Complex Monitoring of Biochemical and Radionuclide Parameters in Patients with Metastatic Renal Cell Carcinoma during Immunotherapy

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Study Objective. To study the effectiveness of complex monitoring of the kidney function, based on biochemical and radionuclide methods in patients with metastatic renal cell carcinoma (mRCC).

Materials and Methods. 41 mRCC patients after nephrectomy received nivolumab ($n=23$) and interferon-$\alpha$ ($n=18$) from 2015 to 2017. At baseline and 2 months after, all patients underwent blood chemistry, urinalysis, Rehberg test, and ELISA to determine serum levels of IL-17A, TGF-$\beta$, and erythropoietin. The monitoring of the renal function and urodynamics by complex renal scintigraphy (CRS) was used for all patients using a dual-detector gamma camera and simultaneous data recording in 2 projections. The interpretation of CRS data used the original SENS CRS technology.

Study Results. Statistically significant correlations were established between IL-17A, TGF-$\beta$, and D (excretion rate of $99m$Tc-technephore from the parenchyma) and Rnfsc (a stable sign of nephrosclerosis), respectively. A significant correlation was established between the parameters of the complex functional monitoring with the prognosis for the risk of renal failure (RF) and efficacy of immunotherapy in mRCC.

Conclusions. All mRCC patients after nephrectomy were recommended to undergo biochemical monitoring with inclusion of TGF-$\beta$ and IL-17A, as well as radionuclide monitoring (CRS) to determine the RF risk at an early stage.

1. Introduction

Literature describe cases of proteinuria and irreversible renal failure during immunotherapy and targeted therapy in patients with mRCC, resulting in a dose reduction and/or drug withdrawal, which can affect the objective response [1–4]. All patients with mRCC after nephrectomy should be assigned to a high risk group for developing chronic kidney disease (CKD).

Until recently, clearance of endogenous creatinine was the most widely used method for determining GFR in clinical practice. However, in moderate to severe renal insufficiency, GFR values calculated from the endogenous creatinine clearance are significantly overestimated, since creatinine is secreted by the proximal tubule in the settings of renal failure and uremia [5, 6].

In order to estimate GFR, formulas such as MDRD (Modification of Diet in Renal Disease), Cockcroft-Gault, CKD-EPI (Chronic Kidney Disease Epidemiology Collaboration), and MCQ (Mayo Clinic Quadratic) are widely put into practice. Currently, there is evidence allowing suggesting that the screening diagnosis of CKD should be based on only a simultaneous estimation of GFR and albuminuria/proteinuria, which allows estimation of the prognosis and risk of cardiovascular events [7].

The KDIGO (Kidney Disease: Improving Global Outcomes) classification, as is the case with RIFLE (risk, injury, failure, loss, and end-stage kidney disease) and AKIN (acute
kidney injury network), based on serum creatinine levels and
the volume of diuresis, allow timely diagnosing acute kidney
damage and have a prognostic value, but do not allow to
take into account causes of kidney damage and, therefore, do
not always help determine preventive and treatment tactics
[8, 9]. In this regard, the search for the most accurate
biomarkers that would allow for early diagnosis of CKD
and establishing the cause of its development is of current
concern. At the moment, a number of biomarkers associated
with nephrotoxicity are known, but FDA (Food and Drug
Administration) and EMEA (European Medicines Agency)
have approved only KIM-1 (kidney injury molecule-1), albumin,
total protein, β2-microglobulin, cystatin C, clusterin, and TFF3 (trefoil factor 3) for routine use in practice [10, 11].

The rate of progression of chronic renal failure (CRF) is
noted to be proportional to the rate of the renal parenchyma
sclerosis, a fundamental component of the CRF pathogenesis
[6]. Transforming growth factor-β (TGF-β) plays the most
prominent role in this process. The signaling pathways of
this growth factor (Smad, p38, Erk1/2, PI3K, JNK, etc.) can
cause glomerulosclerosis and tubulointerstitial fibrosis via
multiple pathological processes [12]. An increase in TGF-
β activity stimulates the cell proliferation and accumulation
of extracellular matrix (ECM) components, such as collagen
types I, III, and IV, laminin, and cellular and plasma forms
of fibronectin, which contributes to the development of
glomerulosclerosis [12].

Hemodynamic effects of glomerular hypertrophy caused
by the loss of the renal mass are closely related to the
mechanisms of persisting kidney inflammation and fibro-
sis via interaction of angiotensin II, TGF-β, and other
growth factors. In addition to hemodynamic effects, pri-
marily a systemic vasoconstrictor action also transmitted to
the glomerular capillaries. So angiotensin II has so-called
inherent nonhemodynamic effects, including the ability to
induce endothelial dysfunction, as well as to enhance local
renal expression of TGF-β. A nonhemodynamic component
of the angiotensin II effect plays the main role in aggra-
vation of proteinuria, which is why the drugs that block
its formation (ACE inhibitors) or interaction with type 1
receptors (angiotensin II receptor blockers) have prominent
antiproteinuric properties [6].

Of interest is investigation of IL-17 concentrations in the
serum, having strong proinflammatory properties and
inducing severe autoimmune pathology, including nephritis
[13, 14].

In recent years, glomerulotrophic radiopharmaceuticals
labeled with radioisotopes are widely used as marker sub-
stances allowing determining GFR [15]. The diagnostic
value of renal purification from nephrotropic substances (99mTc-
MAG3, 123I-hippuran, and 99mTc-DTPA) closely correlates
with inulin clearance [16–18]. However, GFR studies using
radioactive isotopes are used only if specialized radiologi-
cal laboratories are available [19, 20]. In this regard, the
Laboratory for Radioisotope Diagnosis of Russian N.N.
Blokhin Cancer Research Center, a federal state-funded sci-
entific institution, has developed the systemic examination of
nephrourological status based on complex renoscintigraphy
(SENS-CRS) and has been using it in pediatric and adult
clinical practice for more than 15 years [19, 21–23]. SENS-CRS
is a high technology implemented in the development of an
automated workplace (AWP) for a radionephrologist (Project
Manager A.P. Alekhin) (Figure 1).

SENS-CRS is designed for a rapid assessment of the
functional reserves of the urinary system and the risk of renal
failure. The CRS method provides not only the monitoring
of the concentration levels in the parenchyma, but also early
detection of relative stagnation in the parenchyma, its edema,
urine stasis in the departments of the pyelocaliceal system
(PCS), and lower urinary tract, that is, at all functional
structural levels. Biochemical parameters of the kidney func-
tion, such as serum creatinine and urea, reflect quite gross
morphologic alterations in the renal parenchyma and become
diagnostically significant when 50 to 70% of active nephron
mass (ANM) have already become dysfunctional [19].

When planning this study, it was assumed that complex
monitoring of the renal function based on biochemical and
RN methods will allow diagnosis of the risk factors for RF
at an early stage, differentiate the structural kidney damage
from the functional one, determine their relationship with
immunotherapy toxicity and efficacy in patients with mRCC,
and timely prescribe concomitant therapy.

2. Materials and Methods

This study included 41 mRCC patients after nephrectomy
within the period from 2015 to 2017. 18 patients were treated
with interferon (IFN-α) and 23 patients were treated with
nivolumab (as part of the BMS expanded access program). Of
18 patients treated with IFN-α, 16 patients (88.8%) received
it as the first line therapy. The median age was 56 years.
Before (within a week) and during the treatment (every 2
months), all patients underwent blood chemistry, urinalysis,
and Rehberg test.

The serum levels of proteins studied were determined
according to the standard procedure prior to the treatment
(within a week) and 2 months after. The serum was obtained
after blood centrifugation at 3000 rpm, 4°C for 10 min using
RS-6 model centrifuge (Technocent, Russia). 300–400 μL
of serum were dispensed in 2 plastic tubes and stored at
−80°C until the analysis. ELISA tests for IL-17 (eBioscience,
USA), TGF-β1 (eBioscience, USA), and EPO (erythropoi-
etin) (Biomerica, USA) were performed using standard kits
for direct immunoassay according to manufacturer's instruc-
tions.

Complex renoscintigraphy (CRS) was carried out on a
dual-detector gamma camera (E-com, Siemens) with simul-
taneous recording in 2 projections, which allowed studying
the entire renal clearance system, starting with the
heart blood flow and ending with the bladder. Diagnostic
simulation of renal clearance from nephrotropic sub-
stances begins with intravenous administration of 99mTc-
technephore. 99mTc-technephore, a Russian product from
the group of bishophonates, has hemodynamics of a glomeru-
lotropic product, concentrating mainly in the nephrons
via filtration, with partial involvement of secretion. The
Figure 1: An automated workstation implementing the SENS-CRS technology on a personal computer by processing CRS DICOM (Digital Imaging and Communications in Medicine) files generated by a modern dual-detector gamma camera. In the upper band a set of tool icons is used in the workstation in the analysis of data of functional radionuclide studies of the kidneys. In the left part there is a set of zones of interest (an interphase in Russian language), selected on scintigraphic images of the kidneys and urinary tract according to the SENS-CRS technology. The curves obtained from the automated workstation represent the dynamics of the concentration of urine labeled with a nephrotrropic radiopharmaceutical agent (99mTc-MAG3) in the structures of the left (L) kidney: the group of small upper calicles, the big upper calicles, the pelvis, and the ureter (delay of labeled urine in the middle third). Scintigrams on the right: top, image of the urinary system in the front projection; bottom, in the back projection. At the top there is a color scale (the red color corresponds to the maximum score on scintigrams). In a separate window (in the center): results of a quantitative estimation on original algorithms of function parameters of urinary system are shown.

Visualization quality (even with a weak kidney function) of 99mTc-technephore is comparable to that of tubulotropic 99mTc-MAG3 (mercaptoacetyltriglycine) and 123I-hippuran, significantly outperforming conventional glomerulotropic 99mTc-DTPA (diethylenetriaminepentaacetic acid) [21]. The data registered in 2 projections is processed based on 2-phase registration: the first step is a 21-minute (1 min, angiophase) basic test with administration of the labeled substance; the second phase is delayed (after a 25-minute break) 21-minute examination (sometimes a 7-minute test) without administration of RP, but with administration of small amounts of water (200–300 ml) and/or an antispasmodic (less frequently diuretic). The bladder emptying by the patient before the baseline test and examination is a mandatory functional test. Complex renoscintigraphy allowed achieving the lowest radiation doses for patients and staff. When CRS is performed, adults are administered intravenously 74 MBq of 99mTc-technephore (an effective equivalent dose of 0.6 mSv), less frequently 99mTc-technephore; children are given a radiopharmaceutical based on their age and body weight. When kidneys and bones are investigated on the same day, adults are given 370 to 555 MBq of 99mTc-technephore (an effective equivalent dose of 3.0–4.5 mSv). The interpretation of CRS data used a concentration-rate model of urinary excretion and the original SENS CRS technology developed in the laboratory of radioisotope diagnostics, Russian N.N. Blokhin Cancer Research Center. The level of concentration of both glomerulo- and tubulotropic radiopharmaceuticals in the parenchyma is proved to be a well reproducible measure of the kidney concentration function [21, 24]. The statistical analysis of the results was performed using software Statistica 13.0 with the Spearman nonparametric method ($R_{sp}$ is a correlation coefficient; the result was considered insignificant at $p \geq 0.05$).

3. Study Results and Discussion

This study included 41 mRCC patients after nephrectomy within the period from 2015 to 2017; 18 patients were treated with IFN-α and 23 patients were treated with nivolumab. Of 18 patients treated with IFN-α, 16 patients (88.8%) received it as the first line therapy. Twelve (12) patients (52%) and 11 patients (48%) in the nivolumab group received nivolumab as their second line and third or further line therapy. Thus, given the large number of previous lines of therapy, the nivolumab group had a higher risk of developing tubulointerstitial nephritis (TIN). Thus, the incidence of stage-3 CKD at the time of treatment was 35% in the nivolumab group and 17% in the IFN-α group.

Inflammation of the renal tubulointerstitium is always clinically characterized by impairment of the renal concentration function and often renal filtration function. Renal glomeruli can be abnormal, but abnormalities have a secondary nature [7]. As a result, the amount of the radiopharmaceutical concentrates reduced during a radionuclide study, which determines the fundamental premise of the
complex renography (CR), the “concentration function” as a total result of all processes in the renal parenchyma [25]. In SENS-CRS, an algorithm was developed that determines the level of compensation and the risk of destabilization of the total renal function in gradations of FSS (Functional Systems Scores), the total prognostic index of the urinary system functional state, and stability. The relationship between different degrees of clinical parameter gradations—CKD according to KDOQI (Kidney Disease Outcomes Quality Initiative) and KDIGO—and RN parameter—total prognostic index (FSS) according to CRS findings with nephrotropic radiopharmaceuticals ($^{99m}$Tc-technephore, $^{99m}$Tc-DTPA, $^{99m}$Tc-technemag, $^{99m}$Tc-MAG3)—are presented in Table 1.

The grades of the FSS index are the same for the above radiopharmaceuticals (the differences between these drugs are taken into account within the SENS-CRS algorithm). A single concentration-rate approach to the study of kidney function and urodynamics of the urinary tract with different nephrotropic radiopharmaceuticals is described in detail in the publication [21, 26].

The rate of irreversible deterioration of the kidney function in most variants of TIN is much slower than that in other chronic progressive nephropathies. In our study, only 1 patient (2.4%) developed acute renal failure (ARF) after 2 injections of nivolumab. It should be noted that this was the only patient who initially had the lowest RN estimate for the total urinary system function, FSS = 3b (significantly reduced). The cause elimination is crucial in the management of patients with TIN. In this particular case, the patient probably developed tubular necrosis as early as during previous therapy with everolimus (for 2 years); however, at the time of the initiation of nivolumab treatment, GFR, calculated by the MDRD formula, was 41 ml/min. Due to the development of acute renal failure, nivolumab treatment was discontinued. The patient was switched to dialysis.

The presence of confounding factors that can increase the severity of renal disease should be taken into account: chronic heart failure; type 2 diabetes; impaired uric acid metabolism. Elderly patients can have a combination of several forms of renal disease (“multimorbidity”), such as analgesic, urate, and diabetic nephropathy, as well as ischemic renal disease (IRD) and chronic pyelonephritis [6]. In our study, hypertension was observed in 15 patients (36.5%), type 2 diabetes in 4 patients (9.7%), urinary infection in 4 patients (9.7%), urolithiasis in 2 patients (4.8%), and obesity in 2 patients (4.8%).

### Table 1: Relationship between CKD and FSS gradation: a radionuclide estimate for the total renal function.

<table>
<thead>
<tr>
<th>CKD stage</th>
<th>Renal function characteristic</th>
<th>GFR level (mL/min/1.73 m²)</th>
<th>FSS characteristic</th>
<th>FSS index gradation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>High and optimal</td>
<td>&gt;90</td>
<td>Status 1</td>
<td>1a</td>
</tr>
<tr>
<td>2</td>
<td>Slightly decreased</td>
<td>60–89</td>
<td>high level slightly decreased</td>
<td>1b</td>
</tr>
<tr>
<td>3a</td>
<td>Slightly decreased</td>
<td>45–59</td>
<td>Status 2</td>
<td>2a</td>
</tr>
<tr>
<td>3b</td>
<td>Significantly decreased</td>
<td>30–44</td>
<td>moderately decreased</td>
<td>2b</td>
</tr>
<tr>
<td>4</td>
<td>Severely decreased</td>
<td>15–29</td>
<td>Status 3</td>
<td>3a</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>moderately to severely decreased</td>
<td>3b</td>
</tr>
<tr>
<td>5</td>
<td>End-stage renal failure</td>
<td>&lt;15</td>
<td>Decompensation</td>
<td>4</td>
</tr>
</tbody>
</table>

### Table 2: The relationship between biochemical and radionuclide parameters in the monitoring of the urinary system function.

<table>
<thead>
<tr>
<th>N = 97</th>
<th>Plasma creatinine level (112.3 ± 24.4 μmol/L)</th>
<th>Plasma urea level (7.3 ± 2.4 mmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RK, compensation level</td>
<td>$R_sp = +0.3, p &lt; 0.01$</td>
<td>Trend $→$ $R_sp = +0.2, p &lt; 0.05$</td>
</tr>
<tr>
<td>FSS, a total prognostic index for the urinary system</td>
<td>$R_sp = +0.3, p &lt; 0.005$</td>
<td>$p &gt; 0.05$</td>
</tr>
<tr>
<td>$G_{rem}$, a measured level of $^{99m}$Tc-technephore in the renal parenchyma</td>
<td>$R_sp = -0.3, p &lt; 0.001$</td>
<td>$R_sp = -0.3, p &lt; 0.01$</td>
</tr>
<tr>
<td>$A[s]$, an arterial index of renal parenchyma</td>
<td>$p &gt; 0.05$</td>
<td>$p &gt; 0.05$</td>
</tr>
<tr>
<td>IF$_{uret}$, a speed index of the ureteral orifice (during examination)</td>
<td>$R_sp = -0.3, p &lt; 0.01$</td>
<td>$p &gt; 0.05$</td>
</tr>
</tbody>
</table>

3.1. **Analysis of Biochemical and Radionuclide Parameters during Immunotherapy.** The analysis of blood chemistry parameters (creatinine, urea), daily Rehberg test, total protein in the urine, and total and partial RN parameters established their relationship, which confirmed the diagnostic value of both methods—biochemical and CRS (Table 2). We analyzed 97 clinical cases (during the monitoring) constituting the group of patients after nephrectomy during therapy with IFN-α and nivolumab.

There was also a statistically significant relationship between the total GFR based on the 24-hour Rehberg test and RN parameters, $D$, % (RP excretion rate from the parenchyma at the “cortex-medulla” level) and $GB_{20}$ (a 20-minute level of
RP concentration in the urinary bladder in the basic CRS test) (Figure 2).

An increase in proteinuria up to the nephrotic level is determined primarily by the loss of the selectivity of the glomerular basement membrane and progressive podocyte dysfunction. This disorder is also accompanied by inappropriate activation of the renin-angiotensin-aldosterone system, typical of many variants of nephrotic syndrome and resulting in sodium retention and osmotically bound water aggravating the edema in addition to the developed resistance of respective nephron segments to natriuretic peptides [7]. Hypercoagulability typical for nephrotic syndrome is determined primarily by the activation of serum and endothelial hemostasis, which results in a trend toward increased proteinuria with the development of CTIN and possible urinary tract infection (UTI).

With regard to the search of diagnostic markers of early stages of renal dysfunction, we analyzed transforming growth factor-\(\beta\) (TGF-\(\beta\)) as a factor of renal parenchyma sclerosis and IL-17 having strong proinflammatory properties and inducing severe autoimmune pathology, including nephritis, in the serum of 40 patients with mRCC prior to immunotherapy (IFN and nivolumab) and 2 months after. Against the backdrop of immunotherapy with the inclusion of INF-\(\alpha\) and nivolumab, there was a significant increase in IL-17A from 0±4.29 (median ± SD) to 0.166±1.714 pg/ml (\(p<0.0005\)) and a trend toward TGF-\(\beta\) growth from 11.3±12.4 to 13±10.1 ng/ml (\(p=0.1\)) (Figure 4). The study was able to compare TGF-\(\beta\) and IL-17 values with RN parameters, Rnfss, a stable sign of nephrosclerosis (presumably sclerosis of interlobar renal arteries), and \(D\), an excretion rate of \(^{99}\)mTc-technephore from the parenchyma (at the “cortex-medulla” level), respectively (Table 3).

Thus, an increase in TGF-\(\beta\) concentrations correlates with Rnfss during CRS, which confirms the diagnostic value of Rnfss as a visual radionuclide sign of “nephrosclerosis”, whose assessment was carried out according to the grading specified in Table 4. Moreover, in the nivolumab group, nephrosclerosis was much more pronounced than in the IFN-\(\alpha\) group, which may be due to a larger number of previous lines of targeted therapy.

At the same time, an increased IL-17 level both prior to immunotherapy and 2 months after corresponded to a decrease in the RP excretion rate from the parenchyma (\(D\)) due to increasing interstitial edema, confirming the importance of cytokine IL-17 in the pathogenesis of autoimmune nephritis.
It should be noted that these biochemical and RN markers not only allow establishing impairment of the renal function at an early stage, but also differentiating the cause of this disease, nephrosclerosis or an autoimmune condition, which is extremely important in determining the treatment strategy for CKD.

Also, in the IFN-α group, endogenous erythropoietin levels were evaluated by ELISA in 15 mRCC patients after nephrectomy prior to the treatment. The findings had a significant correlation with creatinine levels before the treatment ($R_S = -0.6, p < 0.05$). Thus, reduced erythropoietin levels correlated with increased creatinine concentrations, which is consistent with the pathophysiological basis of the renal function (Figure 5).

In the SENS-CRS technology, radionuclide (CRS with 99mTc-technetium or 99mTc-MAG3, rarely with 99mTc-DTPA) images of the kidneys in the gray scale are obtained using a $64 \times 64$ matrix, chosen to minimize the dose of the radiopharmaceutical administered and the patient's radiation load.

### 3.2. The Prognostic Value of Laboratory Diagnostic and Biochemical Parameters.

At the first stage, we evaluated the effect of RN and biochemical parameters on the RF risk estimated based on an increase in creatinine and urea levels during immunotherapy with IFN-α and nivolumab. The data of the nonparametric correlation analysis for parameters potentially significant for the prognosis of RN risks are presented in Table 5.

Statistically significant correlations are established between an increase in creatinine levels and IL-17, FSS, $G_{eff}$, and Rnfss (Figure 6) and an increase in urea and protein levels in the urine, IL-17, $D$, $T_{ev}$, and $T_{pelv}$ (Figure 7). Thus,
Table 4: The rating scale of an RN visual sign of “nephrosclerosis” in points.

<table>
<thead>
<tr>
<th>Rnfsc, a visual sign of nephrosclerosis</th>
<th>Points</th>
<th>Typical scintigrams</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left kidney</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basic test (left), examination (right)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right kidney</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basic test (left), examination (right)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>None (?)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Contradictory picture of initial changes</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>Unsure sign of “nephrosclerosis”</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Picture of irreversible “nephrosclerosis”</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

Figure 6: Correlation between a 1.2-fold increase in creatinine levels (±0.4) during immunotherapy and $G_{\text{eff}}$ prior to the treatment initiation.

Figure 7: Correlation between a 1.6-fold increase in urea levels (±0.6) during immunotherapy and $D$, % prior to the treatment initiation.

the initial IL-17 value in the serum can be an early predictor for RF development in CKD during immunotherapy, whereas serum creatinine and urea levels gave no statistically significant results in terms of the RF prognosis.
Table 5: An effect of biochemical and radionuclide parameters on the prognosis for RF risk during immunotherapy.

<table>
<thead>
<tr>
<th>Before initiation of immunotherapy</th>
<th>A 1.2-fold increase in plasma creatinine level (±0.4)</th>
<th>A 1.6-fold increase in plasma urea level (±0.6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameters of laboratory blood and urine tests (n = 40)</td>
<td>Parameters of laboratory blood and urine tests (n = 40)</td>
<td></td>
</tr>
<tr>
<td>Plasma creatinine level</td>
<td>p &gt; 0.05</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>Plasma urea level</td>
<td>p &gt; 0.05</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>Protein in urine</td>
<td>Trend →</td>
<td>R_sp = +0.5, p &lt; 0.05</td>
</tr>
<tr>
<td>Plasma TGF-β_1</td>
<td>Trend →</td>
<td>Trend →</td>
</tr>
<tr>
<td>Plasma IL-17</td>
<td>R_sp = +0.4, p &lt; 0.05</td>
<td>R_sp = +0.4, p &lt; 0.05</td>
</tr>
</tbody>
</table>

Parameters of complex renoscintigraphy with 99mTc-technephore (n = 31)

| FSS, a total prognostic index for the urinary system | R_sp = +0.4, p < 0.05 |
| G_eff, an effective index of the renal concentration function | R_sp = +0.4, p < 0.05 |
| D, %, an excretion rate of 99mTc-technephore from parenchyma | R_sp = +0.5, p < 0.005 |
| T_cr [min], the time of start of excretion of the labeled urine from the pyelocaliceal system | R_sp = +0.4, p < 0.05 |
| T_pe [min], the time of start of excretion of the labeled urine from the renal pelvis | R_sp = +0.3 (p = 0.06) |
| Rnfss, a visual sign of nephrosclerosis | R_sp = +0.4, p < 0.05 |

Table 6: An effect of biochemical and RN parameters on the immunotherapy effectiveness.

<table>
<thead>
<tr>
<th>Before initiation of immunotherapy (n = 40)</th>
<th>Treatment effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameters of laboratory blood and urine tests</td>
<td>Parameters of laboratory blood and urine tests</td>
</tr>
<tr>
<td>Plasma creatinine level</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>Plasma urea level</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>Protein in urine</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>TGF-β_1</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>IL-17</td>
<td>p &gt; 0.05</td>
</tr>
</tbody>
</table>

Also, a correlation was observed between the RN sign of nephrosclerosis (Rnfss) and an increase in creatinine levels. Thus, despite the lack of a statistically significant correlation between serum TGF-β_1 and an increase in creatinine levels, but given the correlation between TGF-β_1 and Rnfss (p < 0.005), TGF-β_1 can be considered as a relative risk factor for RF development during immunotherapy in patients with mRCC who underwent nephrectomy.

At the next stage, we evaluated the effect of RN and biochemical parameters on the effectiveness of immunotherapy estimated according to the RECIST criteria (Table 6).

As it turned out, the CRS findings have prognostic significance in relation to not only the RF risk, but also the efficacy of immunotherapy in patients with mRCC after nephrectomy. It should be noted that most of correlations were obtained due to statistically significant relationships in the IFN-α group. Thus, patients with mRCC with better kidney function parameters have a better prognosis regarding the efficacy of immunotherapy.

4. Conclusion

Thus, immunotherapy has no pronounced nephrotoxicity. All mRCC patients after nephrectomy were recommended, prior to the treatment initiation, to undergo biochemical monitoring with inclusion of TGF-β_1 and IL-17, as well as radionuclide monitoring (SENS-CRS) to determine the RF risk at an early stage and to establish prognosis for the underlying disease and timely adjust the treatment in order to improve their response to immunotherapy.

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.
References


