Comparative study of Apparent Diffusion Coefficient Values and Ultrasonography in Evaluation of Adult Chronic Kidney Disease Patients

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ABSTRACT

Introduction: Diffusion-weighted magnetic resonance imaging (DWI) is a non-invasive method sensitive to water motion in the tissue. Diffusion weighted magnetic resonance imaging (DW-MRI) in evaluation of renal diseases is an evolving field and its utility is to be understood and it has the potential to become an effective imaging biomarker. Study objective was to evaluate the role of diffusion weighted MRI in staging chronic kidney disease by apparent diffusion coefficient values and correlating ADC values with CKD staging and Ultrasound grading.

Material and Methods: A prospective study done in adult patients with chronic kidney disease from January 2019 to August 2019. A total of 50 cases were included in the study. Serum markers of renal failure were noted, CKD staging was done by calculating eGFR using MDRD formula followed by Ultrasound evaluation for parenchymal Grading. Later, MRI was done and ADC values of parenchyma were determined. CKD staging and ultrasound grading of parenchyma was correlated with ADC values of parenchyma.

Results: In our study, out of total 50 cases, majority number of cases were seen in male population (64%), with younger agegroup (21 – 30 years) predominance -16 cases (32%). Clinically based on eGFR calculation, CKD stage I cases were 4(8%), Stage II – 2(4%), Stage III – 8(16%), Stage IV-14(28%) and stage V cases 22(44%). On ultrasound grading, Grade I cases - 10(20%), Grade II cases -16(32%), Grade III cases – 14(28%) and Grade IV cases-10(20%). ADC values of stage IV and V were correlating with CKD staging and Ultrasound Grading (90 to 100%) whereas clinically CKD stage I and Stage II, and USG grade I and II cases ADC values are not correlating as most of cases were showing Stage III ADC values. Linear correlation was seen with ADC values and eGFR and inverse correlation was seen with serum markers.

Conclusion: ADC values of renal parenchyma is a useful tool in evaluating renal dysfunction, which will make MR imaging of kidney as one step modality for renal evaluation.

Keywords: Apparent Diffusion Coefficient, Chronic Kidney Disease, Ultrasound, eGFR.

INTRODUCTION

Chronic kidney disease is recognised as world wide leading health problem. Serum markers such as creatinine and blood urea nitrogen level and estimated glomerular filtration rate (eGFR) are useful parameters for estimating renal function, however they depend on age and body mass index of patient and cannot be used to evaluate single kidney function.¹² Hence because of these limitations, imaging techniques such as ultrasonography, diffusion weighted magnetic resonance imaging (DW-MRI) are gaining importance in evaluation of renal function, Though ultrasonography is preferred modality, the lack of specificity in assessing renal disease emphasises on using better imaging modality. The apparent diffusion coefficient is a quantitative parameter calculated from DW-MRI images and represents water diffusion in extracellular and extra vascular space and capillary perfusion, Diffusion weighted imaging (DWI) in renal diseases is an evolving field and many investigators used it to characterize renal parenchymal disease.³⁴⁵⁶⁷⁸ The purpose of this study is to determine the apparent diffusion coefficient (ADC) values of renal parenchyma and its relation with ultrasonographic grading of CKD, serum.
markers of renal function and staging of chronic kidney disease.

**MATERIAL AND METHODS**

This was the prospective single institution study approved by institute ethical committee and was done after informed consent was taken from patients. Our study was conducted from January 2019 to August 2019. The total number of patients included were 50, with male predominance (64% male 36% female) with a mean age group of 42 years. For eGFR calculation, patients demographic data and serum investigations were collected from records.

**CKD grading**

eGFR was calculated from MDRD formula using body surface area, age, sex, race, serum creatinine level Formula

\[ \text{eGFR} = \frac{194 \times \text{creatinine} - 1.094 \times \text{age} - 0.287 (0.739 \text{ if female})}{1.73} \]

Patients were classified into 5 stages with eGFR value.

- **Stage 1**: eGFR ≥ 90 mL/min/1.73 m² (kidney damage with normal or increased eGFR)
- **Stage 2**: eGFR 60–89 mL/min/1.73 m² (kidney damage with a mild reduction in eGFR).
- **Stage 3**: eGFR 30–59 mL/min/1.73 m² (moderate reduction in eGFR).
- **Stage 4**: eGFR 15–29 mL/min/1.73 m² (severe reduction in eGFR).
- **Stage 5**: eGFR < 15 mL/min/1.73 m² (kidney failure).

**Ultrasoundography**

Ultrasound was done on GE voluson P5 and renal parenchymal echogenicity was evaluated. Grading of renal parenchymal echogenicity was done.

- **Grade 1** - Normal sized kidney, cortical echogenicity is same as liver, with well maintained cortico-medullary differentiation.
- **Grade 2** - Normal sized kidney, cortical echogenicity is greater than that of liver, with maintained cortico-medullary differentiation.
- **Grade 3** - Normal sized kidney, cortical echogenicity is more than that of liver, decreased cortico-medullary differentiation.
- **Grade 4** - Reduced renal length, cortical echogenicity is more than that of liver, with poorly maintained cortico-medullary differentiation.

**MRI**

All the patients underwent MRI on 1.5 T (semensavanto Germany) using phased array body coil. Image protocols included were T1 axial, T2 axial, coronal, sagittal planes. DWI was done at b values 0, 500, 1000 s/mm. ADC maps derived automatically on voxel by voxel basis. The ADC values are expressed as mean ± standard deviation as \( A \times 10^{-3}\) mm²/s up to 4 decimal places. Region of interest (ROI) for quantitative measurement of ADC were placed on renal parenchyma with out preference for cortex /medulla. Values for each kidney were recorded and relationship between ADC values and stage of CKD, ultrasonography grade of renal parenchyma were evaluated for each patient.

**RESULT**

In our study, out of total 50 cases, majority number of cases were seen in male population-32 cases (64%), with younger age group (21 – 30 years) predominance-16 cases (32%). Clinically based on eGFR calculation, CKD stage I cases were 4(8%), Stage II – 2(4%), Stage III – 8(16%), Stage IV –14(28%) and stage V cases 22(44%). On ultrasound grading, Grade I cases- 10(20%), Grade II cases-16(32%), Grade III cases – 14(28%) and Grade IV cases-10(20%). ADC values of stage IV and V were correlating with CKD staging and Ultrasound Grading (90 to 100%) whereas clinically CKD stage I and Stage II, and USG grade I and II cases ADC values are not correlating as most of cases were showing Stage III ADC values. Linear correlation was seen with
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Graph-4: Pie diagram showing eGFR CKD grading: (% of patients)

Graph-5:

Graph-6: ADC values and renal function

Graph-7: ADC values and stages of CKD

ADC values and eGFR and inverse correlation was seen with serum markers
Out of total 50 cases, majority number of cases were seen in male population-32 cases (64%) (fig-1)
Majority of cases were seen in younger age group population that is between 21-30years of age constituting 32% (16 out of 50 cases) (fig-2).
On ultrasound grading, Grade I cases- 10 (20%), Grade II cases-16 (32%), Grade III cases – 14(28%) and Grade IV cases-10 (20%) (fig-3).
Clinically based on eGFR calculation, CKD stage 1 cases were4(8%), Stage II – 2(4%), Stage III – 8(16%), Stage IV-14(28%) and stage V cases 22(44%) (fig-4).
The adc values($x 10^{-3}\text{mm}^2/\text{s}$) of stage 3 were in the range of 2.170-2.410, stage 4: 1.630-2.050 and stage 5 : 1.340-1.700 which were taken as reference from previous studies (table-1).
ADC values of stage IV and V were correlating with CKD staging and Ultrasound Grading (90 to 100%) whereas clinically CKD stage I and Stage II, and USG grade I and

Graph-8: ADC values and serum markers of renal function

Graph-9: ADC values and serum markers of renal function
Figure-1: A 65 year old male presented with decreased urine output and bilateral loin pain
Serum creatinine: 2.1 mg/dl
eGFR: 36 ml/min/1.73 m² - CKD Stage III
USG: Grade III
MRI: Mean ADC values of 2218/2027 - MRI ADC stage III

Figure-2: A 32 year old male presented with decreased urine output with giddiness, k/c/o young hypertension
Serum creatinine: 4.1 mg/dl
eGFR: 17 ml/min/1.73 m² - CKD grade IV
USG: Grade III
MRI: mean ADC values of 1752/1792 - MRI ADC stage IV

Figure-3: A 45 year old female with generalised edema, decreased urine output, shortness of breath
Serum creatinine: 12.8 mg/dl
eGFR: 3.2 ml/min/1.73 m² - CKD stage V
USG: Grade IV
MRI: Mean ADC values of 1509/1557 - MRI ADC Stage V

Figure-4: A 40 year old female with loin pain
Serum Creatinine: 0.9 mg/dl
eGFR: 90 ml/min/1.73 m² - CKD stage I
USG: Grade I
MRI: mean ADC of 2115/2150 - MRI ADC stage III

different stages of CKD showing decreasing ADC values with increasing stage of CKD
The mean ADC values of different stages of CKD were significantly different from each other and showed decreasing trend with increasing stage (2.01504 ± 0.1243 \(\times 10^{-3}\) mm\(^2\)/s) for stage-3, 1.8263 ± 0.2117 \(\times 10^{-3}\) mm\(^2\)/s) for stage-4, and 1.2208 ± 0.1853 \(\times 10^{-3}\) mm\(^2\)/s) for stage-5 (fig-7).

Within the CKD study group \((n = 50)\), a significant linear correlation was found between renal parenchymal ADC values and eGFR (fig-8). There was a significant inverse correlation between ADC values of renal parenchyma and \(S\) Cr levels (Fig- 9).

**DISCUSSION**

According to our study, chronic kidney disease is more common in males with mean age of 42 years, which is similar to observations in previous studies.

By grading the renal parenchymal changes, the severity of
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**Figure-5:** A 44-year-old male presented with bilateral loin pain. Serum Creatinine: 1.3 mg/dl eGFR: 68 ml/min/1.73 m2 - CKD Stage II USG: Grade II MRI: Mean ADC values of 2061/1958 - MRI ADC Stage III

Table-2: Diagram 5: Ultrasonographic grading and ADC values correlation

<table>
<thead>
<tr>
<th>Ultrasound grading-% of patients</th>
<th>CKD staging (eGFR)-% of patients</th>
<th>ADC values-% of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade I - 20</td>
<td>CKD stage I - 8</td>
<td>MRI stage III - 12</td>
</tr>
<tr>
<td></td>
<td>CKD stage II - 4</td>
<td></td>
</tr>
<tr>
<td>Grade II - 32</td>
<td>CKD stage III A - 8</td>
<td>MRI stage III - 16</td>
</tr>
<tr>
<td>Grade III - 28</td>
<td>CKD stage III B - 8</td>
<td>MRI stage IV - 28</td>
</tr>
<tr>
<td></td>
<td>CKD stage IV - 28</td>
<td></td>
</tr>
<tr>
<td>Grade IV - 20</td>
<td>CKD stage V - 44</td>
<td>MRI stage V - 44</td>
</tr>
</tbody>
</table>

In our study, clinically CKD stage I and Stage II, and USG grade I and II cases ADC values are not correlating as most of the cases were showing Stage III ADC values. ADC values of stage IV and V were correlating with CKD staging and Ultrasound Grading (90 to 100%) which is similar to Goyal, et al.

The mean ADC values of renal parenchyma in patients with CKD was significantly lower than in patients with normal renal function, which is similar to Namimoto et al. reported that ADC values in both the cortex and the medulla of the kidneys of acute and chronic kidney disease patients were significantly lower than the values in normal population in previous investigations.

Low ADC values of renal parenchyma explained by reduced perfusion, reduced water diffusion, glomerulosclerosis, tubular atrophy and interstitial fibrosis. Low ADC (2.035 x 10^-3) were seen in renal dysfunction while high ADC values (2.451 x 10^-3) were seen in normal function. The mean ADC values in different stages of CKD were different from each other showing decreasing trend with increasing stages of CKD.

Significant difference was seen with stage 4, 5 but not with stage 1, 2, 3 of CKD. There was a changing trend observed in incidence of CKD, with renal parenchymal changes observed in younger age groups.

The mean ADC values of different stages of CKD were significantly different from each other and showed decreasing trend with increasing stage (2.01504 ± 0.1243 x 10^-3) for stage-5, 1.8263 ± 0.2117 x 10^-3 for stage-4, and 1.2208 ± 0.1853 x 10^-3 for stage-5) which is similar to Goyal, et al.

Within the CKD study group a significant linear correlation was found between renal parenchymal ADC values and stages of CKD. Similar observations were made by Xu, et al. who found a linear correlation between renal ADC values and stage of CKD.

There was a significant inverse correlation between ADC values of renal parenchyma and serum creatinine levels. Similar to our study, Xu, et al. and Goyal, et al. found a negative correlation between renal parenchyma ADC values and serum creatinine levels.

**Limitations**
We did not recruit healthy volunteers for comparison with renal dysfunction; instead we have taken ADC values in patients who underwent abdominal MRI for various pathologies with normal renal function.

**CONCLUSION**
ADC values may serve as additional tool to identify and estimate degree of renal dysfunction as well as to monitor disease progression. ADC values may help to guide the decision to inject gadolinium based contrast into patients previously known to have renal disease. DWI is not a substitute to serum markers or renal scintigraphy to assess
renal dysfunction, rather it is an additional tool. Addition of DWI to existing MRI protocol provide additional imaging information with minimal increase in acquisition time and MRI can be one stop modality for renal evaluation, as it can evaluate Morphology(T1, T2),Pelvicalcyeal system(MR UROGRAPHY),Vascularity (MR ANGIOGRAPHY) and Function (DWI). ADC values of stage IV and V were correlating with CKD staging and Ultrasound Grading (90 to 100%) whereas clinically CKD stage I and Stage II, and USG grade I and II cases ADC values are not correlating as most of cases were showing Stage III ADC values. Linear correlation was seen with ADC values and e GFR and inverse correlation was seen with serum markers. We conclude that ADC values can play a role in the evaluation of renal dysfunction. Cut off values that we obtained may be useful for stage 3 CKD patients that classified in early stages of disease and respond to treatment.

REFERENCES