The Clinical Efficacy of Low-Dose Tacrolimus Combined with Tripterygium to Treat the Steroid-Resistant Nephrotic Syndrome

Hongqi Ren1, Guofang Chen2, Xuan Zhou1, Yan Li1, Qing Cai1, Shujing Han1, Rui Wang1
1Department of Nephrology, Huaihai Hospital of Xuzhou Medical College, Xuzhou, China
2Department of Neurology, Central Hospital of Xuzhou, Xuzhou, China
Email: sznk2005@yahoo.com.cn

Received September 30, 2012; revised November 2, 2012; accepted November 15, 2012

ABSTRACT

Objective: To observe the clinical efficacy and safety of low dose tacrolimus (TAC) combined with tripterygium (TW) in treatment of steroid resistant nephritic syndrome (SRNS). Method: The patients, who were diagnosed with mesangial proliferative glomerulonephritis (MesPGN) and focal segmental glomerulosclerosis (FSGS) by biopsy and failed to respond to a 3-month treatment with prednisone (1 mg/kg·d), were randomly divided into 2 groups (TAC + TW Group and TW Group). Initially TAC + TW group took TAC 0.05mg/(kg·d) 2 h after meal at 12 h interval. The plasma TAC level was examined after 3 days and was kept at 1.5 - 4 ng·ml; meanwhile, TW was given at 60 mg/d before meal. TW group only took TW (60 mg/d). The efficacy, adverse reactions and plasma TAC levels were observed in each group. Results: 1) Totally 20 SRNS patients completed the trial, 11 of TAC + TW Group and 9 of TW Group. There is no statistical difference between the two groups in terms of age, gender, duration since onset of the disease, blood pressure, 24 h UPQ, serum albumin, creatinine, cholesterol, triglyceride, FBG, kidney pathological categories, time of taking prednisone etc.; 2) Urine protein started to decrease after 1 month treatment in both of TAC + TW and TW groups. By the 12th month of treatment, TAC + TW group showed 8 cases of complete remission (72.7%), 2 cases of partial remission (18.2%) and 1 case of no improvement (9.1%), while those of TW groups were 2 (22.2%), 4 (44.5%) and 3 (33.3%), respectively; 3) With treatment, the TAC + TW Group patients’ plasma protein was significantly higher than that of pretreatment stage and recovered to normal level after 6 month of treatment. However, there was no significant plasma protein increase in TW Group. No obvious changes were observed on serum creatinine level of patients of both the two groups; 4) The incidence of adverse reactions was not significantly different between the two groups. Conclusion: TAC + TW reduced proteinuria of SRNS patients, increased clinical remission rate and was tolerant to SRNS patients. We conclude that TAC + TW treatment is an effective way to treat patients with SRNS.

Keywords: Steroid-Resistant Nephrotic Syndrome; Tacrolimus; Tripterygium; Treatment Outcome

1. Introduction

As an immune disease, primary nephrotic syndrome is usually treated with adrenocortical hormone and cytotoxic drugs in clinic. For patients whose clinical symptoms are not remitted completely or partially after being treated with hormone of standard dosage (1 mg/kg·d) for at least 3 months, they are diagnosed with steroid-resistant nephrotic syndrome (SRNS) [1]. SRNS is usually presented, pathologically, as minimal change disease (MCD), mesangial proliferative glomerulonephritis (MesPGN) and focal segmental glomerulosclerosis (FSGS), with a few of membranous nephropathy and individual of non-IgA mesangial proliferative glomerulonephritis, which is difficult to be treated and generally immunosuppressive agents such as cyclophosphamide and cyclosporine A are needed for the treatment in clinic [2,3]. Tacrolimus (TAC) is a new immunosuppressive agent, by suppressing calmodulin, the activation of T-cell, hyperplasia of T-helper-dependent B cell and the expression of lymphocyte factor and receptor to achieve immunodepression [1,2,4]. In 1990, Mccauley [5] reported children HRNS cases caused by the treatment of FSGS with TAC for the first time. Up to now, most of the treatment on primary nephrotic syndrome with TAC are still about the report of cases and only a few are about retrospective study and prospective study. And the treatment objects are hormone resistance and (or) dependent nephrotic syndromes patients mostly attacked FSGS and MCD [6,7]. In the 1980s, Leishi Li first used common three-wingnut root, a traditional medicine on the treat-
ment of renal glomerular diseases and in the later severe
diseases, through clinical application, it has been found
that tripterygium (TW) presents its unique curative ef-
facts in the treatment of many renal glomerular diseases
such as minimal change disease, IgM nephropathy and
IgA nephropathy. Researches showed that common three-
wingnut root has obvious functions of anti-inflammation
and immunosuppression; In addition, a series of basic
researches in recent years have proved that common three-
wingnut root has unique functions of podocyte protection
and can reduce the urine protein amount of glomeru-
lonephritis patients [8,9].

Currently, certain curative effects have been achieved
through the treatment of SRNS with sufficient and long-
treatment TAC and double-dosage TW in clinic, but TAC
is expensive and has certain side effects, so it is difficult
to promote it in clinic. On the other hand double-dose
TW has serious side effects such as liver dysfunction and
paramecia for long-term use. The disease may easily re-
cur if TW is reduced to single dose.

Therefore, it is not feasible for TW to be used in clinic
for a long time. Based on the previous researches, we ob-
served the effectiveness and safety of treatment of SRNS
with small-dose TAC combined with single-dose TW, aiming to explore a cost-affordable method to treat SRNS.

2. Target and Method

2.1. Case Selection

20 patients hospitalized in our hospital from January
2007 to September 2009 are selected and all of the cases
meet the following conditions: 1) Age: 18 - 85, gender:
unlimited; 2) Clinically being diagnosed as glomerular
mesangial hyperplasia cases and glomerular FSGS cases
through kidney biopsy; 3) Clinical manifestation: brine
protein ration >3.5 g/d, serum creatinine (SCr) <132.6 μmol/L; 4) No curative effects
are observed on the patients after their taking metacor-
tandracin (1 mg/kg/d and maximum 60 mg/d) for above
three months. The following situations are excluded: 1) Clear secondary cases such as systemic lupus erythema-
tosus, hepatitis B, hepatitis C virus infection, thyroid
gland and tumor in other parts and poisoning of organic
solvent and heavy metal; 2) Patients with serious complica-
tions, such as infection, cardiac insufficiency, liver
dysfunction and disorders of galactose metabolism; 3) Patients who have received the treatment of cyclophos-
phamide (CTX), cyclosporin A (CsA), leflunomide and
mycophenolate mofetil. This research has been approved
by the Ethics Committee of our hospital and all patients
have signed the Informed Consent before being arranged
in the groups.

2.2. Research Design and Treatment Plan

2.2.1. Random Grouping

Before participating in the research, patients were ap-
plied to reduced dosage of metacortandracin (20 mg/d, reducing 5 - 10 mg each week). Those patients were di-
vided into TAC + TW Group and a TW Group randomly,
with the group number of 11 and 9 respectively. The fol-
low-up visit observation time is 12 months.

2.2.2. Treatment Plans

TAC + TW group: TAC produced by Astellas China Phar-
maceutical Co., Ltd. (0.5 mg/pill and 1 mg/pill) is applied
in the test. Patients took TAC 0.05 mg/kg/d 2 hours after
meal every 12 hours. The whole blood TAC concentra-
tion was detected (by ELISA method, PROTrac II kit
produced by DiaSorin Corporation) 3 days later. Later,
TAC concentration was examined every month and the
blood concentration was maintained at 1.5 - 4 ng/ml.
During the period of TAC treatment, 60 mg/d TW (pro-
duced by Taizhou Pharmaceutical Factory, 10 mg/pill
and 100 pills/bottle) were taken by patients be fore 3
meals. TW group: only TW was taken by patients before
3 meals, patients whose conditions are completely or
partially remitted are required to continue taking it up to
12 months.

Reduction of metacortandracin: for TAC + TW Group,
metacortandracin was reduced by 5 mg/d each week.
Later, it was reduced by 5 mg every other day each week
when it was reduced to 20 mg/d and 2.5 mg/d every two
weeks when reduced to 20 mg every other day; and re-
duced gradually till after it was stopped. For TW group,
metacortandracin was reduced by 5 mg every two weeks;
changed as reduction by 5 m every other day when it is
reduced to 20 mg/d and 2.5 mg every two weeks when
reduced to 20 mg every other day; and gradually reduced
till it is stopped.

If liver dysfunction (alanine or glutamic-oxalacetic trans-
aminase exceeds twice of the normal value) occurs or
Peripheral blood leukocyte reduces to <3 × 10^9/L after
decine taking, TAC would be suspended and TW was
taken continuously for a week in the original dosage; if
the situation mentioned above can return to normal, TAC
and TW can be taken continuously in treatment dosage. If
a patient shows serious complicating infection, the medi-
cine shall be suspended and then be used continuously
after the infection is controlled for 2 weeks and the base-
line standard shall be re-evaluated. In the treatment, pa-
tients stopping taking specified medicine for 2 weeks for
various reasons shall not be considered anymore.

Other treatment: low-salt diet (<3 g/d sodium salt in-
take). If a patient has high-blood pressure, calcium an-
tagonist was preferably used to control the blood pres-
ture to 120/80 mmHg; if ACEI and ARB have been
adopted before treatment, they can be used continuously; diuretic can be used for patients with obvious edema and statin lipid-lowering drugs can be used for patients with serious hyperlipidemia.

2.3. Follow-Up Observation and Observation Indicators

Follow-up observation: Follow-up observation must be carried out in the 2nd week, and the 1st, 3rd, 6th and 12th month after the start of the treatment. To observe patient’s conditions, adverse reactions and laboratory examination were performed.

Laboratory examination: The 24 h urine protein quantitation was examined at the base line and each follow-up visit, and average value at each follow-up visit was calculated. In the 3 days before taking 24 h urine, patients were subject to regular diet, and were not allowed to take diuretic or transfuse albumin.

Efficacy observation: the urine protein level was observed at the 2nd week, and the 1st, 3rd, 6th and 12th month after the beginning of treatment, and the values of plasma protein, liver function, serum creatinine, blood sugar and blood fat were recorded.

Adverse events: All adverse events in therapeutic process should be recorded in details, such as anaphylactic reaction, infection, gastrointestinal reaction, renal function impairment, liver function impairment, blood system impairment and female menses condition, etc.; if the values of glutamic-pyruvic transaminase or glutamic-oxaloacetic transaminase are twice of the upper limit of normal value, it can be defined as elevation of transaminase.

2.4. Assessment of Efficacy [9,10]

Complete remission (CR): urine protein concentration < 0.4 g/24h, serum albumin > 35 g/L, and stable renal function.

Partial remission (PR): urine protein level decreases and exceeds 50% of the base value, ≤3.5 g/d and stable renal function.

No remission (NR): urine protein concentration is lower than 50% of the base value, or concentration >3.5 g/d, or serum creatinine increases and exceeds 50%.

Effective rate = CR rate + PR rate.

3. Statistical Methods

The inspection data were processed with SPSS 13.0 statistical software, and quantitative data were indicated by mean ± standard deviation. Comparison among groups adopts t inspection. Enumeration data were indicated in the number of cases (percentage); comparison among groups adopts $\chi^2$ for inspection, in which $P < 0.05$ represents that the difference has remarkable statistical significance and $P < 0.01$ represents extraordinary statistical significance.

4. Results

4.1. General Conditions

20 SRNS patients were selected, with 9 to TW Group and 11 to TAC + TW Group. There were no obvious statistical difference between the two groups in age, gender, duration since onset of the disease, blood pressure, 24 h UPQ, serum albumin, creatinine, cholesterol, triglyceride, FBG, kidney pathological categories, time of taking prednisone etc. (Table 1).

<table>
<thead>
<tr>
<th>No.</th>
<th>Patient</th>
<th>TW</th>
<th>TAC + TW</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>40.7 ± 11.9</td>
<td>42.8 ± 13.5</td>
<td>0.36</td>
<td></td>
</tr>
<tr>
<td>Gender (M/F)</td>
<td>6/3</td>
<td>7/4</td>
<td>0.00</td>
<td></td>
</tr>
<tr>
<td>Duration since onset of the disease (Mon)</td>
<td>5.2 ± 2.9</td>
<td>4.0 ± 2.2</td>
<td>0.11</td>
<td></td>
</tr>
<tr>
<td>BP (mmHg)</td>
<td>126.0 ± 15.3</td>
<td>125.3 ± 15.7</td>
<td>0.46</td>
<td></td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>75.1 ± 11.8</td>
<td>77.7 ± 9.7</td>
<td>0.29</td>
<td></td>
</tr>
<tr>
<td>Urine protein (g/24hr)</td>
<td>5.6 ± 1.9</td>
<td>6.0 ± 1.4</td>
<td>0.28</td>
<td></td>
</tr>
<tr>
<td>SCr (umol/l)</td>
<td>87.8 ± 9.9</td>
<td>89.1 ± 9.6</td>
<td>0.39</td>
<td></td>
</tr>
<tr>
<td>Alb (g/l)</td>
<td>24.3 ± 2.6</td>
<td>23.0 ± 3.3</td>
<td>0.18</td>
<td></td>
</tr>
<tr>
<td>Cho (mmol/l)</td>
<td>8.9 ± 1.4</td>
<td>8.4 ± 1.6</td>
<td>0.23</td>
<td></td>
</tr>
<tr>
<td>Tg (mmol/l)</td>
<td>3.4 ± 0.9</td>
<td>3.0 ± 0.6</td>
<td>0.18</td>
<td></td>
</tr>
<tr>
<td>Histology</td>
<td>MesPGN</td>
<td>3</td>
<td>4</td>
<td>0.88</td>
</tr>
<tr>
<td></td>
<td>FSGS</td>
<td>6</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>Duration of 1 mg/kg/d prednisone (weeks)</td>
<td>8.3 ± 0.8</td>
<td>9.2 ± 2.0</td>
<td>0.11</td>
<td></td>
</tr>
</tbody>
</table>
4.2. Clinical Efficacy

For TAC + TW Group, 1 month after the treatment, the urine protein of patients started to decrease, with 0 case of complete remission, 3 cases of partial remission (27.3%), and the total effective rate (27.3%); 3 months after treatment: 2 cases of complete remission (18.2%), 7 cases of partial remission (63.6%), 2 cases of ineffectiveness (18.2%), and the total effective rate was 81.8%; 6 months after treatment: 3 cases of complete remission (27.3%), 7 cases of partial remission (63.5%), 2 cases of ineffectiveness (9.1%) and the total effective rate was 90.9%; when the follow-up visits were carried out till the 12th month: 8 cases of complete remission (72.7%), 2 cases of partial remission (18.2%), 1 case of ineffectiveness (9.1%) and the total effective rate was 90.9%. When the follow-up visits were over, the complete remission rate of TAC + TW Group was higher than that of TW Group (Table 2).

For TW Group, 1 month after the treatment, urine protein of patients also started to decrease, with 0 case of complete remission, 2 cases of partial remission; 3 months after treatment: 0 case of complete remission, 5 cases of partial remission (55.6%), 4 cases of ineffectiveness (44.4%), and the total effective rate was 55.6%; 6 months after treatment: 1 case of complete remission (11.1%), 5 cases of partial remission (55.6%), 3 cases of ineffectiveness (33.3%), and the total effective rate was 66.7%; when the follow-up visits were carried out till the 12th month: 2 cases of complete remission (22.2%), 4 cases of partial remission (44.5%), 3 cases of ineffectiveness (33.3%), and the total effective rate was 66.7%.

4.3. Changes of Observation Indicators of the Two Groups before and after Treatment

For TAC + TW Group, comparing with the base line values, the urine protein values of patients have decreased obviously to 28.3%, 69.5%, 82.3% and 86.2% at the 1st, 3rd, 6th and 12th month after the beginning of treatment, which are obviously higher than those of TW Group (Table 4, Figure 1(a)). For TW Group, the urine protein values of patients have also decreased to 28.3%, 44.7%, 52.4% and 59.2% at the 1st, 3rd, 6th and 12th month after the beginning of treatment. The conclusion is: for TW Group, the urine protein value decrease examined one month after treatment has statistical significance; the decrease examined at the end of the test, although still has statistical significance comparing with the base line values, is at a low level of only 59.2% (Table 3).

Table 2. Efficacy comparison of follow-up visits until the end of the trial.

<table>
<thead>
<tr>
<th></th>
<th>PR</th>
<th>CR</th>
<th>NR</th>
<th>Total effective rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>TW</td>
<td>2 (22.2%)</td>
<td>4 (44.5%)</td>
<td>2 (33.3%)</td>
<td>66.7%</td>
</tr>
<tr>
<td>TAC + TW</td>
<td>8 (72.7%)</td>
<td>2 (18.2%)</td>
<td>1 (9.1%)</td>
<td>90.9%</td>
</tr>
</tbody>
</table>

The complete remission rate of TAC+TW group is higher than that of TW group when follow-up visits are ended, P = 0.024 < 0.05.

Table 3. Changes of urine protein and biochemical indicators of TW group before and after treatment.

<table>
<thead>
<tr>
<th></th>
<th>Pre-treatment</th>
<th>2 w</th>
<th>1 m</th>
<th>3 m</th>
<th>6 m</th>
<th>12 m</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urine protein (g/24hr)</td>
<td>5.6 ± 1.9</td>
<td>5.2 ± 1.0</td>
<td>4.4 ± 0.9*</td>
<td>3.1 ± 1.4**</td>
<td>2.7 ± 1.3**</td>
<td>2.3 ± 1.2**</td>
</tr>
<tr>
<td>Scr (umol/l)</td>
<td>87.8 ± 9.9</td>
<td>86.5 ± 6.9</td>
<td>83.2 ± 10.6</td>
<td>80.3 ± 12.2</td>
<td>84.1 ± 5.8</td>
<td>79.9 ± 9.2</td>
</tr>
<tr>
<td>Alb (g/l)</td>
<td>24.3 ± 2.6</td>
<td>25.5 ± 2.3</td>
<td>26.2 ± 2.4*</td>
<td>27.8 ± 1.8**</td>
<td>29.3 ± 1.9**</td>
<td>30.9 ± 2.1**</td>
</tr>
<tr>
<td>Cho (mmol/l)</td>
<td>8.9 ± 1.4</td>
<td>8.3 ± 1.5</td>
<td>7.4 ± 1.0*</td>
<td>5.5 ± 2.1*</td>
<td>4.4 ± 1.4**</td>
<td>5.0 ± 1.7**</td>
</tr>
<tr>
<td>Tg (mmol/l)</td>
<td>3.4 ± 0.9</td>
<td>2.9 ± 0.6</td>
<td>2.9 ± 0.7</td>
<td>2.2 ± 0.5**</td>
<td>2.3 ± 0.5*</td>
<td>2.2 ± 0.6*</td>
</tr>
<tr>
<td>GLU (mmol/l)</td>
<td>5.7 ± 0.7</td>
<td>5.3 ± 0.5</td>
<td>5.4 ± 0.5</td>
<td>5.5 ± 0.7</td>
<td>5.1 ± 0.8</td>
<td>5.2 ± 0.4</td>
</tr>
</tbody>
</table>

*Compared with conditions before treatment P < 0.05; **Compared with conditions before treatment P < 0.01.

Table 4. Changes of urine protein and biochemical indicators of TAC + TW group before and after treatment.

<table>
<thead>
<tr>
<th></th>
<th>Pre-treatment</th>
<th>2 w</th>
<th>1 m</th>
<th>3 m</th>
<th>6 m</th>
<th>12 m</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urine protein (g/24hr)</td>
<td>6.0 ± 1.4</td>
<td>5.1 ± 1.4</td>
<td>4.3 ± 1.3**</td>
<td>1.8 ± 1.4**</td>
<td>1.1 ± 1.0**</td>
<td>0.9 ± 0.9**</td>
</tr>
<tr>
<td>Scr (umol/l)</td>
<td>89.1 ± 9.6</td>
<td>83.5 ± 6.6</td>
<td>82.9 ± 10.4</td>
<td>84.4 ± 5.1</td>
<td>81.3 ± 12.2</td>
<td>81.6 ± 10.8</td>
</tr>
<tr>
<td>Alb (g/l)</td>
<td>23.0 ± 3.3</td>
<td>24.1 ± 2.4</td>
<td>29.8 ± 2.3*</td>
<td>31.5 ± 5.5**</td>
<td>37.4 ± 1.8**</td>
<td>37.1 ± 1.3**</td>
</tr>
<tr>
<td>Cho (mmol/l)</td>
<td>8.4 ± 1.6</td>
<td>8.0 ± 1.1</td>
<td>6.6 ± 1.2**</td>
<td>4.9 ± 1.1**</td>
<td>5.4 ± 1.9**</td>
<td>4.8 ± 1.1**</td>
</tr>
<tr>
<td>Tg (mmol/l)</td>
<td>3.0 ± 0.6</td>
<td>2.8 ± 0.5</td>
<td>2.8 ± 0.5</td>
<td>2.3 ± 0.4**</td>
<td>2.2 ± 0.6**</td>
<td>1.9 ± 0.4**</td>
</tr>
<tr>
<td>GLU (mmol/l)</td>
<td>5.8 ± 0.4</td>
<td>5.4 ± 0.8</td>
<td>5.4 ± 0.6</td>
<td>5.6 ± 0.6</td>
<td>5.3 ± 0.7</td>
<td>5.7 ± 0.4</td>
</tr>
</tbody>
</table>

*Compared with conditions before treatment P < 0.05; **Compared with conditions before treatment P < 0.01.
After treatment, plasma protein of TAC + TW Group increased obviously; when treatment was carried out for 6 months, plasma protein basically recovered to the normal level (Table 4, Figure 1(a)); however, while urine protein of TW Group decreased, although plasma albumin increased, the increase was obviously lower than that of TAC + TW group (Figure 1(b)). The serum creatinine level of patients in two groups had no obvious change before and after treatment (Tables 3 and 4, and Figure 1(c)). In the follow-up visits of TAC + TW group, no patient’s serum creatinine has exceeded 50% of the basic value. The cholesterol and triglyceride values of patients in two groups decreased obviously compared with those values before treatment (Tables 3 and 4), but the decline range of cholesterol of patients in two groups were basically similar with no obvious statistical difference (Figure 1(d)). The fasting blood-glucose level of patients in two groups had no obvious change.

4.4. TAC Concentration Change

The TAC blood concentration of TAC + TW group was maintained at 1.5 - 3 ng/ml, and the average blood concentration was relatively stable during treatment, but 6 months after treatment, the TAC content of patients of complete remission was reduced to 0.5 mg/2/day (Figure 2).

4.5. Adverse Reactions

Generally, adverse reactions are not severe and will maintain for a short period of time. 2 cases of nausea and 1 case of herpes zoster happened to TAC + TW group, and 1 month after treatment, there were 3 cases with slightly increase of glutamic-pyruvic transaminase, and menstrual disorder happened to 2 cases of 4 female cases; no increase of serum creatinine exceeding 50% of baseline value, and the fasting blood-glucose had no ob-

Figures 1. Change of urine protein and plasma albumin of TAC + TW group and TW group before and after treatment. (a) Follow-up consultation and comparison of urine protein of TAC + TW group and TW group; (b) Follow-up consultation and comparison of plasma albumin of TAC + TW group and TW group; (c) Follow-up consultation and comparison of serum creatinine of TAC + TW group and TW group; (d) Follow-up consultation and comparison of cholesterol of TAC + TW group and TW group. *Compared with conditions before treatment P < 0.05; **Compared with conditions before treatment P < 0.01.
vious increase. 1 case of nausea and 1 case of laxative-ness happened to TW group, and after treatment of 1 month, there were 2 cases of slightly increase of glutam-amic-pyruvic transaminase, and menstrual disorder hap-pened to 1 case of 3 females. Patients with increase of transaminase in two groups were given bifendate for symp-tomatic treatment and patients’ conditions were improved; all patients were kept taking medicines and the treatment as well as follow-up visits were not affected.

5. Discussion

Therapeutic methods available for SRNS include the application of cyclophosphamide, cyclosporine A, implosive therapy of high-dose methylprednisolone, ACEI or ARB and mycophenolate mofetil, etc. Most researches reported that the effective rate of SNRS treatment is about 50% - 60%, and patients who have no response to treatment are apt to develop to end-stage renal failure [10].

TAC is a calmodulin inhibitor, which is mechanisti-cally similar to CsA, and its inhibiting effect on cell fac-tor is significantly stronger than that of CsA. In recent years, there were many reports about treating SRNS with TAC. Gulati et al. [11] treated 19 SRNS patients with the combination of TAC and hormone. When the treatment was carried out for 3 months, 16 patients were completely remitted with a complete remission rate of 84%; 2 cases partial remission with a partial remission rate of 10.5%. Li and other researchers (references needed here) treated SRNS patients with TAC combined with low-dose hormone, with the results indicating that the complete and partial remission rate were 64.7% and 17.6% after treatment of 24 weeks in TAC group. However, the TAC concentrations were relatively high, which fluctu-ated at 9.54 ± 5.13 ng/ml, and SRNS is prone to recur if the dose is reduced.

TW plays a unique role in the treatment of many in-flammations, autoimmune diseases and organ transplan-tation rejections. In recent years, in vitro experiments have indicated that triptolide can stabilize the skeleton structure of podocyte and antagonize the damage of puromycin aminonucleoside (PAN) to podocyte, and when podocyte is damaged, triptolide can reverse the damage of podocytes and recover the skeleton structure of podo-cytes and the expression of relevant molecules [8]. Cur-rently, the influence of triptolide on podocytes is consid-ered to be an important factor for the decrease of urine protein of patients with glomeru-lonephritis [8,9]. With the research on the influence of glomerular podocytes on proteinuria production, people have recognized that for some diseases of the renal glomeruli which are charac-terized in a great quantity of proteinuria and accompa-nied by the damage to podocyte such as minute lesion, focal segmental glomerulo-sclerosis and membranous nep-hropathy, TW works well on reducing proteinuria to all the above diseases [9,12].

However, the problems exist: hormonotherapy which is necessarily to be used in combination with TAC and TW in the treatment of SRNS always cause acne, central obesity and other side effects because of the high dose of hormone and long period of application [13-17]; In addition, high-dose TAC and double-dose TW also can cause moreside effects [4,9]. Therefore, first of all, among do-mestic institutions, we studied the feasibility of the treatment of SRNS with low-dose TAC (blood concentra-tion 1.5 - 4 ng/ml) combined with single-dose TW.

The research results are: 3 months after treatment with low-dose TAC combined with TW, there were 2 cases of complete remission out of 11 cases (18.2%), and 7 cases of partial remission (63.6%); 6 months after treatment there were 3 cases of complete remission out of 11 cases (27.3%), and 7 cases of partial remission (63.5%); when the follow-up visits were carried out till the 12th month, 8 cases were completely remitted out of 11 cases (72.7%) and 2 cases were partially remitted (18.2%). The complete remission rate was obviously higher than that of TW group. In addition, our research also discovered that after the treatment with TAC combined with TW, the plasma protein of SRNS patients increased obviously, and when treatment was carried out for 6 months, it ba-sically recovered to the normal level; despite that urine protein of TW group decreased and plasma albumin in-creased, the increase range was significantly lower than that of TAC + TW group. Therefore, it is indicated that the treatment of SRNS with low-dose TAC combined with TW would reach the expected efficacy.

The main renal histopathological symptoms of patients with primary nephrotic syndrome are: minimal change disease (MCD), focal segmental glomerulosclerosis (FSGS), membranous nephropathy (MN), MSPGM and MPGN. Current researches have showed that MCD is caused by damage to podocytes which is caused by nephrotoxic cell factors mediated and generated by abnormal immune
response of T cells; FSGS is not only caused by abnor-
mal permeability of slit membrane of podocytes which is
casted by genetic defect, the occurrence of some FSGS
is similar to that of MCD, which is related to the abnor-
mal immune response of T cells; and MesPGN is a path-
ological change involving many Immune factors [18-21].
Therefore, we applied TAC to reduce the generation of
immune complex by inhibiting the abnormal immune
response of T cells; meanwhile, TW was combined not
only to strengthen TAC anti-inflammation and anti-im-
munization, but also to reverse the damage to podocytes
and recover the skeleton structure of podocyte and the
expression of relevant molecules, so as to realize better
efficacy [18]. Moreover, it is a better choice for patients
who have clinical side effects caused by hormone as well
as those who are hormone-phobic.

For the obvious increase of plasma protein of SRNS
patients who received the treatment with low-dose TAC
combined with TW, the plasma protein basically recov-
ered to the normal level when treatment was carried out
for 6 months; but the plasma protein of TW group was
not obviously increased. One potential mechanism may
be that TAC can assist the liver to produce albumin;
However, TW has inhibiting effect on the synthesis of
albumin of liver, and the exact mechanism is to be ex-
plored [4,12].

Common side effects of TAC include: gastrointestinal
reactions, immunosuppression, infection, elevated blood
pressure, anemia, renal toxicity, increased blood sugar,
etc. [4,22]. Adverse reactions of TW mainly include nau-
sea, vomit, anorexia, abdominal distention, reduction of
leukocyte, liver dysfunction and influences on menstrual
cycle of female patients of childbearing age [9,12,17].
In our research, side effects in low-dose TAC + TW group
included 2 cases of nausea and 1 case of herpes zoster, 3
cases of slightly increase of glutamic-pyruvic transami-
nase after treatment of 1 month, and 2 cases menstrual
disorder out of 4 female cases; there was no discovery of
serum creatinine exceeding 50% of the base line, and the
fasting blood-glucose did not increase obviously. Patients
with increased transaminase were given bifendate for
symptomatic treatment and patients’ conditions improved,
all patients kept taking medicines and the treatment and
follow-up visits were not affected. Our research indicated
that the treatment of SRNS with low-dose TAC com-
bined with TW would not generate serious side effects.
In conclusion, low-dose TAC + TW group may effec-
tively reduce the proteinuria of SRNS patients, increase
the level of plasma albumin, has obvious effective remis-
sion rate, does not obviously increase side effects, which
indicate that low-dose TAC + TW may be applied to the
treatment of SRNS. However, since this research is a sin-
gle-center research, and the number.

REFERENCES

oid-Sensitive and Steroid-Resistant Idiopathic Nephrotic
Syndrome,” Pediatric Nephrology, Vol. 26, No. 6, 2011,

[2] X. F. Sun and X. M. Chen, “Clinical Application of Tac-
rolimus in Primary Nephrotic Syndrome,” Chinese Jour-
nal of Practical Internal Medicine, Vol. 28, No. 5, 2008,
pp. 323-325.

Nephrotic Syndrome with Tacrolimus,” The Journal of

liminary Study of Tacrolimus (FK506) Combined with
Low-Dose Prednisone in Patients with Membranous Lu-
pus Nephropathy,” Chinese Journal of Nephrology, Di-
alysis & Transplantation, Vol. 13, No. 2, 2004, pp. 101-
106.

in the Management of Transplant-Related Nephrotic Syn-
drome and Steroid-Resistant Nephrotic Syndrome,” Trans-
plantation Proceedings, Vol. 23, No. 6, 1991, pp. 3354-
3356.

Safety of Tacrolimus versus Cyclosporine in Children
with Steroid-Resistant Nephrotic Syndrome: A Random-
ized Controlled Trial,” American Journal of Kidney Dis-
doi:10.1053/j.ajkd.2008.11.033

with Steroid- and Cyclophosphamide-Resistant Nephrotic
Syndrome and Normal or Mildly Reduced GFR,” Ameri-
can Journal of Kidney Diseases, Vol. 54, No. 1, 2009,
pp. 51-58. doi:10.1053/ajkd.2009.02.018

duces Proteinuria in Experimental Membranous Neph-
ropathy and Protects against C5b-9-Induced Podocyte In-
pp. 974-988. doi:10.1038/ki.2010.41

ous Nephropathy with Tripterygium Combined Low-
Dose Hormone: A Prospective Randomized Control Trial,”
Chinese Journal of Nephrology, Dialysis & Transplanta-

with Steroid-Resistant Nephrotic Syndrome Treated with
Tacrolimus,” Pediatric Nephrology, Vol. 25, No. 6, 2010,
pp. 1117-1124. doi:10.1007/s00467-010-1471-8

New Therapy for Steroid-Resistant Nephrotic Syndrome

ggium in Patients with Diabetic Nephropathy: A Prospec-
tive Randomized Control Clinical Trial,” Chinese Journal
of Nephrology, Dialysis & Transplantation, Vol. 19, No.

Copyright © 2012 SciRes.


