Metastatic Extra-Adrenal Pheochromocytoma with Single Kidney and Renal Compromise: A Case Report of Excellent Response, Tolerability, and Outcome to a Modified Regimen of $^{131}$I-mIBG, and Decision-Making between $^{131}$I-mIBG Therapy and PRRT

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Abstract

Determining the choice and the goal is key element for decision-making of a systemic radionuclide therapy. They should be clearly defined in deciding and individualizing the dose and regimen. For iodine-131 metaiodobenzylguanidine ($^{131}$I-mIBG) therapy, the important considerations during dose fractionation include disease burden, tumor biology, functional symptoms, and associated comorbidities, all of which are important determinants for the intent and course of treatment. Herein, we present the case of a 67-year-old elderly female with highly functional metastatic recurrent extra-adrenal pheochromocytoma (presenting 42 years after the primary surgery and 32 years following excision of pararenal recurrence) with multiple comorbidities including single kidney and borderline renal compromise, treated successfully with a relatively lower dose of $^{131}$I-mIBG (cumulative dose of 22.2 GBq in four cycles with a mean dose of 5.7 GBq per therapy cycle). The excellent tumor burden reduction, hormonal tumor marker response, and most importantly asymptomatic status could be achieved with the administered dose. On follow-up, none of the pretherapeutic parameters (including renal function) showed any further derangement compared with the baseline during next 24 months following the treatment. All cycles were well tolerated with only reversible hematological toxicity that normalized without any active intervention. The report is intended to provide some guidance for future therapeutic regimens.

Keywords

- $^{131}$I-mIBG therapy
- metastatic pheochromocytoma
- paragangliomas
- quality of life
- single kidney
- treatment response

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**Introduction**

Neural crest cell-derived tumors are extremely diverse; of them tumors consisting of sympathetic-adrenal cell lineage comprise pheochromocytomas (PCCs), paragangliomas (PGLs), neuroblastomas, and others. The majority of PCCs/PGLs are benign, with 10 to 20% extra-adrenal PGLs showing malignant course of disease and metastasis. The prognosis in malignant PGL/PCC is poor with overall 5-year survival <50%. The diagnostic workup includes blood/urinary levels of catecholamines or their metabolites, functional and anatomical imaging including iodine-131 metaiodobenzylguanidine (131I-mIBG) and fluordeoxyglucose positron emission tomography/computed tomography (FDG-PET/CT) and somatostatin receptor (SSTR)-based imaging, and tissue diagnosis with IHC markers. The management in metastatic cases remains mainly palliative and includes multimodality approach comprising 131I-mIBG therapy, SSTR-targeted therapies, chemotherapy, surgical debulking (where feasible), and novel antineoplastic agents.

As mIBG is an analog of norepinephrine and has high affinity to chromaffin cells and derived tumors, it has been used for molecular imaging and targeted therapy with success for these tumors since 1984. Its uptake and retention take place both by an active, ATP-dependent process known as “uptake-1” (major pathway) and energy-independent passive diffusion and have proven capability of selectively delivering substantial radiation dose to the tumor cells in both primary tumor and metastatic lesions. 131I-mIBG therapy is usually well tolerated even in the pediatric population when compared with chemotherapeutic regimens. The immediate adverse effects are nausea, vomiting, transient myelosuppression—prominently isolated thrombocytopenia (more common with chemotherapy), and rarely delayed effects—thyroid hypofunction and persistent myelosuppression. 131I-mIBG is usually administered as a fixed activity/according to patient weight alternative approach is to modify according to whole-body absorbed dose. As in myelosuppression, the dosages are also modified in impaired renal function as 131I-mIBG is principally excreted in the urine. Herein, the authors present the gratifying response and excellent tolerability profile following 131I-mIBG therapy with modified dose regimen including highlighting the typical therapy management involving pretherapy workup and post-treatment follow-up in a patient of metastatic extra-adrenal PCC with single kidney and baseline renal compromise and multiple other comorbidities.

**Case Report**

A 67-year-old female, known diabetic and hypertensive, with a history of open laparotomy due to abdominal symptoms 42 years ago concluded as left extra-adrenal tumor underwent mass excision first and again had left pararenal tumor recurrence for which she underwent excision and left nephrectomy 10 years after first surgery; the histopathology report inferred as extra-adrenal PCC. After 30 years, she had no significant symptoms and had been followed up in the initial years with yearly X-rays and clinical examination and abdominal ultrasound.

In her present history, around 18 to 20 months prior to the first cycle of 131I-mIBG therapy cycle, she presented with cough (for >3 months) and diarrhea; on biochemical investigations she showed raised 24-hour urinary vanillylmandelic acid (VMA), which was 9.5 mg/24 hours (normal <7), plasma-free metanephrine (PFM) was 537.9 pg/mL (normal <60 pg/mL), and plasma-free normetanephrine (PFN) was 1227.0 pg/mL (normal <140 pg/mL). Her blood pressure (BP) was 180/100 mm Hg; evaluation with contrast-enhanced CT abdomen and chest showed a 2.2 × 1.5 cm enhancing, soft tissue lesion around the left adrenal gland, multiple rounded lung nodules, largest ~1.4 × 1.4 cm, and abdominal lymph nodes (largest ~1.4 cm × 1.0 cm, pre-aortic). 18F-FDG PET/CT scan showed no significant tracer concentration in any of these lesions.

The patient commenced long-acting somatostatin analog (injection Sandostatin 30 mg LAR intramuscularly) per 3 weeks and other antihypertensive and antidiabetic medications including insulin by a local medical oncologist. Within 3 months the BP was 130/80 mm Hg and biochemical tests also showed reduced values of VMA to 6.1 mg/24 hours, PFM to 52.0 pg/mL, and PFN to 750.0 pg/mL. After this initial period of 15 to 18 months on medication, the patient showed few spikes in BP as the highest noted was 180/100 mm Hg with VMA raised to 27.9 mg/24 hours. Further 3 months

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Abbreviation: mIBG, metaiodobenzylguanidine.
later, there was raised PFM of 130.7 pg/mL and PFN of 2143 pg/mL. She did not show relief in her diarrhea and lost > 6 kg of weight (from ~63 to 56 kg) in the previous 1 year (prior to mIBG therapy). She also showed deterioration of her renal functions (serum creatinine from 1.02 to 1.5 mg/dL) and got referred to nuclear medicine for the consideration of appropriate radionuclide therapy. She underwent 1 mCi diagnostic 131I-metaiodobenzylguanidine (mIBG) whole-body scan (►Fig. 1), which showed multiple foci of increased tracer uptake in the bilateral lung fields and midabdomen, which on further single-photon emission CT (SPECT/CT) correlation showed tracer concentrating multiple abdominal nodules and bilateral lung nodules.

In view of age, Eastern Cooperative Oncology Group performance status of 2, and nephrectomized status (with serum creatinine value 1.5 mg/dL and estimated glomerular filtration rate [eGFR] of 34.0 mL/min), she was first considered for 131I-mIBG therapy rather than peptide receptor radionuclide therapy (PRRT). She was treated with 5,254 MBq (142 mCi) 131I-mIBG by intravenous infusion over 3.5 hours. Posttherapy planar scan (►Fig. 2) showed multiple foci of increased tracer uptake was noted in bilateral lung fields and midabdomen, VMA, PFM, and PFN showed 300 to 400% reduction from pretherapy values (from 27.9 to 1.6 mg/24 hours for VMA, 130.7 to 22.0 pg/mL for PFM, and 2,143 to 492.4 pg/mL for PFN) and controlled hypertension after 3 months of follow-up. She was treated with 131I-mIBG two times subsequently at an interval of 12 weeks with a dose of 6512 MBq (176 mCi) and 5,476 MBq (148 mCi). All therapies were well tolerated, with no significant adverse effect in the first 24 hours of treatment. In view of clinically asymptomatic status and excellent response, the fourth cycle of therapy was planned after some spacing around 9 months after third cycle and 5,735 MBq (155 mCi) of dose was given; this time patient showed clinically asymptomatic grade 3 hematological toxicity after 6 weeks of therapy that got reversed but over a period of 8 to 12 weeks. The VMA and PFM levels were constantly within normal range after first cycle. PFN levels were higher than normal limits, but 400 to 500% less than pretherapy values (from 2,143 to 371 pg/mL; ►Fig. 2). Adequate thyroid blockade using SSKI solution started from 1 day before therapy that is continued up to 14 days post-therapy and absence of 131I-mIBG uptake interfering medications was ensured at every therapy; this was verified with posttherapy scans and the thyroid function values were normal.

The pretherapy 131I-mIBG scan (►Fig. 3) before the last therapy showed tracer-avid bilateral lung fields and only small portion of left paravertebral regions. SPECT/CT correlation showed tracers concentrating left subcentimetric para-aortic lymph node and multiple bilateral lung nodules; no new sites were revealed. Comparing with first pretherapy scan, overall suggestive of partial response to therapy, multiple lesions showed reduced size and disappearance of few abdominal nodal lesions. On her follow-up from baseline to recent visit (6 months after the last therapy), significant symptomatic improvement was documented with complete disappearance of her primary complaint of diarrhea and cough (►Table 1). She did not show any paroxysmal hypertensive episodes with maintained BP in the range of 140 to 110 (systolic BP)/90–76 (diastolic BP). She had also gained 8 kg of weight in this period of 1 year (►Fig. 4). Assessment of quality of life using specially made intuitional questionnaire showed improvement in all subscales (►Fig. 5).

The follow-up hematological profiles, renal function, and LFTs tests were performed fortnightly for 12 weeks following treatment and during the pretherapy workup of subsequent therapies. The serum creatinine remained within control (varying between 1.10 and 1.60) while corresponding eGFR values varied between 45.0 and 30.6), improving following the 131I-mIBG therapy when compared to their baseline data (►Fig. 6). These revealed asymptomatic grade 1 hematological toxicity (according to the NCI CTCAE [National Cancer Institute; Common Terminology Criteria for Adverse Events]
Grading System) as predominantly thrombocytopenia 4 weeks after therapy, which returned to normal range in 6 to 8 weeks after the therapy (► Fig. 7).

Discussion

The neural crest tumors are relatively uncommon and widely heterogeneous class of tumors. Although malignant variants are less common, they contribute to high mortality and morbidity owing to their functional status and variable response to different therapies. It is observed that among neural crest tumors in adults, the PCCs and the PGLs arising in the abdomen secrete catecholamines and contribute to added morbidity. Being associated with several molecular markers such as VEGF, N-cadherin, and telomerase catalytic subunit, this subclass of metastatic tumors is generally considered poor responder to multiple therapeutic options available. Such tumors may show recurrence even 8 to 10 years after initial complete response, which was seen in our case (42 years after first surgery and 32 years after recurrence). It necessitates stringent surveillance for long period even after complete response. Systematic review and meta-analysis have proven role of $^{131}$I-mIBG therapy in these tumors and combination of $^{131}$I-mIBG therapy with chemotherapy did not show any superior result, but an increased toxicity and hence not recommended.

There is no definite agreement on policy about dosage of $^{131}$I-mIBG in this group of patients; the review of literature shows varying dosage from 3.7 to 9.5 GBq. One of the widely used dosage schedules is median initial dose of 7.4 GBq (200 mCi) and median cumulative dose of 22.2 GBq with a range of 6.8 to 81.4 GBq depending on the case. Few investigators have also tried with escalated dosage of $^{131}$I-mIBG with a median single treatment dose of 30 GBq or 11.5 mCi/kg (range, 5.6–18.3 mCi/kg) that showed promising results but with significant toxicities. Multiple comorbidities including one-sided nephrectomy with renal compromise, hyperuricemia, and elderly age as presented here require personalized approach with rigorous workup throughout the therapies.

The excretion of $^{131}$I-mIBG is not well understood at the molecular level, but recent studies revealed that it is transported by organ transporter 2/toxin extrusion proteins hMATEs (multidrug and toxin extrusion proteins) pathway involved in active secretion. $^{131}$I-mIBG excretion rate over the first 24 hours is only partially related to the GFR. This makes $^{131}$I-mIBG quite safer in renal compromise, especially when compared with other radionuclide therapies such as PRRT ($^{177}$Lu/DOTATATE). Furthermore, $^{131}$I-mIBG is theoretically superior owing to the physical characteristics of harder $\beta$-emission of $^{131}$I compared with $^{177}$Lu ($E_{\beta_{\text{max}}} 0.6$ vs. 0.497 MeV). The impression of renal safety was reflected in our case where renal parameters remained in pretherapy ranges (► Fig. 6) without any special measure except dose adjustments (with a median dose of 5.55 GBq per cycle). The patient received a cumulative dose of 23 GBq in four cycles with a mean dose of 5.7 GBq per therapy cycle. At 24 months following the first cycle of $^{131}$I-mIBG therapy, she is asymptomatic with normal daily activities.

The hematological complications are most common side effects and generally recover spontaneously; the same
Fig. 3  The fourth pretherapy iodine-131 metaiodobenzylguanidine (131I-mIBG) scan (A) showing small foci of increased tracer uptake in the left paravertebral region noted in the bilateral lung fields. Post fourth cycle of 131I-mIBG therapy planar scan (B) with multiple foci of increased tracer uptake noted in the bilateral lung fields and midabdomen (note reduced sites and intensity of uptake compared with first posttherapy scans in Fig. 1). (C) Post-131I-mIBG therapy anterior planar scans of all four cycles with multiple foci of increased tracer uptake noted in the bilateral lung fields and midabdomen (note reducing sites and intensity of uptake with each cycle).

Fig. 4  Line graph showing weight in kg over the period of therapy and during follow-up.
findings got reflected in our case, showing typical pattern of deranging hemograms at every cycle, especially platelets (also hemoglobin) and spontaneous correction within a specific time period of 6 to 10 weeks (Fig. 7).

The goal of each therapy and cycle should be even clearly defined and plays a pivotal role in individualizing the $^{131}$I-mIBG therapy. The important considerations during dose fractionation in $^{131}$I-mIBG therapy should be tumor biology,
disease burden, and functional symptoms, thereby determining the intent of therapy. In the above-mentioned case with highly functional, progressive symptomatic and metastatic disease in an elderly patient with renal compromise, early symptom control and disease stabilization were achieved with repetitive treatments with smaller doses. Further studies with larger number of cases with dose adjustment and tolerability guidance, especially in deranged renal, hematological, and hepatic functional status, are merited.

Conclusion

Thus, in this case report, the authors describe the use of $^{131}$I-mIBG imaging and therapy to treat recurrent metastatic PCC. Taking into account the aims of treatment in a patient of advanced years with complicating factors, the patient was treated with four cycles of $^{131}$I-mIBG, resulting in clinical response and diminution of symptoms at up to 6 months after the fourth cycle. Dose was determined based on imaging data and patient-specific comorbidity factors and was at the lower range of the published range (23GBq compared with 6.8–81.4 GBq). The report is intended to provide some guidance for future therapeutic regimens for systemic radionuclide therapies in patient-specific manner.

Declaration of Patient Consent

The authors certify that they have obtained all appropriate patient consent forms.

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Nil.

Conflict of Interest

None declared.

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