Nephroprotection in Unani Medicine through Herbal Medicine & their research scope: A Review

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ABSTRACT

Plants are very potential & vital source for human health and also used therapeutically in the various kidney disorders due to presence of phytochemical constituents. There are numerous herbs existing that have more pharmacological activities including with nephroprotective activity. Now days herbal drugs are verified & proved as nephroprotective agents and used for improving renal health, even reverse renal damages. The current review is targeted to explain the Unani & modern concept of kidney disorders, nephrotoxicity and the list of scientifically proved medicinal plants having nephroprotective activity medicinal plants which used in the treatment of renal disorders.

Keywords: Nephroprotection, Unani Medicine, Kidney disorders

INTRODUCTION:

The waste products produced in the body due to various metabolic activities must be excreted from the body (1). If they are allowed to accumulate they are injurious to health (2). Kidney is an important organ of the body, which has a natural function of excretion of waste products from the body. Apart from this it controls the volume and composition of the body fluids, for water and virtually all electrolytes in the body, balance between intake and output is maintained in large part by the kidneys (3). Kidneys are most important channel of excretion for majority of drugs and its metabolites (4), so it always opens up to receive the drugs and other toxins which cause renal damage or renal toxicity (5). Prominent Unani scholars were very much concern about renal disorders. Chronic kidney failure has been described in the classical Unani literature as za'uf Kulya (6), (7), (8), (9), (10). In Unani System of medicine kidney diseases are caused due to three reasons; Su'i Mizaj-i Kulya, Su'i Tarkeeb, Ta Farruq wa Ittesal (11). There are a unique concept of muqawiyat (tonics) in unani system of medicine. According to Ibn-e-Rushd (1980) functions of the kidney depends upon its Quwwa. Whenever any Quwwa becomes weak kidney disorders appear. These Quwwa are as follows, Quwwat-e-Hazima, Quwwat-e-Jaziba, Quwwat-e-Masika, Quwwat-e-Dafia, Quwwat-e-mumayyazaa. (7). So herbal drugs are capable to tune up the kidney or other important organs as well as also provide strength to them. In modern medicine concept of kidney disorders can be defined as “a heterogeneous group of disorders which affected the kidney structure and its functions. Now a days, it is accepted that even minor abnormalities in measures of kidney structure and function are related with higher risk for other organ systems as well as mortality, all of which happen far more frequently than kidney failure. Acute & chronic kidney failure comes in the category of kidney disease which can be life threatening. Reduction in GFR (Gomerular filtration rate) that decreases renal excretory functions is called as renal failure. This is conveyed to a changeable variable by failure of erythropoietin production, vitamin D hydroxylation, and regulation of acid-base balance, regulation of salt & water balance and blood pressure (12). Nephrotoxicity occurs when kidney-specific detoxification and excretion do not work properly due to the damage or destruction of kidney function by exogenous or endogenous toxicants (effect of toxic chemicals or drugs) (13), (14). Plants are very potential & vital source for human health and also used therapeutically in the various kidney disorders due to presence of...
phytochemical constituents. There are numerous herbs existing that have more pharmacological activities including nephroprotective activity. Now days herbal drugs are verified & proved as nephroprotective agents and used for improving renal health, even reverse renal damages. The present review is aimed to elucidate the list of nephroprotective medicinal plants, which are scientifically proved in treating renal disorders.

### Unani Perspective of Kidney Diseases:

According to Ibn-e-Rushd (1980) functions of the kidney depend upon its Quwwa. Whenever any Quwwa becomes weak kidney disorders appear. These Quwwa are as follows:

- Quwwat-e-Hazima
- Quwwat-e-jaziba
- Quwwat-e-Masika
- Quwwat-e-Dafia
- Quwwat-e-mumayyaza

In Unani System of medicine kidney diseases are caused due to three reasons; (15), (7):

1. **Su’i Mizaj-i- Kulya (Renal dyscrasia):** any alteration in temperament of kidney from normal called as *Su’i Mizaj-i- Kulya*. If alteration is found in blood (dam) or bile (sofra) called as *Su’i Mizaj-i- Kulya Haar* (Renal hot dyscrasia) and if alteration in phlegm (balgham) or black bile (sauda) known as *Su’i Mizaj-i- Kulya Barid* (Renal cold dyscrasia) due to this altered temperament there are following diseases found in kidney such as:

   - Huzāl al-kulya (Reduction in the size of kidney or Renal hypotrophy)
   - Du’āl-Kulya (Weakness of kidney)
   - Dhayabitus Hārr (Diabetes Mellitus), Dhayabitus bārid (Diabetes Insipidus)
   - Hasāh al-kulya (Renal caculus)
   - Sudad al-kulya (Renal obstruction)
   - Waram al-kulya (Nephritis)
   - Dubayala al-kulya (Renal Abscess)

2. **Su’i Tarkeeb:** any variation in the formation of kidney at the time of intra uterine life (abnormality in genome), *Amraze shakal* such as *Marz Akyas-v-e Kulya* (Polycystic kidney disease), *Amraze miqdar* such as *Huzāl al-kulya* (Renal hypotrophy), *Amraze adad* such as Renal agenesis.

3. **Tafarruq wa Ittesal (Renal Damage):**

   - Quruh al-kulya (Renal ulcers)
   - Jarab al-kulya (Renal irritability)
   - Rih al-kulya
   - Waja’ al-kulya (Renal pain)

### Amraaze Kulya (Renal Diseases), Causes, Sign & Symptoms

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Name of the disease</th>
<th>Causes</th>
<th>Sign &amp; Symptoms</th>
</tr>
</thead>
</table>
| 1      | Huzāl al-kulya (Reduction in the size of kidney or Renal hypotrophy) | Impaired temperament (hot or cold) of kidney | -Urine become colorless
         |                     | Weakness of digestive faculties (Quvvate hazima) | -Poly urea |
| 2      | Du’āl-Kulya (Weakness of kidney) | Impaired temperament | Frequency of urine increases
         |                     | Dilatation of calyces | Oliguria
         |                     | Excessive use of diuretics, More physical efforts. | Loss of libido
         |                     | | Headache |
| 3      | Dhayabitus Hārr (Diabetes Mellitus), Dhayabitus bārid (Diabetes Insipidus) | Impaired hot temperament | Excessive thirst
         |                     | Expulsive faculty (Quvvate da’fe) becomes strong & weakening of holding faculty (Quvvate masika) | Excessive & colorless urination
         |                     | Dilatation of calyces | |
| 4      | Hasāh al-kulya (Renal caculus) | Retaining of Viscous humor inside the kidneys | Pain in lumber region
         |                     | Weakening of expulsive faculty of kidney | associated with vomiting
         |                     | Any hindrance due to kidney’s impaired temperament | Frequency of urination increases
         |                     | | Dysuria & anuria |
         |                     | | Colorless urination |
| 5      | Sudad al-kulya (Renal obstruction) | Accumulations of mucilaginous viscous humors | Pain with renal heaviness
         |                     | Inflammations (Warm al-kulya) | Decreased urine output
         |                     | | Fever due to inflammation |
| 6      | Waram al-kulya Hārr (Acute Nephritis) | Viscous blood (Ghaliz Dam), diluted bile (Raqeeq Safra) or pure bile (Khalis safra) | Fever with chills & rigor
         |                     | suppurated Phlegm (Muta’affin Balgham) or | Pain & heaviness at the site of inflammation
         |                     | suppurated blood & Phlegm both | Irritation & restlessness |
         |                     | | Dysuria or Anuria |
| 7      | Waram al-kulya Bārid (Chronic Nephritis) | Predominance of Phlegm (Balgham) & Black bile (Sauda) | Mild Fever
         |                     | | Edema |
Kidney disease differs from the diseases of other organ systems, these diseases habitually "quiet" or sometimes having limited symptoms. The symptoms of renal diseases remains often nonspecific and until late in the course of disease. Thus, there are few kidney-specific "clinical events," which results in a dependence on laboratory procedures to define the major clinical syndromes, so documentation or implication of the duration of abnormalities in laboratory measures are suggested the duration of kidney disease.

### Principles for the Definitions of Kidney Diseases and Disorders (18), (19)

<table>
<thead>
<tr>
<th>Disease/Disorder</th>
<th>Functional criteria</th>
<th>Structural Criteria</th>
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<tbody>
<tr>
<td>Acute kidney Injury (AKI)</td>
<td>Increase in S.Cr by 50% within 7 d, or increase in S.Cr by 0.3 mg/dL within 2 d, or oliguria</td>
<td>No criteria</td>
</tr>
<tr>
<td>Chronic Kidney Disease (CKD)</td>
<td>GFR &lt; 60 mL/min for &gt;3 mo.</td>
<td>Kidney damage for &gt;3 mo.</td>
</tr>
<tr>
<td>Acute Kidney Injury (AKD)</td>
<td>AKI, or GFR &lt;60 mL/min/1.73 m2 for &gt;3 mo. or decrease in GFR by ≥35% or increase in S.Cr by &gt;50% for &lt;3 mo.</td>
<td>Kidney damage for &lt;3 mo.</td>
</tr>
<tr>
<td>No known kidney disease (NKD)</td>
<td>GFR ≥60 mL/min/1.73 m2, stable S.Cr</td>
<td>No damage</td>
</tr>
</tbody>
</table>

The following most common Kidney disorders and include (20) (19)

- Acute kidney injury (AKI) or Acute renal failure (ARF)
- Nephritic syndrome
- Nephrotic syndrome
- Tubulointerstitial disease
- Vascular disease of the kidney
- Papillary necrosis
- Chronic kidney disease (CKD)

### Nephrotoxicity:

Kidneys are performed many physiological functions of the body such as protection of homeostasis as well as regulation of extracellular atmosphere for example detoxification and elimination of toxic substances (metabolites) and therapeutic agents. These functions are more challenging task for the kidney, that's why kidneys are the most important passage for the entry of any exogenous toxicant into human body. Nephrotoxicity occurs when kidney-specific detoxification and excretion do not work properly due to the damage or destruction of kidney function by exogenous or endogenous toxicants (effect of toxic chemicals or drugs) (13), (14). Nephrotoxins differ in their chemical structure, having different cellular targets and mechanism of cellular injury within the kidney and all of them produce varying degree of kidney damage.

There are many drugs which are responsible for impairment of kidney functions & leads to renal perfusion or direct nephrotoxicity including antibiotics such as aminoglycoside, vancomycin, antifungal such as amphotericine B, immunomodulators such as calcineurin inhibitors, chemotherapeutic agents, cyclosporine, tacrolimus etc. Kidney injury can also occurs within days of initiating therapy with antihypertensive drugs such as ACE inhibitors, β-blockers, NSAIDS / selective COX-2 inhibitors, Cocaine, occupational toxins (21), (22). Approximately 20 % of hospitalized and non-hospitalized cases are suffering from acute renal failure (ARF) due to drug induced nephrotoxicity (23), (24), (25). The prevalence of drug-induced nephrotoxicity may be 66 percent high in the older adults (26).
Mechanisms of actions of drug induced nephrotoxicity:

Drug induced nephrotoxicity can be explained by the following different pathogenic mechanism such as (27), (28).

- Changed intra-glomerular hemodynamics,
- Tubular cell toxicity,
- Inflammation,
- Crystal nephropathy,
- Rhabdomyolysis,
- Thrombotic microangiopathy

I. **Changed intra-glomerular hemodynamics**: The pathogenic mechanism of changed by the following flow chart;

```
Anti-prostaglandin drugs       Anti-angiotensin-II

Interfere with autoregulation of glomerular pressure

Vasoconstriction

Decreased blood flow

Ischemic injury

Decreased GFR

• Nephrotoxic Drugs: NSAIDs, ACE Inhibitors, Cyclosporine, Tacrolimus (29),

II. **Tubular cell toxicity**: The following flow chart explained the pathogenic mechanism of tubular cell toxicity; (30), (31)

Nephrotoxic drug (Amphotericin B, antiretrovirals)

Impairing mitochondrial function

Interfering with tubular transport,

Increasing oxidative stress or free radicals formation

Tubular cell Toxicity

• Nephrotoxic drugs: Antimicrobials, Amphotericin B, Beta-lactam antibiotics, Rifampicin, Adefovir, Cidofovir, and Contrast dye, Zoledronate (32), (33), (34), (35), (31), (36)

III. **Crystal nephropathy**: Nephrotoxic drugs caused renal impairment by the production of urine insoluble crystals within the distal tubular lumen explained following flow chart (31),

Urine insoluble Crystal’s production

Precipitations of crystals (within the distal tubular lumen)

Obstruction in urine flow

Production of an interstitial reaction

Renal impairment

Crystal nephropathy
• **Nephrotoxic Drugs**: Fosinopril, Methotrexate, Triamterene, Ganciclovir (35), (31), (37).

IV. **Rhabdomyolysis**: It is the condition in which rapidly damaged & break down of skeletal muscles tissue, these damaged skeletal muscles cells such as myoglobin released into bloodstream and causes kidney injury. The pathogenic mechanism explained through following flow chart (38)

![Flow Chart for Rhabdomyolysis](image)

- Skeletal muscle injury
- Lysis of the myocyte
- Releasing intracellular contents (myoglobin and creatine kinase) into the plasma.
- Renal injury

(Direct renal toxicity, tubular obstruction, and modifications in GFR.)

- **Nephrotoxic drugs**: Amitriptyline, Diphenhydramine, Doxylamine, Benzodiazepines, Statins, Methotone (39), (38)

V. **Thrombotic Microangiopathy**: Thrombotic microangiopathy is due to formation of thrombi in the microcirculation which leads to renal injury, following flow chart described pathogenic mechanism of thrombotic microangiopathy (40)

![Flow Chart for Thrombotic Microangiopathy](image)

- Formation of thrombus in the microcirculation
- Immune-mediated reaction
- Direct endothelial toxicity

- **Nephrotoxic drugs**: Cyclosporin, Clopidogrel, Mitomycin C, Quinidine (41), (40)

VI. **Inflammation**: Inflammatory changes occurs mostly in the glomerulus, renal tubular cells, and the surrounding interstitium due to nephrotoxic drugs & fibrosis and renal scarring takes place. Glomerulonephritis is an inflammatory condition caused primarily by immune mechanisms and is often associated with proteinuria in the nephrotic range (35).

- **Nephrotoxic drugs**: Allopurinol, Aspirin, Acetaminophen, Ayclovir, antibiotics, Beta-lactam, Chinese medicines Quinolones, Rifampicin, Sulfonamides, Cisplatin, Loops & thiazide diuretics (35), (31), (42)

Research scope of herbal medicine:

Plants are very good for human health and used therapeutically in the various kidney disorders because of their chemical components and medicinal properties (43). Lots of herbs have been proven to be capable as Renoprotective agents but some of them are claimed to be Nephrotic. There are so many researches available which proves the Nephroprotective effect of these herbs such as

1. **Nephroprotective Action of Phoenix dactylifera in Gentamicin-Induced Nephrotoxicity**

Aqueous extract of date flesh has a prophylactic action on gentamicin induced nephrotoxicity in pretreated animals, proved by a significant decrease in the plasma concentrations of creatinine and urea & in an increase in the body weight of the animals during the treatment period. Aqueous extract of date flesh also has a curative effect when it was given combined with gentamicin which proved by a significant decreased in increased biochemical parameters. Acute and chronic toxicity study also has been done by an oral administration of graded doses of the aqueous extracts of the date flesh and depths to male Wistar rats, proved that the given extract was safe for the animal (44).

2. **Nephroprotective Effect of Kahab Chini (Piper cubeba) in Gentamicin-Induced Nephrotoxicity**

The pre- and post-treated animal groups was taken powder of test drug in suspension form at 810 mg/kg and 1220 mg/kg orally. The assessment parameters for nephroprotective effect are based on biochemical markers such as serum urea and creatinine levels as well as histopathological analysis of the treated kidney. Biochemical markers of renal function were found to be significantly lower in the pre-treated group in comparison the control group, while mild tubular damage shown in histological analysis. In the same way, there were a significant decrease in serum urea and creatinine levels in the post-treated animals in comparison to negative control group, moderate degree of tubular necrosis seen in the histopathological examination. So KC has a significant amount of nephroprotective activity which proved by the examination of biochemical markers of kidney function and the histopathological features pre-treated and post-treated groups (45).

3. **Nephroprotective Action Of Peucedanum Grande Against Cadmium Chloride Induced Renal Toxicity In Wistar Rat**

Nephroprotective effect of P. grande against CdCl2 induced nephrotoxicity was assessed by the estimation of biochemical markers, antioxidant enzyme activities and histopathological changes. Animals were taken pre-treatment of P. grande at a dose of 60 and 120 mg/kg b.wt orally against the renal nephrotoxicity produced by CdCl2 at a dose of 3mg/kg b.wt. CdCl2 induced deteriorative effects were prevented by Pretreatment with P. grande, through a protective mechanism, which involved Improved deterioration i.e. increased oxidative stress as well as
histopathological changes i.e. damaged renal tissues against CdCl2 administration. BUN, creatinine significantly lower in pretreated rats with *P. grande* in comparison to that animals which getting only CdCl2. The present study concluded *P. grande* has potent nephroprotective activity through a protective mechanism & this activity of *P. grande* might be due to its free radical scavenging activity (46).

4. Diuretic and nephroprotective effect of Jawarish Zaroooni Sada—a polyherbal unani formulation

Ethanol and aqueous extracts of jawarish zaroooni sada (JZS) at 300 mg each were examined as diuretic activity by calculating the total urine output over a period of 6 h in the present study. Sodium and potassium level in urine sample was also estimated. Nephrotoxicity was investigated by administration of JZS along with gentamicin at high dose (40 mg/kg) and elevated biochemical markers such as urea and serum creatinine he study proved that JZS has significant diuretic and nephroprotective properties (47).

5. Protective effects of 'Khamira Abresham Hakim Arshad Wala', a unani formulation against doxorubicin-induced cardiotoxicity and nephrotoxicity

Doxorubicin produced a marked nephrotoxicity, as shown by in the elevation serum enzymes such as of AST, LDH, kidney biochemical markers i.e. BUN, creatinine, and, depletion in stress marker such as MDA, GSH level & catalase activity Khamira abresham Hakim Arshad wala showed nephroprotective effect against doxorubicin-induced nephrotoxicity, there was a significant enzymatic changes in serum as well as damaged renal tissues in pretreated group of animals. The results of present study reveals that the elevation & depletion in above serum enzymes, biochemical and stress markers as well renal damage were significantly (p<0.01) improved after the treatment of Khamira Abraham Hakim Arshadwala (48).

6. Nephroprotective Effect of Echinodorus macrophyllus Micheli on Gentamicin-Induced Nephrotoxicity in Rats

Current study on "Nephroprotective Effect of Echinodorus macrophyllus Micheli on Gentamicin-Induced Nephrotoxicity in Rats" shown that gentamicin induced nephrotoxicity increased biochemical markers such as polyuria, decreased in GFR & morphological changes also. On treatment with extract *Echinodorus macrophyllus Micheli* (EM) along with gentamicin reversed all above altered parameters & morphological changes were also not detected. EM extract also produced a dose-dependent decrease in urine elimination. So the study concluded that EM has nephroprotective potential & EM can used therapeutically an incidence of GM-induced acute kidney injury (49).

7. Phytochemical, pharmacological evaluation of Morinda pubescens J.E.Sm. Bark extract for nephroprotective activity

Water extract *Morinda pubescens* J.E. Sm. (WEMp) of silymarin, and its isolated ethyl acetate fraction compound (ISLTD mp-B) significantly increased the biochemical markers of kidney such as albumin, protein, body weight, and urine volume & decreased kidney serum creatinine, urea, uric acid, blood urea nitrogen, kidney weight with a significant reduction in gentamicin induced toxicity. The study also reveals that (WEMp) and (ISLTD mp-B) treated group increased urine output than normal which helped to dilutes the concentration of kidney function parameters such as S. creatinine, urea, uric acid, and blood urea nitrogen . Thus, kidney's toxins, increased biochemical markers or waste products are flushed out via the urine and reduced the chances or prevent the renal toxicity. The Renal protective activity of bark extracts may be due to the antioxidant activity, due to the existence of secondary metabolites i.e. tannins, flavonoids and phenolic compounds, etc. (50).

8. Evaluation of Nephroprotective Activity of Ethanolic Extract of Annona reticulata in Gentamicin and Cisplatin Induced Nephrotoxicity in Rats

In the study “Evaluation of Nephroprotective Activity of Ethanolic Extract of Annona reticulata in Gentamicin and Cisplatin Induced Nephrotoxicity in Rats” toxicant group (2nd and 5th) gentamicin & cisplatin treated has severe nephrotoxicity evidence by an increment concentration of serum, urea, Creatinine, Uric acid, Total protein and Urine Urea, uric acid, creatinine in comparison to normal control (group 1). The increased biochemical parameters improved by the treatment of ethanolic extract of aerial parts of *Annona reticulata* & showed Nephroprotection because of presence of phyto chemical constituents (51).


Treatment with the A. cepa Linn. Has shown significant (p<0.01 and p<0.001) dose-dependent improvement in the body weight at the dose of 200 and 400 mg/kg and also shown significant improvement by protecting the kidney from the oxidative stress. It is also identified that treatment with A. cepa significantly lowered the level of serum creatinine, total protein when compared with the toxic group. Conclusion: Nephroprotective activity of EEAC treatment was found compared with the standard group (Vitamin E – 250 mg/kg) and control group against the toxic control group animals in parameters including serum creatinine, total protein, kidney weights, and body weights. The histopathological studies were also evinced the protective effect of EEAC, (52).

10. Nephroprotective Effect of Aqueous Extract of Pimpinella anisum in Gentamicin Induced Nephrotoxicity in Wistar Rat

Co-administration of Pimpinella anisum extract with gentamicin decreased the following rise parameters such as serum urea, serum uric acid, and serum creatinine and blood urea nitrogen in a dose dependent manner. Gentamicin treated group shown nephrotoxicity proved by elevated serum urea, serum uric acid, serum creatinine and blood urea nitrogen (107.5±16.92mg/dl, 0.8±0.09 mg/dl, 3.05±2.29 mg/dl, and 47.8±9.07 mg/dl) respectively in comparison to the saline treated groups. Gentamicin treated group also shows histopathological changes such as epithelial loss with deep granular degeneration in rats, while water extract of Pimpinella anisum decreased the severity of gentamicin-induced renal damage. So the study concluded, that water extract of Pimpinella anisum reveals nephroprotective effect in gentamicin induced renal damage (53).

11. Nephroprotective Activity of Methanolic Extract of Lantana camara and Squash (Cucurbita pepo) on Cisplatin-Induced Nephrotoxicity in Rats and Identification of Certain Chemical Constituents of Lantana camara by HPLC-ESI-MS.

Methanolic (defatted) extract of the two plants (*Lantana camara* and *Cucurbita pepo*) have nephroprotective effect on cisplatin-induced nephrotoxicity in rats. Ethyl acetate and
butanolic derivative of methanolic extract of L. camara also had high nephroprotective activity. The study also revealed that the nephroprotective effect of these extract because of the presence secondary metabolites or plants phytochemical constituents such as phenolic acid derivatives, flavonoids, phenylenethanoids and Iridoids etc and phytochemicals analyses by the use of HPLC-ESI-MS technique (54).

12. Nephroprotective Effect of Sonchus oleraceus Extract against Kidney Injury Induced by Ischemia-Reperfusion in Wistar Rats

Sonchus oleraceus Extract (S.O.e) significantly reduced the higher level of kidney biochemical parameters such as urea nitrogen (BUN), creatinine, malondialdehyde (MDA), and pro-inflammatory cytokines & lowered the SOD level in Ischemia-Reperfusion (I/R) treated animal groups than the sham group. Tubular epithelial necrosis in the medulla and cortex were seen on histopathological examination in I/R treated group. While pretreatment with S.O.e & treatment in combination of S.O.e and I/R reduced the I/R induced increased renal biochemical parameters included BUN, creatinine, MDA, and pro-inflammatory cytokines induced, SOD as well as discontinuous necrosis in the medulla but no necrosis in the cortex on histopathological examinations. So the present study concluded that S.O.e has neither nephrotoxicity nor hepatotoxicity in spite of Nephroprotection pretreatment with S.O.e (55).

13. Nephroprotective Effect of Camel Milk and Spirulina platensis in Gentamicin-Induced Nephrotoxicity in Rats

Gentamicin treated animal group significantly showed increased levels of urea, creatinine and malondialdehyde, decreased level of glutathione (GSH) & extensive tubular necrosis present in histopathological examination. While extract of Camel Milk (CM) and Spirulina platensis (CP) and their combination (SPCM) treated animal group showed a significant improvement in these increased biochemical parameters, stress marker & histopathological changes (56).

14. Phytochemical and Nephroprotective Activity of Eclipta prostrata against Gentamicin Induced Nephrotoxicity in Wistar Rats

In the present study "Phytochemical and Nephroprotective Activity of Eclipta prostrata against Gentamicin Induced Nephrotoxicity in Wistar Rats" showed a significant nephrotoxicity in nephrotoxic group (Group II) by increased in S. urea, S. creatinine, and S. uric acid, BUN & kidney weight, while a major reduction has been shown in all these parameters in extract treated group which compared with standard (Cystone) group (Group V). Preliminary phytochemical screening of leaves of Eclipta prostrata reveals the presence of secondary metabolites such as terpenoids, glycosides, alkaloids, sterol, flavonoids, volatile oils and Saponins etc (57).

15. Acute nephroprotective and antioxidant activities of aqueous leaf extract of Plecetranthus ambionicus (Roxb.) grown in Sri Lanka

The current study has been shown that ADR produced nephrotoxicity by the elevated biochemical parameters such as S. in rats. The elevated S. creatinine level significantly decreased after treatment with aqueous extract of leaf of P. ambionicus (400 mg/kg body wt.) in comparison to control group. Glomerular and tubular functions of the kidney was measured by the β2 -microglobulin. Concentration of β2 -microglobulin increased 50% in the control (nephrotoxic) group rats in comparison to the control group (healthy) (p<0.05). The results of the study concluded that, P. ambionicus is reduced very fast the raised level of β2 – microglobulin in the comparison & extract of plant leaf also reduced the increased levels of S. Albumin, total protein as compared to standard drug, proved nephroprotectivity of the leaf extract of P. ambionicus against ADR induced nephrotoxicity (58).

16. Antioxidant and nephroprotection activities of Combretum micranthum G. Don in cisplatin induced nephrotoxicity in rats: In-vitro, in-vivo and ex vivo studies

End stage renal disease is a most common & serious complications of diabetic nephropathy. A significant dose dependent inhibition (P < 0.001) was found in lipid peroxidation induced by ferric chloride-ascorbic acid after treatment with CM extract. Human embryonic kidney cell (HEK-293) was used as in-vitro model for glucose induced toxicity for diabetic nephropathy in this study. The present study revealed that a significant morphological changes such as cell shrinkage, rounded cell shape and cytoplasmic vacuolation after taking the high glucose (100 mM) for 72 h, but the cell viability (10 to 23%) improved after taking treatment with 1 and 25 mg/mL CM extract in comparison to high glucose control. The results of study proved that C. micranthum has potent antioxidant and nephroprotective activity (59).

17. Nephroprotective activity of Combretum micranthum G. Don in cisplatin induced nephrotoxicity in rats: In-vitro, in-vivo and in-silico experiments

Recent study on "Nephroprotective activity of Combretum micranthum G. Don in cisplatin induced nephrotoxicity in rats: In-vitro, in-vivo and in-silico experiments" reveals that Cisplatin produced a significant increase in cell death with changes in normal cellular morphology in HEK-293 cells. The study also revealed that CP and CM extract treated HEK-293 cells shows a significant improvement in cell growth which indicate CM-extract has high cytoprotective activity in comparison to CP extract while CP-treated rats showed a characteristic clinical and pathological symptoms i.e. increased relative kidney weight, altered kidney function parameters such as creatinine, urea, uric acid, total protein, albumin and electrolytes against cisplatin induced cytotoxicity, hence proved that CM extract contain Nephroprotective activity (60).

Conclusions:
The above review concludes that herbal drugs are very useful for kidney health. It is mentioned in classical Unani literature that various single as well as compound formulations are used extensively as diuretics (Mudir e baul), kidney and bladder tonics ( Muqawwi e Gurdwa wa Masana) and lithotriptic (mufattit hisaat) since ages. A number of researches has been performed in this field but further validation with proper pre-clinical & clinical trials are needed so the herbal formulations as well as the single drugs could be accepted globally and help the mankind with least adverse effects in various renal disorders.

References:
5. M.A.P. Renal vulnerability to drug toxicity. Clinical Journal of the
10. MA. K. Iseere Azam (Urdu translation) Rawalpindi Pakistan: Tibbi company; 1940.
15. Roohi Zaman AAMJGSA. Journal of Pharmacognosy and Phytochemistry..


54. Abdel-Hady H, ESMM, AHA AHMM, AHAMAT, AHIES, ALESA MEA. Nephroprotective Activity of methanolic extract of Lantana camara and squash (Cucurbita pepo) on cisplatin-induced nephrotoxicity in rats and identification of certain chemical constituents of Lantana camara by HPLC-ESI. Pharmacognosy Journal. 2018; 10(1).


73. Ahmad F ASAA0AMA 2, 6(2):4. Phytochemical and Nephroprotective Activity of Eclipta prostrata against Gentamicin Induced Nephrotoxicity in Wistar Rats. International Journal of Pharma Research and Health Sciences..