External dose assessment from the patients treated by $^{177}$Lu-DOTATATE

E. Mahmoudi¹, M. Amoui², M.R. Deevband¹*, E. Pirayesh³, M. Ghorbani Rad³

¹Biomedical Engineering and Medical Physics Department, School of Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran
²Nuclear Medicine Department, Shohada_e Tajrish Hospital, School of Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran
³Nuclear Medicine Department, Shohada_e Tajrish Hospital, Tehran, Iran

► Short report

ABSTRACT

Background: The present study was done to establish release criterion in the treated patients and to determine external radiation doses received by personnel and caregivers of patients with metastatic neuroendocrine tumors (NETs) during peptide receptor radionuclide therapy (PRRT) by means of Lutetium-$^{177}$Lu DOTATATE.

Materials and Methods: For this purpose, 30 patients were enrolled in the study who received 5.5±1.1 (in a range of: 3.7-7.4) GBq of $^{177}$Lu-DOTA-tyr³-octreotate. Dose rate was analyzed at distances 0, 0.25, 0.5, 1.0, and 2.0 m in different times after termination of infusion using an ionization chamber. Results: Mean dose rate at distance of 1 m from the patient, approximately 5 h after injection was considered as discharge limit. The maximum dose to caregivers in the first 48 h was equal to 340±29 µSv. Mean dose to the nurse was estimated as 6.3±0.4, and 7±0.5 µSv per patient with and without lead shield, respectively. Discussion: According to our findings, approximately a time delay of 5 h after injection is recommended as release criterion for patients treated by Lutetium-$^{177}$Lu DOTATATE. For a total of 30 patients, external radiation dose to staff was found to be within permissible levels. Conclusion: The use of protective equipment is recommended at all stages of procedure for staff.

INTRODUCTION

One of the most important radionuclides in the field of therapy with nuclear medicine is Lu-177, which is a new and promising tool for control of non-operable metastatic neuroendocrine tumor when it is combined with somatostatin analogues (1-3). Currently most centers offering $^{177}$Lu-DOTA-tyr³-octreotate treatment perform it only in hospitals; because isolation and hospital admission are among controversial subjects. Conversely, in some countries, patients are discharged within a few hours after treatment when a determined outpatient’s release criterion is achieved according to local regulations (for example 30 µSv/h in Turkey and 25 µSv/h in Australia) (4-5). Since, treatment with $^{177}$Lu-DOTA-tyr³-octreotate is costly, additional costs of hospitalization are incurred by many patients. Long-term isolation in the hospital may also cause emotional disturbances to the patient. The use of different types of shield is questionable due to production of specific X-rays of lead following collision of high-energy gamma rays (6). However, various reports on methods of reducing the dose received by staff have been published by some national and international organizations (7). Therefore, the present study is designed to firstly focus on quantification of the caregiver’s mean dose, and secondly establishment of release criterion for patients treated with $^{177}$Lu-DOTATATE.

MATERIALS AND METHODS

The current study was approved by the Ethics Committee of the University (IR.SBMU.MSP.REC.1400.292), and an informed written consent was obtained from all the patients then, it was carried out according to provisions of the Declaration of Helsinki. Inclusion criteria were having over 35 years of age, being diagnosed with metastatic neuroendocrine tumor, and being candidate to be treated with $^{177}$Lu-DOTA-tyr³-octreotate. Mean age of patients was equal to 52.6 years (in a range of: 38-60 years) old. A total of 30 patients undergoing treatment with $^{177}$Lu-DOTA-tyr³-octreotate, were enrolled in the current study from March to August 2019. Patients were admitted in Department of Nuclear Medicine, Shohada_e Tajrish Hospital, Tehran Province, Iran. Infusion of 1,500-2,000 mL of normal...
saline and amino acids, mixture of 5% lysine HCl (50 mg) and 10% L-arginine HCl (50 mg) was carried out for 4 h to reduce radiation exposure of kidneys and subsequent adverse effects. The procedure was started 30 min after administration of 5.5 ± 1.1 GBq (in a range of: 3.7-7.4 GBq) of $^{177}$Lu-DOTATATE. All the administered patients were positioned in an isolated room with an area about 30 m². The room included 4 beds located in 4 corners of the room, with about 2 m of distance from each other.

### Dose rate measurement

The dose limit recommended by European guidelines for discharge of patients after iodine-131 therapy was set as the basis for discharge (20 µSv/h at 1 meter) (8-9). Using equation (1), cumulative dose can be estimated, $E$, to a caregiver standing from the patient for an unlimited time, assuming that only physical decay occurs. It was assumed that rate of initial dose is $D_0 = 20 \text{ µSv/h}$ at 1 m of distance, with half-life of $^{177}$Lu, 6.7 days, which is represented by $t_{1/2}$. Following calculation it was found that $E = 4.6 \text{ mSv}^{(10)}$.

$$E = \int_0^\infty D_0 \times e^{-\ln(2) \cdot \frac{t}{t_{1/2}}} \, dt \quad (1)$$

This study was carried out using an ionization chamber (Thermo, FH 40G-L10, made in Germany) calibrated by the secondary standard dosimetry laboratory (SSDL). Energy response of the dosimeter is equal to 30 keV-4.4 MeV and it has the capability to measure dose rate in the range of 10 nSv/h-100 mSv/h. Dose rate was measured on chest position at distances of 0, 0.25, 0.5, 1, and 2 m from the patient who received 5.5 ± 1.1 (in a range of : 3.7-7.4) GBq of $^{177}$Lu-DOTA-tyr³-octreotate with and without 2-mm lead shield after the mentioned time (0, 1 h, 2 h, 3 h, 4 h, 5 h, 18 h and 24 h). Mean dose to staff was estimated by recording the time interval and dose rate at the mentioned distances from the patient (11). Finally, mean dose to staff and related SDs were calculated based on µSv/GBq.h. The demographic information of staffs included in the current study is presented in table 1.

### Determining behavioral pattern for caregivers

Authors defined a pattern of behavior for the patient’s caregivers (a member of the family). Due to different conditions of patients, caregivers were categorized into three groups. The details are given in table 2. Caregivers’ mean dose was determined according to their behavioral pattern, duration of time spent with the patient, and measured dose rate.

### Table 2. Caregivers’ Behavior Pattern in different groups.

<table>
<thead>
<tr>
<th>Number of group</th>
<th>Pattern of behavior</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First group</strong></td>
<td>The patient was able to perform individual activities The patient was kept in a separate room and used a separate toilet The patient slept alone The caregiver visited his patient on average 5-10 minutes for the first 5 hours</td>
</tr>
<tr>
<td><strong>Second group</strong></td>
<td>The patient was able to perform individual activities The caregiver visited his patient on average 10-15 minutes for the first 5 hours The patient was slept 2 meters of the patient The patient and caregiver used a shared vehicle to drive</td>
</tr>
<tr>
<td><strong>Third group</strong></td>
<td>The patient was not able to perform any personal tasks after treatment The caregiver was present during the first 5 hours of treatment at the closest distance to the patient The caregiver was slept 1 meters of the patient The patient and caregiver used a shared vehicle to drive</td>
</tr>
</tbody>
</table>

### Statistical analysis

Data processing and fitting were performed using Microsoft Excel (Microsoft office professional plus, 2013) and SPSS (ver. 16.0, IBM Corp.) softwares were used for statistical analysis. For this purpose, the K-S (Kolmogorov-Smirnov) test was used to investigate normal distribution of data. A value of p-value of ≤0.05 was assumed as statistically significance. Data were presented as mean and standard deviation unless stated otherwise.

### RESULTS

Mean dose rate at a distance of 1 m from the patients treated with $^{177}$Lu-DOTA-tyr³-octreotate in different times after administration was measured and results are presented in figure 1. As shown in figure 1, dose rate was gradually decreased due to excretion of activity from the body. According to figure 1 for $^{177}$Lu-DOTA-tyr³-octreotate therapy, equation (2) was obtained from the curve, which was fitted to the data.

$$y = 28.442 \cdot e^{-0.067x} \quad (2)$$

The x and y indicates the time (h) and dose rate (µSv/h), respectively. Mean dose rate at a distance of 1m from the patient, approximately 5 h after the injection was measured as the discharge criterion.

According to the results of current study, mean
dose rate (µSv/GBq·h) at a close distance to a patient was obtained equal to 13.6 (SD=1.2) at 0 m, 8.9 (SD=1.2) at 0.25 m, 4.1 (SD=0.6) at 0.5 m, 1.3 (SD=0.2) at 1 m, and 0.6 (SD=0.2) at 2 m.

Dose to staff with and without lead shield

The mean, minimum, and maximum doses to staff in charge of caring the patients under treatment by $^{177}$Lu-DOTA-tyr$^3$-octreotate, were measured according to the time and distance from the patient with and without lead shield. The results are summarized in table 3. The annual mean dose of the staff, the distance and duration of time, in which they were in direct contact with the patient during each treatment period are listed in table 4. About 240 treatment sessions are performed annually at the Nuclear Medicine Center in shohada-e Tajrish Hospital (Tehran, Iran). The annual mean dose for staff in all treatment cycles was evaluated (patients selected for study as well as excluded patients). The annual mean dose received by the nurse was higher than all staff, which was approximately by 1.80 mSv. When lead shield is used, this value is estimated as 1.4 mSv. This value is estimated to be 1.7 and 0.8 mSv, for staff in charge of radiopharmaceutical injection and imaging staff, respectively. The annual mean dose was 1.3 and 0.6 mSv in case of not using and using lead-shield, respectively. Physician and physicist have the lowest annual dose received with and without using lead shield, by 0.5 and 0.6 mSv, respectively. The calculations were done based on the number of treatment sessions performed over a year assuming that nuclear medicine personnel participate in all treatment sessions.

#### Table 3. Mean, minimum and maximum dose (µSv per patient) of the staff during PRRT with $^{177}$Lu-DOTATATE, without and with the use of lead shield.

<table>
<thead>
<tr>
<th>Staff</th>
<th>Without lead shield</th>
<th>With lead shield</th>
<th>Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min dose per patient (µSv)</td>
<td>Max dose per patient (µSv)</td>
<td>Mean ± SD</td>
</tr>
<tr>
<td>Staff in charge of radiopharmaceutical injection</td>
<td>6.1</td>
<td>8.0</td>
<td>6.5±0.5</td>
</tr>
<tr>
<td>Staff in charge of imaging</td>
<td>3.0</td>
<td>4.0</td>
<td>3.3±0.5</td>
</tr>
<tr>
<td>Physician</td>
<td>2.0</td>
<td>3.2</td>
<td>2.5±0.4</td>
</tr>
<tr>
<td>Physicist</td>
<td>2.2</td>
<td>3.0</td>
<td>2.7±0.2</td>
</tr>
<tr>
<td>Nurse</td>
<td>6.5</td>
<td>8.5</td>
<td>7.5±0.5</td>
</tr>
</tbody>
</table>

#### Table 4. Estimated annual mean dose to staff in treatment by $^{177}$Lu-DOTATATE.

<table>
<thead>
<tr>
<th>Staff</th>
<th>Mean annual dose using lead shield (mSv)</th>
<th>Mean annual dose without lead shield (mSv)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staff in charge of radiopharmaceutical injection</td>
<td>1.3</td>
<td>1.7</td>
</tr>
<tr>
<td>Staff in charge of imaging</td>
<td>0.6</td>
<td>0.8</td>
</tr>
<tr>
<td>Physician</td>
<td>0.5</td>
<td>0.6</td>
</tr>
<tr>
<td>Physicist</td>
<td>0.5</td>
<td>0.6</td>
</tr>
<tr>
<td>Nurse</td>
<td>1.4</td>
<td>1.8</td>
</tr>
</tbody>
</table>

Dose to caregivers

Mean dose to 30 caregivers was found to be within a range of 36-390 µSv in $^{177}$Lu-DOTA-tyr$^3$-octreotate therapy after infusion of 5.5±1.1 (in a range of: 3.7-7.4) GBq. Mean dose of the first caregiver group was estimated to be 47.3 ± 8.4 µSv (in a range of: 36-60 µSv). Mean dose of the second group was equal to 184 ± 29 µSv (in a range of: 150-220 µSv). The third group had the highest mean dose, by 340.5±29 µSv (in a range of: 300-390 µSv).

DISCUSSION

Two important features of clinical treatment are cost-effectiveness and availability of treatment. Many nuclear medicine centers are not able to offer this treatment because of lack of facilities for hospitalization, increasing waiting time for patients to receive $^{177}$Lu-DOTA-tyr$^3$-octreotate treatment. Based on results of the current study presented in figure 1, mean dose rate at a distance of 1m from the patient, approximately 5 h after the injection was considered to be lower than the discharge limit. According to figure 1, equation (2) would also be used to achieve release criterion which was defined less than 20 µSv/h at a distance of 1m, and can be considered as a dose limit for discharge from the patients treated with $^{177}$Lu-DOTA-tyr$^3$-octreotate. In a similar study on iodine therapy, Ahmadi

![Figure 1. External dose rate (µSv/h) for patients treated with $^{177}$Lu-DOTATATE therapy at 1 m distance.](image-url)

\[ y = 28.422e^{-0.66x} \]

\[ R^2 = 0.8045 \]
Jeshvaghane et al., (9) showed that the maximum and minimum doses after release were equal to 21 (SD-18) and 11 (SD-4.0) µSv/h at a distance of 1 m from the patients, respectively.

Mean dose for caregiver in three groups was measured below the level recommended by the International Commission on Radiological Protection (ICRP) for each patient [5 mSv] in each treatment period (10). The amount of activity received by the patients and duration of the caregiver’s presence at close distances to the patient are important parameters regarding the caregiver’s mean dose. Abuqbeitah et al., (8) in a study reported the limit of 20 µSv/h for hospital discharge. They estimated a dose rate of 100-200 µSv for caregiver of the patients receiving 177Lu-DOTA-tyr3-octreotate treatment (8), Calais et al., (5) estimated the mean total dose to 25 caring sessions during day of therapy and they reported a dose rate of 10-470 µSv for surrounding people while taking the patient to home within a period up to 5 days after treatment. However, some nuclear medicine clinics prefer to admit patients in hospital to monitor the probable side effects. In a study by Kurt et al., (12), patients were admitted in two centers to be hospitalized for 48 and 72h. The maximum dose to individual members in the public per treatment cycle was ~ 250 ± 55 and ~ 190 ± 36 µSv when the patients were discharged after 48 and 72 h, respectively. But our findings, showed no need for intensive radiation monitoring of caregivers who had a distance more than 1 m from the treated patients. Abuqbeitah et al., (8) showed that the maximum mean dose received by radiopharmaceutical injection was equal to 4 ± 1.9 µSv per patient. Calais et al., (5) reported the maximum mean staff dose (of 33 µSv) for the nurse. Results of the current study showed the maximum mean dose of 7.5 µSv per patient for nurse. Table 5 presents comparison of the results for staff. The nurse’s high mean dose is due to the fact that the nurse has the most contact with the patient during treatment. The difference between results of studies may be due to experience, skills, and promptness of the staff. No measured dose to hospital staff or family members exceeded the limits.

**Table 5.** Comparison of the results of current study with international results.

<table>
<thead>
<tr>
<th>Staff in charge of patient imaging</th>
<th>Abuqbeitah et al. (8) µSv Per therapy treatment day with one patient</th>
<th>Calais et al. (5) µSv Per therapy treatment day with four patients</th>
<th>Current study µSv Per therapy treatment day with one patient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staff in charge of patient imaging</td>
<td>3.0 ± 0.9</td>
<td>7.0 ± 0.5</td>
<td>3.3 ± 0.5</td>
</tr>
<tr>
<td>radiopharmaceutical</td>
<td>4.0 ± 1.9</td>
<td>17.0</td>
<td>6.5 ± 0.5</td>
</tr>
<tr>
<td>Physician</td>
<td>2.0 ± 0.8</td>
<td>9.0</td>
<td>2.7 ± 0.2</td>
</tr>
<tr>
<td>Nurse</td>
<td>5.0 ± 1.1</td>
<td>33.0</td>
<td>7.5 ± 0.5</td>
</tr>
</tbody>
</table>

**LIMITATION**

The limitation of this study was low number of patients included in the study. Also there were no lead and syringe shields of varying thicknesses for general assessing the effect of protective equipment on the received dose.

**CONCLUSION**

Results of the present study showed that the dose rate from the injected patients was decreased to lower than the specified threshold of 20 µSv/h at a distance of 1 m after approximately 5 h. which was considered as release criterion for patients treated with 177Lu-DOTA-tyr3-octreotate. Due to the effect of lead shield on reducing staffs mean dose, it is recommended that protective device should be used in all treatment stages. In summary, no measured mean dose to hospital personnel, caregivers, family or member of the public exceeded the annual related dose limits.

**Ethical consideration:** Provide The current study was approved by the Ethics Committee of the University (IR.SBMUMSP.REC.1400.292).

**Declaration of Interests:** Declared none.

**Funding:** The present research is financially supported by research department of the school of medicine, Shahid Beheshti University of Medical Sciences (Grant No. 25569).

**Author contribution:** (E.M): Contribution to the conception and design of the work, acquisition, collection of data, measurements, analysis, interpretation of data for the work and drafting the report. (M.A): Contribution to the conception of the work, interpretation of data, revising the report and final approval of the version to be published. (M.R.D): Contribution to the conception and design of the work, interpretation of data, revising the report and final approval of the version to be published as corresponding author. (E.P): Contribution to the conception of the work, interpretation of data, revising the report and final approval of the version to be published. (M.Gh) Rad: Contribution to the collection of data and measurements.

**REFERENCES**
