Fig. 2g) and negative for vimentin (Fig. 2h) by immunohistochemistry, consistent with characteristics of the primary tumor; its surgical margin was negative. The patient’s serum CEA level normalized to 4.6 mg/dL soon after the left adrenalectomy (Fig. 3), and an additional four cycles of systemic treatment of immunotheraphy (CBDCA, PEM, and Bev) were performed. However, 9 months after surgery, the serum CEA level had increased to 22.2 mg/dL. A 60 × 24-mm–sized mass in the right adrenal cortex was observed with 18F-FDG uptake on PET/CT (Fig. 1c,d); this was suspected to be an asynchronous AM of NSCLC. Recently, in an analysis of a limited series or several case reports, the long-term survival of patients who underwent a surgical resection of isolated AM, including laparoscopic procedures, was published. Patients showed an average 5-year survival of 25%. However, metachronous bilateral AM, as in our case, was rare. In an analysis of 43 patients who underwent the surgical removal of a solitary AM of NSCLC, only one case in 11 metachronous AM of NSCLC showed a recurrence in the contralateral adrenal gland. In view of the surgical strategy for AM of NSCLC, no unified guidelines exist on whether a radical or partial adrenalectomy should be performed. In the clinical course of our case, a partial adrenalectomy of the left AM could have been completed the first time. Therefore, an AE1/3 immunostain of the left adrenal tumor specimen was performed (Fig. 4). As a result, the dissemination of multiple tumor cells positive for AE1/3 (Fig. 4c,d), was recognized, apart from the mass, as not being detected by HE staining (Fig. 4a,b). Therefore, it is reasonable to select a radical adrenalectomy from the viewpoint of cancer control. In addition, an intraoperative renal capsule injury led to a high risk of retroperitoneal dissemination after the surgical removal of oral hydrocortisone. After an additional four cycles of immunotherapy (Ram, Doc), he was monitored as an outpatient, and was in remission for more than 4 years after the last surgery without any adrenal insufficiencies.

**Discussion**

Unilateral AM identified with initial treatment of NSCLC were generally classified as synchronous metastases; metastases not present when initial treatment was started were categorized as metachronous. Up to 4% of patients with operable NSCLC are estimated to have a synchronous unilateral AM. However, no consensus exists for the management of isolated AM of NSCLC. Recently, in an analysis of a limited series or several case reports, the long-term survival of patients who underwent a surgical resection of isolated AM, including laparoscopic procedures, was published. Patients showed an average 5-year survival of 25%. However, metachronous bilateral AM, as in our case, was rare. In an analysis of 43 patients who underwent the surgical removal of a solitary AM of NSCLC, only one case in 11 metachronous AM of NSCLC showed a recurrence in the contralateral adrenal gland. In view of the surgical strategy for AM of NSCLC, no unified guidelines exist on whether a radical or partial adrenalectomy should be performed. In the clinical course of our case, a partial adrenalectomy of the left AM could have been completed the first time. Therefore, an AE1/3 immunostain of the left adrenal tumor specimen was performed (Fig. 4). As a result, the dissemination of multiple tumor cells positive for AE1/3 (Fig. 4c,d), was recognized, apart from the mass, as not being detected by HE staining (Fig. 4a,b). Therefore, it is reasonable to select a radical adrenalectomy from the viewpoint of cancer control. In addition, an intraoperative renal capsule injury led to a high risk of retroperitoneal dissemination after the surgical removal of

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**Fig. 1** Contrast-enhanced CT (a) revealed a 15 × 12-mm–sized substantial mass in the left adrenal cortex with positive accumulation on FDG–PET/CT (b) (white arrowheads). CT showed a 60 × 24-mm–sized mass in the right adrenal cortex (c) with positive accumulation on FDG–PET/CT (d) (black arrowheads).
Therefore, reflecting on the clinical course of our case and findings, a radical wide resection of the mass, including the adrenal gland, should be considered first to control this rare malignant entity.
In the treatment of AM of NSCLC, surgical removal without systemic treatment is not usually recommended, although the evidence is unclear. Additional treatment after surgical removal did not have a positive effect on prognosis. However, cases were limited, and the retrospective nature of the study and selection bias may have led to an overestimation of efficacy. From a combination treatment point of view, the addition of systemic chemotherapy after adrenalectomy, rather than chemotherapy alone, could prolong survival for patients with AM of NSCLC. In a retrospective analysis of 37 patients with AM that mainly originated from the lung, more than 6 months of disease-free survival with systemic treatment after radical surgery of the primary lesion could be a favorable prognostic factor compared with that of less than 6 months. Undoubtedly the pathogenesis of AM of NSCLC is a very aggressive one; thus, a combination of sequential treatment of surgical removal following systemic immunochemotherapy might be needed to control the disease, as in our case. Further prospective trials are required in future to assess such combination strategic approaches in patients with AM of NSCLC.

**Conclusion**

We experienced a case of bilateral metachronous AM of NSCLC that was controlled by surgical resections and systemic treatment. This suggests that the prognosis of this disease can be extended by the appropriate surgical indication combined with systemic treatment.

**Author contributions**

Nami Tomiyama: Conceptualization; data curation; writing – original draft. Taku Naiki: Conceptualization; writing – review and editing. Aya Naiki-Ito: Data curation. Tatsuya Kawai: Data curation. Toshiki Etani: Data curation. Masakazu Gonda: Data curation. Maria Aoki: Data curation. Toshiharu Morikawa: Data curation. Yosuke Sugiyama: Data curation. Takahiro Yasui: Supervision; writing – review and editing.

**Conflict of interest**

The authors declare no conflict of interest.

**Approval of the research protocol by an Institutional Reviewer Board**

The ethics committees of Nagoya City University Graduate School of Medical Sciences gave approval for this study (#1574) and written informed consent was obtained in accordance with the World Medical Association Helsinki Declaration.

**Informed consent**

The patient gave written informed consent for the publication of this article and associated images.

**Registry and the Registration No. of the study/trial**

Not applicable.

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Editorial Comment

Editorial Comment on A rare case of long-term survival from metachronous bilateral adrenal metastasis of lung adenocarcinoma after combined surgical removal and immunochemotherapy

In clinical practice, urologists encounter adrenal gland metastasis from other malignancies occasionally. Most cases are presented with unilateral metastatic lesions in the adrenal gland, although limited cases are demonstrated with bilateral adrenal metastasis. Usually, unilateral resection of the adrenal gland with metastatic lesion could be indicated in case the patient only has unilateral adrenal gland metastasis and clinically permitted general condition and acceptable prognosis due to adrenal function preservation even after unilateral adrenal surgical removal. Based on experience, patients with lung or gastric cancer are frequently consulted by the urologic department for adrenal metastasis treatment. Recently, both cancers have been attempted to be treated with immunootherapy and combined immunochemotherapy using a molecular target drug.1,2 The national cancer comprehensive network guideline for nonsmall cell lung cancer has defined to use immunotherapy or immunochemotherapy drugs due to exhaustive gene expression analysis. For instance, described genetic abnormalities include sensitizing EGFR mutation, ALK rearrangement, ROS1 rearrangement, BRAF V600E mutation, NTRK gene fusion, and more than certain PD-L1 expression percentage. Each genetic abnormality corresponds to one to six recommended treatment regimens orderly.3 Therefore, lung cancer could be positioned as a systematized disease regarding the standardization of genetic examination for personalized medicine treatment decision. From the perspective of clinical sample acquisition for genetic testing, adrenal metastasectomy is worthwhile for respiratory and oncology physicians who are mainly responsible for the patients. Besides lung cancer, other cancers have also started to be treated using the same methods. In this case study, the patient underwent combined systemic lung cancer treatment and laparoscopic right adrenalectomy. Subsequent laparoscopic left adrenalectomy and systemic treatment led to the 4-year progression-free survival.4

The case study reminds urologists to have courage to conduct resection of both sides of the adrenal glands to achieve a cancer-free condition with permanent adrenal hormone replacement after screening the patient’s background, general conditions, and drug compliance.

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Conflict of interest

The authors declare no conflict of interest.