Plasma exchange or immunoadsorption for recurrent focal segmental glomerulosclerosis: clear differences in vitro

Susan T. Veissi¹, Bart Smeets², Joanna A.E. van Wijk³, Thea van der Velden¹, Lambertus P. W. J. van den Heuvel¹, 4, 5*, and Michiel F. Schreuder², 6*

¹Department of Pediatric Nephrology, Amalia Children’s Hospital, Radboud university medical center, Radboud Institute for Molecular Life Sciences, Nijmegen, The Netherlands, ²Department of Pathology, Radboud University Medical Center, Radboud Institute for Molecular Life Sciences, Nijmegen, The Netherlands, ³Department of Pediatric Nephrology, Amsterdam University Medical Center, Amsterdam, The Netherlands, ⁴Department of Laboratory Medicine, Radboud University Medical Center, Radboud Institute for Molecular Life Sciences Nijmegen, The Netherlands and ⁵Department of Development and Regeneration, University Hospital Leuven, Leuven, Belgium

*These authors contributed equally to this work.

Correspondence to: Michiel F. Schreuder; E-mail: Michiel.Schreuder@radboudumc.nl

Therapy-resistant nephrotic syndrome in children is generally based on primary focal segmental glomerulosclerosis (pFSGS) and will lead to deterioration of kidney function. Kidney transplantation in such patients may result in a recurrence of FSGS (rFSGS), which has been reported in up to 50% of children [1]. Plasma circulating permeability factors (CPFs), the nature and origin of which still remain elusive, have been designated a pathological role in both pFSGS and rFSGS. Clinical evidence including posttransplant recurrence of the disease and response to extracorporeal therapies supports the presence of CPFs in patients. Moreover, in vitro experimental data with sera from rFSGS patients that induce glomerular permeability and podocyte injury highlights the pathophysiological role of CPFs in FSGS [2]. Furthermore, podocytopenia following podocyte injury is linked to elevated reactive oxygen species (ROS) levels that result in podocyte cell detachment and/or death [3]. Interestingly, higher levels of oxidative markers have been measured in children with active nephrotic syndrome [4].

FSGS patients with a putative CPF often respond well to immunoadsorption (IA) or therapeutic plasma exchange (TPE). More importantly, preemptive IA or TPE has been shown to reduce the risk of rFSGS after transplantation [5]. Although TPE has a success rate of approximately 70% in pediatric patients, a semi-selective IA therapy removing immunoglobulins (Igs) but preserving other plasma proteins is increasingly used [2, 6]. The effectiveness of IA may suggest that the CPF could be a circulating Ig or lie in the Ig fraction, but this concept is not yet supported by in vitro experimental data. Here, we aimed to purify Ig fractions from the plasma of FSGS patients with a presumed CPF and investigate the effect of these fractions on human conditionally immortalized podocytes (hciPods) in vitro.

The study was conducted according to the recommendations of the appropriate version of the Helsinki Declaration and all subjects had given informed consent. TPE samples were collected from three FSGS patient with presumed CPF at active disease (Act 1, Act 2, Act 3). From one of these patients the remission plasma was also available (Rem) and included as control, as well as two non-kidney disease TPE patients (NR-Ctrl 1 and 2, both with neuromyelitis optica). Heparin plasma from seven healthy individuals was collected and pooled, and used as a healthy control plasma (hCtrl pool). Ig fractions were extracted from these seven samples using a protein A/G affinity chromatography column (Sigma). In addition to the column-bound Ig fractions (elution), the flow-through fractions of each sample were also collected. For in vitro assays, Ig fractions were concentrated to original plasma volume and the flow-through samples were corrected for the dilution factor. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) was performed to confirm that plasma Igs were isolated. Ig concentrations in the TPE and through samples were assessed fluorometrically with a CM-H2DCFDA probe.

SDS-PAGE performed on the Ig fractions and flow-throughs showed two bands of 25 and 50 kDa only in the Ig fractions (Fig. 1A). The observed 25 and 50 kDa bands indicate the light chain and the glycosylated heavy chain of the IgG class of antibodies, respectively. Elevated ROS levels in hciPods in response to Ig fractions and flow-throughs was assessed fluorometrically with a CM-H2DCFDA probe.

© The Author(s) 2022. Published by Oxford University Press on behalf of the ERA. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial License (https://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com
lead to oxidative stress in cultured podocytes. Furthermore, approximately 2% Iggs was measured with nephelometry in the flow-through fractions obtained from three FSGS patients with presumed CPF at active disease when compared with the Ig fractions. Together, these data show that the putative CPFs in FSGS do not lie in Ig fractions, at least not in the three patients included in the current study.

Management of rFSGS in clinical practice remains a therapeutic challenge. Hence, currently there are no clear guidelines for the prevention of rFSGS after transplantation; treatment is mainly based on TPE or IA in combination with high dose of CD20-depleting antibody rituximab [7]. As no study has compared IA with TPE and case series have indicated a comparable efficacy, IA is increasingly used in various clinics to treat rFSGS. However, our results show that the putative CPF associated with FSGS does not seem to lie in Ig fractions of the three patients studied here. This suggests that IA solely may not be enough to efficiently remove CPFs. In a patient resistant to IA, it may be of benefit to use TPE to achieve remission of rFSGS. For the future, it may be beneficial to design randomized and prospective clinical trials to compare IA with TPE for the management of rFSGS.

ACKNOWLEDGEMENTS

We thank Prof. Moin Saleem for the conditionally immortalized human podocyte cell line.

FUNDING

This study was performed on behalf of the LEARNs (Levamisole Adjuvant therapy to Reduce relapses of Nephrotic Syndrome) Consortium, an interuniversity collaboration in the Netherlands, funded by the Dutch Kidney Foundation (CP 16.03).

CONFLICT OF INTEREST STATEMENT

All the authors declared no competing financial interests.

REFERENCES

1. Trachtman R, Sran SS, Trachtman H. Recurrent focal segmental glomerulosclerosis after kidney transplantation. Pediatr Nephrol 2015;30:1793–802. https://doi.org/10.1007/s00467-015-3062-1
2. Ha T-S. Circulating permeability factors in idiopathic nephrotic syndrome. Child Kidney Dis 2019;23:7–21.
3. Mundel P, Shankland SJ. Podocyte biology and response to injury. J Am Soc Nephrol 2002;13:3005–15. https://doi.org/10.1097/01.asn.0000039661.06947.fd
4. Reddy P, Sindgikar S, Shenoy R et al. Oxidative stress in childhood steroid sensitive nephrotic syndrome and its correlation with DNA damage. Int J Contemp Pediatrics 2016;3:768–72. http://dx.doi.org/10.18203/2349-3291.ijcp20161853
5. Gohh RY, Yango AF, Morrissey PE et al. Preemptive plasmapheresis and recurrence of FSGS in high-risk renal transplant recipients. Am J Transplant 2005;5:2907–12. https://doi.org/10.1111/j.1600-6143.2005.01112.x
6. Ponticelli C. Recurrence of focal segmental glomerular sclerosis (FSGS) after renal transplantation. Nephrol Dial Transplant 2010;25:25–31. https://doi.org/10.1093/ndt/gfp538
7. Weber LT, Tönshoff B, Grenda R et al. Clinical practice recommendations for recurrence of focal and segmental glomerulosclerosis/steroid-resistant nephrotic syndrome. Pediatr Transplant 2021;25:e13955. https://doi.org/10.1111/petr.13955

Received: 31.5.2022; Editorial decision: 31.8.2022