Evaluating the Efficacy of Bone Tracer $^{99m}$Tc-MDP in Detecting Individual Renal Function Compared to Renal Cortical Imaging Agent $^{99m}$Tc-DMSA

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Abstract

This study was performed to evaluate the usefulness of bone tracer $^{99m}$Tc-MDP in detecting renal disorders by imaging of renal cortex for patients with cortical defect and estimation of split renal function (SRF) as compared with renal cortical imaging agent $^{99m}$Tc-DMSA. In this study 23 patients (6 males, 17 females) with range age of 16y -75y (mean 45.6y ± 17.4y), and a total of 46 studies (using both $^{99m}$Tc-DMSA and $^{99m}$Tc-MDP per patient for left and right kidneys) were performed for detecting renal disorders. Both studies were performed within 48 hours between each other. SRF was calculated in all cases from renal uptake of the tracers into the region of interest by using computer acquired data of gamma camera. For detecting renal cortex abnormalities, all cortical scars and defects scans seen on $^{99m}$Tc-DMSA scan were also detected by $^{99m}$Tc-MDP.

The results showed that the non visualized kidney on $^{99m}$Tc-DMSA scan was also non visualized on $^{99m}$Tc-MDP. Also, there was a high correlation between SRF values obtained from $^{99m}$Tc-MDP and $^{99m}$Tc-DMSA, where (p>0.01, r= 0.93 for left kidney and r=0.0.92 for right kidney).

The similar SRF values of $^{99m}$Tc-MDP and $^{99m}$Tc-DMSA allow to use of $^{99m}$Tc-MDP tracers for detecting both of bone and renal abnormalities.

Keywords: $^{99m}$Tc-MDP; $^{99m}$Tc-DMSA; Renal cortex; Split renal function

Introduction

Estimation of split renal function is a particularly important role of renal scintigraphy [1] and one of the major contributions of radionuclide renal studies to the practice of urology [2]. Functional and morphologic investigations with radionuclide studies play a prominent role in the diagnosis and follow-up of various renal disorders [3-7]. $^{99m}$Tc- DMSA is slowly cleared from the blood and concentrates in the renal cortex; 42 % of the injected dose remains in the renal cortex at 6 hours [8]. DMSA is an excellent agent for detecting focal abnormalities of the renal cortex. Because of its high kidney uptake, it has been suggested that $^{99m}$Tc-DMSA may be the best technetium agent for determining the relative functional renal mass. Technetium-$^{99m}$dimercaptosuccinic acid ($^{99m}$Tc-DMSA) was introduced in 1974 and is still the agent of choice for static renal scintigraphy [9,10], and $^{99m}$Tc MDP(methylene diphosphate) labeled phosphate complexes were first introduced in 1971 as a major advance in skeletal image. Bone scintigraphy, so far, is still a useful examination for the clinical diagnosis, especially in evaluating and following up the status of cancer patients with suspicious bony metastasis. Many authors [10-14] found that there are a number of renal disorders detected while doing bone scintigraphy because $^{99m}$Tc-MDP (is excreted through the kidneys to provide adequate visualization of the urinary tract. Furthermore, damage to the kidney is caused by chemotherapy and/ or radiation therapy when the kidneys are included in the radiation field [15]. Kirkinen [16] and other investigators [17] indicated that further that urethral obstruction from radiation therapy might occur as early as three weeks or as late several years after therapy, it was potentially curable. Therefore, the estimation of renal function as well as the detection of bone metastases is useful in patients that received radiation therapy for detecting undermine renal disorders, and prevent those with high risk of renal abnormalities from going to irreversible stage.

Material and Methods

Subject

A total of 46 studies for 23 patients (within ages ranged from 16 years-75years old) were performed between 2014 and 2015, in Nuclear Medicine division–Radiology department at Zagazig University Hospital, (Egypt). Because of various renal disorders and for routine indications, such as evaluation of renal cortex after urinary tract infection, the split renal function assessment (SRFs) were calculated using both $^{99m}$Tc-MDP and $^{99m}$Tc-DMSA renal scintigraphy. All studies were performed using both radionuclides for each patient to evaluate the usefulness of bone seeking radiopharmaceutical in detecting renal abnormalities as compared with renal cortical agent. All radionuclide studies were carried out using a Dual-headed gamma camera equipped with a low-energy, high-resolution parallel-hole collimator (GE Healthcare Unveils Discovery NM 630 SPECT).
During imaging, the collimator was set as close as possible to the patient’s table.

**99mTc-DMSA scintigraphy**

There is no preparation for patients during a DMSA Scan, they can eat and drink normally. The radiopharmaceutical was prepared according to the manufacturer’s instruction with the kit. Patients were injected with activity 5.0 mci of the radiopharmaceutical followed by infusion of 20 ml of normal saline. After 2-3 hours intravenous injection, 99mTc-DMSA static images were acquired in 256 x 256 matrix with the patients in a supine position that appears the best position to minimize renal depth difference, thus improving the accuracy of split renal function measurement and with gamma camera’s detectors placed in a posterior and anterior views (250 kcounts/view or 5 minutes/view). SRFs were calculated, semi-automatically, by the geometrical mean of the anterior and posterior images, using GE Xeleris software.

**99mTc-MDP scintigraphy**

99mTc-MDP Static images were performed on another day, for comparison with 99mTc-DMSA Static images for the same patients. Data collection and analysis were repeated under the same conditions, and SRFs for all patients were calculated by the same method.

**Data Analysis**

Using the posterior and anterior digital images, regions of interest are placed around both kidneys and below both kidneys for background subtraction. Then, the background corrected counts in each kidney and the percent of total counts (DMSA uptake) in each kidney are calculated. Cross-correlation coefficients were calculated by Pearson correlation for comparison of repeated measurements of SRF for both kidneys using both radiopharmaceuticals. Statistical analysis was performed using the SPSS 14.0 software program.

**Results**

SRFs measured by 99mTc-MDP and 99mTc-DMSA renal scintigraphy are shown in Table 1. In this study, there are 9 patients had normal SRF (45%-55%) as shown in Figure 1, and 14 patients had abnormal SRF, and non visualized kidneys on 99mTc-DMSA scan were also non visualized on 99mTc-MDP. For detecting renal cortex abnormalities, all cortical scars and defects scars seen on 99mTc-DMSA scan were also detected by 99mTc-MDP as shown in Figures 2-4. In correlation analysis, there was a high correlation between the SRF obtained from the 99mTc-DMSA and 99mTc-MDP for both left and right kidneys (p>0.01 and r=0.93, r=0.92 left and right kidney, respectively) as shown in Figures 5 and 6.

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**Table 1**: SRFs measured by 99mTc-MDP and 99mTc-DMSA renal scintigraphy.
Figure 2: Abnormal study for female with age 22 years with left cortical defects seen on both studies. A: SRF derived from static 99mTc-MDP scintigraphy: Left/Right_2/98%. B: SRF derived from (2-3 hours) static 99mTc-DMSA scintigraphy: Left/Right_1/9%.

Figure 3: Abnormal study for female with age 34 years with left cortical defects seen on both studies. A: SRF derived from static 99mTc-MDP scintigraphy: Left/Right_8/92%. B: SRF derived from (2-3 hours) static 99mTc-DMSA scintigraphy: Left/Right_8/92%.

Figure 4: Abnormal study for male with age 19 years with left cortical defects seen on both studies. A: SRF derived from static 99mTc-MDP scintigraphy: Left/Right_6/94%. B: SRF derived from (2-3 hours) static 99mTc-DMSA scintigraphy: Left/Right_13/87%.

Figure 5: Correlation between left Split Renal Function (SRF) using 99mTc-MDP versus 99mTc-DMSA.

Figure 6: Correlation between Right Split Renal Function (SRF) using 99mTc-MDP versus 99mTc-DMSA.
Discussion

99mTc-DMSA scanning has been regarded as the best method for assessing SRF because the radiotracer is retained [18], primarily in the proximal convoluted tubules, for a sufficiently long time to allow static imaging of tubular activity which can be performed over several minutes [19]. It is well known that information about renal abnormalities can be obtained from bone scintigraphy because 99mTc-MDP is excreted through the kidneys to provide adequate visualization of the urinary tract [20]. These abnormalities have included absent renal activity, small or displaced kidneys, urinary obstruction, focal renal parenchymal abnormalities, unilateral decrease in renal function, and asymmetric uptake [10-14]. In addition, detailed views of bladder can be obtained. Based on the data of animal and human studies, 99mTc-MDP, unlike chelates (e.g. 99mTc-DTPA) which are handled by glomerular filtration [20,21], is probably excreted as simple phosphates by renal tubular [22]. Thus, accompany with sufficient excretory amount of MDP, it provides excellent images on kidney at the time of bone scanning. So 99mTc-MDP and 99mTc-DMSA examinations should be performed on different days.

Technetium-99m DMSA scintigraphy is the gold standard for evaluation of split renal function (SRF) [23-25]. 99mTc-dimercaptosuccinic acid (99mTc-DMSA) is an agent that is actively taken up by the proximal and distal renal tubular cells, directly from per tubular vessels, not secreted to the tubular lumen [26] and accumulates in the renal cortex [27]. This modality is primarily used for imaging functioning cortical mass and individual renal function [28] since the abnormal kidney function was set at <45% for one kidney [29]. It is the most reliable method for assessing chronic cortical scarring [19]. In the 99mTc-DMSA studies, SRF is calculated using the geometric mean and taking kidney depth into account by

Conclusion

The result confirmed that 99mTc-MDP scintigraphy can simultaneously estimate split renal function and skeletal lesions, it is valuable for saving time, effort and money because the patients have 99mTc-MDP scintigraphy did not need another renal examination if the individual renal function is the clinical question but assessment of global renal function as well as GFR cannot evaluated with these modalities and further evaluation with more specific renal imaging agents is recommended.

Compliance with Ethical Standards

Funding: This study was funded by the authors.

Conflict of Interest

There is no any conflict interest among the authors.

Ethical Approval

All procedures performed in studies involving human participants were in accordance with the ethical standards of the national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed Consent

Informed consent was obtained from all individual participants included in this study.

References


