Tc-99m Ethylenedicysteine and Tc-99m Dimercaptosuccinic Acid Scintigraphy-Comparison of the Two for Detection of Scarring and Differential Cortical Function

Abstract
Context: The differential cortical function obtained by Tc-99m EC is comparable to that of Tc-99m DMSA. However, identification of scars on Tc-99m EC images needs to be studied. Aims: The aim of the study is to evaluate role of Tc-99m EC for detection of scarring and differential cortical function by comparing with Tc-99m DMSA. Settings and Design: Prospective observational study of recurrent UTI; minimum 6 weeks after acute episode; when urine examination is negative for pus cells. Materials and Methods: Forty-seven children with normal positioned kidneys underwent Tc-99m EC and DMSA scintigraphy. The DRF and cortical phase images of both studies in the same image matrix size were evaluated by two independent observers for scarring; Tc-99m DMSA was considered as the gold standard. Statistical analysis used: MS Excel 2007 and GraphPad Instat V3.1 and ROC analysis. Results: There was no significant difference in the detection of scarring using two studies with Cohen’s kappa coefficient (κ) 0.932. The sensitivity and specificity of Tc-99m EC for detection of scarring was 98.75% and 99.15%, respectively. There was good agreement between the differential cortical function calculated using two studies. Conclusions: The summed Tc-99m EC images with an acceptable high image contrast allow detection of cortical scarring in patients with normal kidney positions. It is an excellent single-modality comprehensive investigational agent for renal parenchymal defects, function, and excretion evaluation with the added advantages of lower cost, convenience, and low radiation exposure to the child.

Key Messages: As the differential renal function and cortical scarring detected using Tc-99m EC is comparable to the gold standard of Tc-99m DMSA; Tc-99m EC can be used as a single modality for obtaining multiple information reducing time, cost, and radiation exposure to the child.

Keywords: cortical scarring, differential renal function, DMSA, EC, UTI

Introduction
Complicated urinary tract infection (UTI) is defined on the basis of clinical and biological criteria, to treat this entity in a more aggressive way than in the case of simple cystitis, to prescribe a prophylactic treatment to protect the kidney from further deterioration and to treat dysfunctional bladders. The workup involves micturating cystourethrogram (MCU) to diagnose the grade of vesico-ureteric reflux (VUR). UTI may affect the urinary bladder (cystitis), the kidneys, and the collecting systems or both. Bacterial infection of the lower urinary tract can be asymptomatic and limited to urinary bladder. But the lower urinary tract infection has a risk of spreading to the kidneys. Infected urine stimulates immunologic and inflammatory response resulting in renal injury and scarring. With grade III, IV, or V VUR along with an episode of febrile UTI, there is 90% chance of acute pyelonephritis on renal scintigraphy.[1] The diagnosis of pyelonephritis and subsequent scarring of renal cortex is made on Tc-99m dimercaptosuccinic acid (DMSA) scanning and is routinely done in complicated UTI.[1,2]

Tc-99m mercapto acetyl triglycine (MAG₃) scintigraphy is indicated whenever the collecting systems are dilated on ultrasound to rule out obstruction. Tc-99m DMSA and MAG scans provide information about the differential renal function (DRF). The DRF provided by two studies has been found to be comparable. However, the detection of scarring on Tc-99m MAG₃ imaging during...
the cortical uptake phase (1-3 minutes) was variable in the studies performed.\textsuperscript{[3-7]}

Tc-99m-N,N-ethylendicysteine (EC) is another tubular secreted agent used for the renogram study. The greatest advantage of Tc-99mEC over Tc-99mMAG3 appears to be the negligible accumulation in the liver.\textsuperscript{[9]} Thus, it can be used in patients with high creatinine with better delineation of kidney cortex. The DRF and detection of scarring using this agent was compared with the same on Tc-99m DMSA. There was good agreement between the DRF obtained using two radiopharmaceuticals. However, the confidence level of detection of areas of scarring was variable.\textsuperscript{[9-14]}

The main disadvantage of Tc-99m DMSA is its slightly higher radiation dose in comparison with other renal agents because of tubular fixation of DMSA.\textsuperscript{[16-18]}

However, about 70% of EC is excreted from the cortex with no tubular fixation.\textsuperscript{[9]}

The techniques used for the Tc-99m EC and Tc-99m DMSA acquisitions are different, the first being a dynamic acquisition and the second being the static acquisition, causing difficulty in selection of comparable images of the cortical phase. The cortical scars are detected qualitatively and need good quality images to identify scars. Most of the studies performed in the past comparing the Tc-99m EC or MAG3 cortical phase images with Tc-99m DMSA images used variable acquisition parameters.

The purpose of the study was to compare Tc-99m EC cortical phase images to that of Tc-99m DMSA using the same image matrix size during acquisition. This exercise provided comparable image quality of Tc-99m EC to the same of Tc-99m DMSA to identify cortical scars.

Materials and Methods

Forty-seven children with mean age of 4.44 years (range 20 days to 10 years) were included in the prospective study conducted in the Department of Nuclear Medicine, over a period of 2 years from June 2014 to July 2016. All the children with urinary tract infection (UTI) with negative urine examinations for pus cells for minimum 6 weeks were included in the study. The children with known ectopic, horseshoe or mal-rotated kidneys, renal agenesis, post-renal transplant, and parents unwilling to sign informed consent were excluded. After obtaining ethical approval from the local ethics committee, an informed consent document was obtained from parents/guardians.

The two studies, i.e., Tc-99m DMSA and Tc-99m EC scintigraphy, were performed on two separate days, maximum 10 days apart. The Tc-99m EC dynamic images were acquired in posterior projection using the 128 × 128 image matrix size after administering a mean activity of 74 MBq/2 mCi (minimum of 15MBq/400 µCi), and Tc99m DMSA static images were acquired in anterior and posterior projection using 128 ×128 as well as the 256 × 256 image matrix 3 hours after intravenous injection of mean 55.5 MBq/1.5 mCi (minimum dose of 18.5MBq/500 µCi).\textsuperscript{[19,20]} The 2-3 minute dynamic images of EC were processed and reframed as 1 minute static cortical phase image set for comparison of cortical scarring with that of DMSA. The wedge shape defect in the cortex with distortion of the cortical contour was considered as the scar. The regions of interest (ROIs) around the kidney contours were drawn manually, and each was divided in three sub-regions (upper, median, and lower segments) for easier and accurate analysis [Figure 1a]. On visual analysis, a photopenic defect > 50% area in a segment was defined as the large scar and ≤50% of cortex in a segment was defined as the small scar. The renal cortical contour was also assessed visually to locate the distortion, which is an indicator of scarring. The Tc-99m DMSA images in 128 × 128 and 256 × 256 matrix were compared to detect scarring with a higher matrix size being considered as the gold standard. The Tc-99m EC and Tc-99m DMSA images in the 128 × 128 matrix size were compared with to identify cortical scars, and the same scarred areas were confirmed on 256 × 256 matrix size images of DMSA (considered as the gold standard). Both images of EC and DMSA were analyzed on a high-resolution monitor meant for gray and color scale presentations, with positive and negative variants. The images were reviewed independently by a two nuclear medicine physicians to eliminate bias due to inter-observer variability and obtain the receiver operator curve (ROC) analysis.

The DRF of both kidneys was calculated using Tc-99m EC
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The distorted cortical contour is representative of the scarring. In total, 42 kidneys had normal contour and 52 had distorted contour that was identified on Tc-99m EC images. There was no significant difference \((P = 0.76)\) noted in the characterization of contour by Tc-99m EC and Tc-99m DMSA imaging [Table 2].

The static images of Tc-99m DMSA acquired using the higher image matrix \((256 \times 256)\) was considered as the gold standard. The comparison of these images with the lower image matrix was considered to be equivalent by demonstrating no statically significant difference in detection of scarring in each segment of the renal cortex [Table 3].

When Tc-99m EC was compared to gold standard Tc-99m DMSA for detection of cortical scarring, the sensitivity was 98.75% (93.23-99.97%), specificity was 99.15% (95.33-99.98%), positive predictive value was 98.75% (93.23-99.97%), and negative predictive value was 99.15% (95.33-99.98%).

The ROC analysis of Tc-99m EC and Tc-99m DMSA images showed Cohen’s kappa coefficient \((\kappa)\) within the range of 0.81 to 1.0 suggesting almost perfect agreement between the two observers for detection of scarring by Tc-99m EC and Tc-99m DMSA imaging [Figure 3].

| Table 1: Comparison of detection of scarring in the image of two matrix sizes of Tc-99m DMSA |
|---------------------------------------------------------------|
| **No of scars**       | DMSA 128 \times 128 | DMSA 256 \times 256 | \(P\) value* |
|------------------------|---------------------|---------------------|--------------|
| \(\leq 50\%\)            | 20                  | 21                  | 27           | 0.76         |
| \(>50\%\)              | 28                  | 27                  |              |              |

*Chi square test-no significant difference between images of two matrix sizes

| Table 2: Comparison of delineation of contour by Tc-99m EC and Tc-99m DMSA |
|---------------------------------------------------------------|
| **No of kidneys**       | Contour | EC (128 \times 128) | DMSA (128 \times 128) | \(P\) value* |
|------------------------|---------|---------------------|----------------------|--------------|
| Distorted              | 54      | 52                  |                      | 0.76         |
| Normal                 | 40      | 42                  |                      |              |

*Chi square test-no significant difference noted in delineation of contour between two scintigraphy methods.

Figure 2: Comparison of scars using Tc-99m EC, Tc-99m DMSA 128 \times 128 image matrix size, and Tc-99m DMSA 256 \times 256 image matrix size: ≤ 50% scar at the upper pole of the right kidney (a, b, c red arrow), detection of multiple 550% and >50% size scars bilaterally (d, e, f), and identification of large scar involving entire right kidney (g, h, i blue arrow)

Figure 3: ROC curves for agreement between two independent observers for identification of scarring: upper segment (a), median segment (b), and lower segment (c) on the 128 \times 128 image matrix size of Tc-99m EC and Tc-99m DMSA and upper segment (d), median segment (e), and lower segment (f) on Tc-99m DMSA (128 \times 128) and Tc-99m DMSA (256 \times 256)

The results

The 47 patients evaluated for UTI had known posterior urethral valves (post-radiofrequency fulguration) in 12 patients, VUR (15 unilateral and 6 bilateral) in 21 patients, pyonephrosis in 7 patients, and recurrent UTI without dilated collecting system in 7. Out of 94 kidneys evaluated in 47 patients; 42 kidneys did not show evidence of scarring with normal contour on the Tc-99m DMSA images with a higher matrix.

Detection of scarring

In 47 patients, 94 kidneys were evaluated to detect the extent of scarring in upper, median, and lower segments of kidney [Figure 1b-Figure 1d; Figure 2]. There was no significant difference noted in the detection of extent of scars \((\leq 50\%\) and \(>50\%\)) using Tc-99m EC in each cortical segment when evaluated with Tc-99m DMSA using the same image matrix size [Table 1].
Comparison of DRF

The DRF calculated by the Tc-99m EC and Tc-99m DMSA for 47 patients and 94 kidneys was comparable as shown by no statistically significant difference (P = 0.99) between the two [Table 4].

Discussion

The DRFs calculated using Tc-99m DMSA and Tc-99m EC were almost similar. This finding was in agreement with the existing literature. Kibar M et al, Raja S et al, Atasever T et al and Buyukdereli G et al suggested that the DRF obtained by the Tc-99m EC and Tc-99m DMSA were comparable; however, Tc-99m DMSA was found to be gold standard for the identification of scarring. The Tc-99m EC cortical images were not enough for identifying scarring in these studies. The image acquisition parameters being used for Tc-99m EC were variable and not comparable with Tc-99m DMSA in these studies. The use of the lower image matrix size for Tc-99m EC scintigraphy reduced the resolution of images. Also, the use of pin hole collimator for acquisition of Tc-99m DMSA improved the resolution of image. The pinhole magnification imaging increased the level of certainty with which renal cortical scintigraphy was interpreted but do not significantly increased the number of studies whose findings were considered abnormal on the basis of high-resolution planar imaging.

The ROC analysis in this study showed excellent agreement between the two observers for identification of scarring. Thus, a simple modification during the acquisition of Tc-99m EC study with use of 128 × 128 image matrix size extended the use of single scintigraphy for obtaining multiple reliable information. The higher matrix size enhanced the image quality to detect the scars even in kidneys with hydronephrosis and impaired cortical function. Narayana et al also noted good agreement between Tc-99m EC and Tc-99m DMSA images for detection of scarring in the presence of raised creatinine and impaired renal cortical function in addition to the DRF. Thus, Tc-99m EC can be used as a preferred scintigraphy technique in UTI even with a dilated collecting system.

This study has evaluated kidneys positioned in the normal location in the clinical scenario of UTI. However, further studies are required for evaluation of scarring in ectopically located kidneys with hydronephrosis as well as impaired renal cortical function.

Conclusion

The DRF provided by Tc-99m EC was comparable with that provided by Tc-99m DMSA. There was excellent agreement between these two studies for detection of scarring in cases of UTI (average κ = 0.932) by two independent observers. The summed Tc-99m EC images with an acceptable high image contrast allowed detection of cortical scarring in patients with kidney in normal anatomical positions. Thus, Tc-99m EC was found to be an excellent single-modality comprehensive investigational agent for renal parenchymal defects, function, and outflow tract evaluation with the added advantages of lower cost, convenience, and low radiation exposure to the child.

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Conflicts of interest

There are no conflicts of interest

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| Extent of scarring | EC (128 × 128 matrix) | DMSA (128 × 128 matrix) | P value* |
|-------------------|-----------------------|--------------------------|----------|
| Upper segment     | ≤50%                  | 21                       | 20       | 0.98     |
|                   | >50%                  | 28                       | 28       | 1        |
| Median segment    | ≤50%                  | 17                       | 15       | 0.88     |
|                   | >50%                  | 18                       | 20       | 0.87     |
| Lower segment     | ≤50%                  | 17                       | 18       | 0.76     |
|                   | >50%                  | 28                       | 25       | 0.88     |

*χ² test-no significant difference noted between the two tests

| DRF | EC | DMSA | P value* |
|-----|----|------|----------|
| Mean| 29.02 | 46.22 |          |
| SD  | 30.22 | 26.72 | 0.99     |

*paired t test-no significant difference noted between the two tests

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