Use of Cyclosporine Therapy in Steroid Resistant Nephrotic Syndrome (SRNS): A Review

Syed Raza Shah¹, Areeba Altaf¹, Mohammad Hussam Arshad², Anum Mari³, Sahir Noorani⁴, Eraz Saeed⁴, Areeb Amir Mevawalla³, Zaiyn Ul Haq³ & Muhammad Ehsan Faquih¹

¹ Department of Medicine, Dow University of Health Sciences (DUHS), Karachi, Pakistan
² Department of Medicine, Aga Khan University of Health Sciences, Karachi, Pakistan
³ Department of Biological Sciences, The Lyceum, Karachi, Pakistan
⁴ Department of Biological Sciences, Karachi Grammar School, Karachi, Pakistan

Correspondence: Syed Raza Shah, Babae Urdu Road, Dow Medical College, Dow University of Health Sciences (DUHS), Karachi, Pakistan. Tel: 92-345-245-4610. E-mail: syedraza91shah@live.com

Received: June 14, 2015   Accepted: July 17, 2015   Online Published: August 6, 2015

doi:10.5539/gjhs.v8n4p136          URL: http://dx.doi.org/10.5539/gjhs.v8n4p136

Abstract

A chronic, progressive disorder Steroid Resistant Nephrotic Syndrome (SRNS) accounts for 10-20% of all children with Nephrotic Syndrome. It is a heterogeneous disorder comprised of persistent edema, proteinuria, hypoalbuminemia and hyperlipidemia. Treatment for steroid-resistant nephrotic syndrome (SRNS) is challenging and children who suffer from SRNS require aggressive treatment to achieve remission. Calcineurin inhibitors have been used more in an empirical manner than on the basis of clear rationale. It was in 1984 when cyclosporine was first considered for the treatment of steroid resistant nephrotic syndrome. Cyclosporin is a calcineurin inhibitor that suppresses immune response by downregulating the transcription of various cytokine genes. Till now many studies have been conducted to determine dosages, duration of therapy, side effects and advantages of cyclosporine. Treatment of SRNS remains a difficult challenge in pediatric nephrology. Treatment should be individualized according to the underlying histopathology, and clinical and environmental conditions of the children. There is an urgent need to distinguish as soon as possible those patients who may benefit from prolonged immnosuppressive treatment from those who will not benefit from such treatment and who will just suffer from its major side effects. The emerging evidence that the majority of genetic forms of SRNS should receive symptomatic treatment only, should also be clinically tested and studies baring its significance should be evaluated in the future.

1. Background and Introduction

A chronic, progressive disorder Steroid Resistant Nephrotic Syndrome (SRNS) accounts for around 10 to 20% of all children with Nephrotic Syndrome (Banerjee, 2002). It is a heterogeneous disorder comprised of persistent edema, proteinuria, hypoalbuminemia and hyperlipidemia. Various studies have shown that over 80% of children with initial episode of Nephrotic Syndrome respond well to steroids but almost 10-20% do not respond to steroids and are known to have SRNS for which thereapeutic other than steroids are used as a treatment. (Bhimma, 2005) Treatment primarily includes steroids but when there is no remission, potent immnosuppressants are used. It has been observed that certain factors have known to contribute to steroid resistance which include hypertension, hematuria, hypertension plus hematuria, proteinuria(>10 g/day), elevated plasma creatinine, black race, presenting in infancy, renal biopsy showing Tubulointerstitial disease, selectivity index >(0.2 and tubular proteinuria (Banerjee, 2002; Bhimma, 2005; Ramjje, Coovadia, & Adhikari, 1997). Unfortunatley patients with SRNS are highly prone to develop complications in comparison to children having steroid sensitive nephrotic syndrome. These complication include acute renal failure (White, Glasgow, & Mills, 1970), chronic renal failure, growth retardation, impaired immunity leading to infections like peritonitis and thrombosis (Bhimma, 2005).

2. Review and Discussion

Children suffering from nephrotic syndrome have shown to have minimal change glomerulonephritis (MCNS), focal segmental glomerulosclerosis (FSGS) or mesangial proliferative glomerulonephritis; hence most are
suffering from Idiopathic Nephrotic Syndrome (Habib, 1993; Churg, Habib, & White, 1970).

SRNS and mainly FSGS have shown to have 50% risk of end stage renal disease within 5 years of diagnosis in case the patient does not manage to have partial or complete remission. Only 10-20% of nephrotic syndrome patients develop resistance to steroids but this fraction contributes disproportionately to end stage renal disease as compared to steroid sensitive nephrotic syndrome. If progressive renal impairment is controlled, complete or partial remission preserves renal function and brings about excellent long term results. However, End stage renal disease in patients with SRNS considerably reduces life expectancy, about 19 years after initiation of dialysis and around 40 years following transplantation (Kidney international Supplement, 2011; Gipson et al., 2006; Butani & Ramsamooy, 2009).

Among pediatric nephrologists there are two definitions of SRNS. The definition introduced by the International Study of Kidney Disease in Children (ISKDC) and used by the Arbeitsgemeinschaft für Pädiatrische Nephrologie (APN) is widely accepted and which states, ‘ No urinary remission within four weeks of prednisone therapy of 60 mg/m2/day’. The other definition, employed by the Society of French Speaking Pediatric Nephrologists says, ‘ No urinary remission following four weeks of prednisone at 60 mg/m2/day followed by three intravenous pulses of methylpredisolone ’.(Brodehl, Krohn, & Ehrich, 1982; Niudet, 1994).

When SRNS is suspected, a meticulous search for the possibility of concurrent infection (e.g skin infection and sinusitis), compliance problem, drug interaction, or inappropriate dosage is necessary. If these secondary conditions are ruled out, tissue diagnosis from a renal biopsy is the next step. Histological findings of SRNS are, and rarely, secondary glomerulopathy such as amyloidosis is unexpectedly found. At the same time, analysis for mutational genes known to cause SRNS is recommended.

Few factors have to be evaluated when treating for SRNS. These include confirmation of resistance to steroids( usually oral prednisone or oral prednisolone), kidney biopsy to rule out secondary reasons of Nephrotic Syndrome, determine GFR at presentation due to long term risk of kidney failure and quantification of proteinuria to check treatment response(Kidney International Supplement, 2011).

Treatment for steroid-resistant nephrotic syndrome (SRNS) is challenging and children who suffer from SRNS require aggressive treatment to achieve remission. Thus, when intravenous high-dose methylprednisolone does not work, calcineurin inhibitors, such as cyclosporine, is used as the first line of treatment (Tejani & Ingulli, 1995). Cyclosporine has shown to have higher rate of remission as compared to other immunosuppressant therapies used for the treatment of SRNS (Tahar & Rachid, 2010). Calcineurin inhibitors have been used more in an empirical manner than on the basis of clear rationale (Tejani & Ingulli, 1995). It was in 1984 when cyclosporine was first considered for the treatment of SRNS. Till now many studies have been conducted to determine dosages, duration of therapy, side effects and advantages of cyclosporine (Adhikari & Coovadia, 1994).

Cyclosporin is a calcineurin inhibitor that suppresses immune response by downregulating the transcription of multiple cytokine genes. The most significant of these cytokines is interleukin-2, which serves as the major activation factor for T cells in numerous immunologic processes. Cyclosporin inhibits cytokine production from T helper cells and also has an inhibitory effect on antigen presenting cells which are the main agents of T cell stimulation. A further effect of IL-2 inhibition is a reduction in B-cell activation and subsequent antibody production. IL-2 levels are known to become elevated during proteinuria and to normalize during remission in adults with idiopathic nephrotic syndrome and in children with Minimal change Nephrotic Syndrome (MCNS) or FSGS (Tejani & Ingulli, 1995). However, this pattern of interleukin-2 activity is felt to be part of a more widespread disorder of cellular immunity that results in nephrotic syndrome rather than being causal of proteinuria.It has been reported that cyclosporin has some antiproteinuric action on glomerular perm-selectivity to proteins that is unrelated to its immunosuppressive properties. Among these are an influence on perm-selectivity and charge selectivity and impairment of GFR. These data come from various human studies (Zietse et al., 1992; Meyrier, Noel, Auriche, & Callard, 1994; Ambalavanan et al., 1996; Heering et al., 2001) and animal models (Heering et al., 2001; Kokui et al., 1992, Schriiver et al., 1995; Desassis et al., 1997) with no immunologically mediated disease. Some studies revealed that lesions from the primary glomerular disease had either not regressed or had continued to progress (Meyrier, Noel, Auriche, & Callard, 1994; Ambalavanan et al., 1996; Chen, 2003).

Treatment of SRNS remains a difficult challenge in pediatric nephrology. At the moment there is no diagnostic marker for children displaying with nephrotic syndrome that can be used as a predictor of steroid resistance or responsiveness. The most important prognostic marker for children with nephrotic syndrome is their response to steroid treatment. However, Initial steroid treatment can be avoided only in patients with a family history of
SRNS or in those who have a known gene mutation. According to a study, the predictors for Cyclosporin non-responsiveness were steroid resistance, non minimal change disease on biopsy and longer duration between onset of nephrotic syndrome and cyclosporin usage, irrespective of the age of onset of the disease (Iyengar, 2006)

Using cyclosporine therapy in SRNS has its consequences. Long-term Cyclosporin therapy in low doses is effective in the treatment of children with idiopathic Nephrotic syndrome (NS), but the rate of relapse is high after drug withdrawal (El-husseini, 2005). Treatment with cyclosporine has significant nephrotoxicity. This is a common effect of all calcineurin inhibitors; hence cyclosporine therapy needs regular monitoring. This can be achieved by measurement of cyclosporine blood concentration at C0 which is pre-dose concentration or C2 levels which is concentration at 2 h post-dose. Better assessment can be done by C2 measurement as C0 represents the trough blood levels which do not significantly represent function of cyclosporine intake. Studies have shown that this drug can be safely used in children with nephrotic syndrome and also in transplant patients. Moreover it has also been proved by yearly assessments of creatinine and creatine clearance that renal function remains stable for up to 20 years (Serkova & Christians, 2003; Midvedt et al., 2003; Fujinaga, 2006; Kandaswamy, 2007).

According to recent researches once a day administration of the drug is more beneficial than the old twice a day administration due to former providing an absorption profile with peak blood concentration of cyclosporine which may cause remission of SRNS and prevent chronic cyclosporine nephrotoxicity (Tanaka et al., 2004; Takeda, 2007).

According to a study, renal insufficiency developed in 6% and hypertension in 10% of patients (most patients with FSGS), (El-husseini, 2005) Furthermore, progression of the previous interstitial fibrosis and tubular atrophy was noted in two patients, suggesting a 17% incidence of cyclosporin nephrotoxicity. This analysis of the long-term risks of cyclosporin for childhood NS has identified two important findings: (Tejani & Ingulli, 1995) combined cyclosporin and alternate-day steroids can be highly effective in inducing complete remission in patients with SRNS and biopsy-proven IgM nephropathy, and (Tahar & Rachid, 2010) long-term use of cyclosporin in moderate doses with closely monitored levels can result in a relatively low incidence of nephrotoxicity. (Gregory et al., 1996) There was a higher incidence of cyclosporin dependence among young responders. Also, patients with cyclosporin resistance are at high risk for significant infections and CRF (Iyengar, 2006).

A number of trials have been done for cyclosporine as a treatment of SRNS. In Three Randomized Control Trials (RCTs) with 49 patients, 26 patients were given cyclosporine while 23 were given control therapy or placebo. It was concluded that complete remission resulted in 31% and partial remission in 38% was seen during a 6 month period of therapy. Similarly another trial which was conducted in 2011 in which cyclosporine was compared to mycophenolate added to high dose of dexamethasone for a 12 month duration resulted in around 19% complete remission and around 26% partial remission (Garin et al., 1988; Lieberman & Tejani, 1996; Ponticelli et al., 1993; Gipson et al., 2011).

In the recent years, experts of the Indian Society of Pediatric Nephrology were involved in a 2-stage process to formulate guidelines, based on recent practices and available evidence, on management of SRNS children. Agreement of at least 80% participants formed an opinion (Gulati et al., 2009). The Expert Group emphasized that while all patients with SRNS should initially be referred to a pediatric nephrologist for evaluation, the subsequent care should be collaborative involving the primary pediatrician and the nephrologist. Following the diagnosis of SRNS (lack of remission despite treatment with prednisolone at 2 mg/kg/day for 4 weeks), all patients (with initial or late resistance) should undergo a kidney biopsy, before instituting specific and targeted therapy. Patients with idiopathic SRNS secondary to MCNS or FSGS should receive similar treatment. Effective therapy regimens include treatment with calcineurin inhibitors (Cyclosporine), combination of pulse corticosteroids with oral cyclophosphamide or intra-venous cyclophosphamide, tapering doses of alternate day corticosteroids. Supportive management comprises therapy with angiotensin converting enzyme inhibitors and statins. These guidelines are expected to enable standardization of care for patients with SRNS worldwide (Gulati et al., 2009).

KDIGO Clinical Practice Guideline for Glomerulonephritis published in 2012, suggested treatment recommendations for SRNS. These were calcineurin inhibitor (CNI) as initial therapy which should be continued for at least 6 months and discontinued only if partial or complete remission of proteinuria is not observed; In case partial remission is achieved by 6 months then CNIs can be continued for minimum of 12 months. Other suggestions in these guidelines were low dose corticosteroid therapy with CNI therapy and ACE-I or ARBs were also recommended. In children who fail to respond to CNI therapy can be treated with mycophenolate mofetil, high-dose corticosteroids, or a combination of these agents in children who fail to achieve complete or partial
remission with CNIs and corticosteroids. It was suggested to not to use cyclophosphamide in these children. These guidelines also mentioned treatment for patients with a relapse of nephrotic syndrome after complete remission, in which therapy should be restarted using oral corticosteroids, returning to previous successful immunosuppressive agent or an alternative immunosuppressive agent to minimize potential cumulative toxicity (Clinical Practice Guidelines KDIGO, 2012).

3. Conclusion

SRNS in children is a difficult disease with significant morbidity and mortality. However, remission is achievable with cyclosporine and other immunosuppressive agents. Although the prognosis of SRNS is complicated, an intensive treatment in the early stages of the disease may achieve remission in more than half of the patients. Thus, timely referral of pediatric SRNS patients to pediatric nephrology specialists for histological and genetic diagnosis and treatment is highly recommended. Treatment should be individualized according to the underlying histopathology, and clinical and environmental conditions of the children. (Kazi & Halawani, 2010) There is an urgent need to distinguish as soon as possible those patients who may benefit from prolonged immunosuppressive treatment from those who will not benefit from such treatment and who will just suffer from its major side effects. The emerging evidence that the majority of genetic forms of SRNS should receive symptomatic treatment only, should also be clinically tested and studies baring its significance should be evaluated in the future.

Competing Interest Section

The authors declare that they have no competing interests

Acknowledgements

None Declared.

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Gipson, D. S., Chin, H., Presler, T. P., Jennette, C., Ferris, M. E., Massengill, S., Gibson, K., & Thomas, D. B.


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