

NZ-ISNCON-2026
30th Annual Conference of the
Indian Society of Nephrology Northern Zone
10th-12th April 2026, Hyatt Centric, Dehradun

Programme for NZ-ISNCON 2026

10th April 2026

Time	Topic	SPEAKERS	CHAIRPERSONS
1 (04:00-04:20 PM)	Rituximab in MN	Dr Avichal Rajpal, PGIMER, Chandigarh	Prof Dipankar Bhowmik Dr Dilip Bhalla Prof Lalit K Pursnani Dr Shobha Sharma Prof S C Tiwari Dr Vijay Kumar Binwal
2 (04:20-04:40 PM)	Thymoglobulin in Kidney Transplant	Dr Vinant Bhargava SGRH, New Delhi	
3 (04:40-05:00 PM)	Multiplex PCR in post-kidney transplant Diarrhoea	Dr Anurag Gupta, SGRH, New Delhi	
4 (05:00 – 05:20 PM)	START trial	Dr Arun Prabhakar PGIMER, Chandigarh	
5 (05:20– 05:40 PM)	Crescentic IgA nephropathy	Dr Pallavi Prasad VMMC, New Delhi	
6 (05:40– 06:00 PM)	Phenotype and Genotype correlation in Indian children with Renal Tubular Acidosis: unravelling the mystery of inheritance.	Dr Suprita Kalra (Base Hospital, Delhi Cant.)	
7 (06:00– 06:20 PM)	AVF angioplasty	Dr Manas Patel, SGPGIMS, Lucknow	
8 (06:20 – 06:40 PM)	Hepatitis B-Post kidney Transplant	Dr Manas Behra, SGPGIMS, Lucknow	
9 (06:40– 07:00 PM)	Management of central venous stenosis	Dr Santosh Kumar, SGPGIMS, Lucknow	

11 April 2026

*“The names in the program have been arranged in alphabetical order.”

Time	Topic	Objective	SPEAKERS	CHAIRPERSONS
8:00-8:15 AM	Welcome address		Dr Puneet Arora Dr Raja Ramachandran	
8:15-9:00 AM	Slippery slopes of Nephrology	To discuss the low-value care in Nephrology and steps to address the same	Dr Ajit Singh Narula Prof Anil Bhalla Prof Dinesh Khuller Prof Harbir Singh Kohli Dr Urmila Anandh Prof Vijay Kher Prof Vinay Sakhuja Prof Vivekanand Jha	
9:00-09:45 AM	Case 1: A 29yr old male with end stage kidney disease of unknown aetiology, on thrice weekly maintenance haemodialysis, underwent an ABOi kidney transplant with his mother (51yr old female) as donor. There were 5/12 mismatches in HLA and both CDC crossmatch and flow cytometry crossmatch were negative before transplant. There were also low-titre donor-specific antibodies detected against DPA1*01:03:DPB1*04:02 (MFI–1070). The baseline anti-B titre (IgG isohemagglutinin) was 1:16, and he underwent pretransplant desensitization with rituximab (200mg) given 2 weeks before transplant. Tacrolimus and MMF were started 1 week before transplant. He received ATG induction (75mg on POD0 and 75mg on POD1) and triple immunosuppression (steroid, MMF, tacrolimus). In the immediate post-operative period he had good urine output with Creatinine reducing to 3.2mg/dl on POD 2. Anti B titer repeated on POD 1 had	Case 1: • How would you manage this patient?	Dr Ajay Kher Dr Jyoti Agarwal Dr Kajaree Giri Dr Lekha Rani Prof Muzafar Wani Dr Namrata Rao Dr Tarun Mittal Dr Uma Kanga Resident Representative (RML)	

reduced to 1:1. However, on the evening of POD 2, the patient developed oliguria and had to be started on diuretics. Labs showed evidence of severe thrombotic microangiopathy (severe anemia, thrombocytopenia and high LDH) with graft dysfunction. Complement factor C3 and C4 were normal and direct comb's test was negative. Tacrolimus was withheld due to suspicion of CNI induced TMA. Doppler of the graft kidney was normal. Serum creatinine continued to rise, peaking at 4.83 mg/dL. He was given 1 session plasmapheresis on POD 3 with full FFP replacement. Over the course of the next few days, he received multiple blood products (10 RDP, 1 SDP, 5 LDPRBC) due to ongoing haemolysis and thrombocytopenia. Despite plasmapheresis there was no clinical improvement. Given the unavailability of eculizumab at that time and the patient's worsening condition, graft nephrectomy was contemplated.

Case 2:
40 y/o Female with ESRD and on dialysis for 1 month with unknown etiology, with her husband as donor. Rest of her workup is normal. After you rule out her husband as donor- due to crossmatch positivity, one of her 2 sons come forth as potential donor though Ankita does not want them to donate. Immunological workup is below.

Sensitization History	2 pregnancies					
HLA typing	A	B	C	DRB1	DQB1	DQA1
Donor (Husband)	02:11, 24:07	08:01, 52:01		03:01, 10:01	02:01, 05:01	01:01, 01:01/04
Elder Son	11:01, 24:07	35:03, 52:01		10:01, 13:01	05:01, 06:03	Not done.
Recipient	11:01, 24:02	35:03, 51:01		04:04, 13:01	03:02, 06:03	01:03, 03:01
XM	T cells	B Cells				
CDC XM (husband)	Positive	Positive				
Flow XM (Husband)	Positive	Positive				
Single Antigen Bead (Ankita Report 1)	Class I : Multiple anti HLA Abs	Class II: Multiple Anti HLA Abs.				

- Case 2:**
- Should we even consider Son as husband has a positive crossmatch?
 - Which of the antibodies are Donor Specific against husband?
 - Role and Importance of DQ and DP antibodies.?
 - How would you evaluate these donors and decide proceeding for transplant? How do you evaluate multiple donors from family?

9:45-10:30 AM

Management of anaemia

- The role of HIF-PHI in the management of anaemia in CKD.
- Role of HIF-PHI in special circumstances like PRCA
- Indian experience in management

Dr Ajay Goyal
Dr Chandani Bhagat
Dr Neeru Aggarwal
Dr Piyush Mathur
Dr PP Varma
Dr Sanjiv Saxena
Dr Vikram Singh

		<ul style="list-style-type: none"> of anaemia using HIF-PHI Strategies to address ESA resistance in CKD 		
10:30-10:45 AM	Tea Break	<ul style="list-style-type: none"> No specific objective 		
10:45-11:30 AM	<p>A 28-year-old man was found to have mildly elevated blood pressure during a routine health check, with a reading of 138/90 mmHg. He had no symptoms and felt well at the time. Initial urine testing showed abnormalities, with 2+ red blood cells and 2+ protein. Further evaluation showed normal kidney function, with a serum creatinine of 0.8 mg/dL. A 24-hour urine test revealed protein excretion of 1.2 g/day, indicating sub-nephrotic range proteinuria. Because of persistent protein in the urine, microscopic blood in the urine, and borderline high blood pressure in a young person with normal kidney function, he was referred to a nephrologist. After clinical evaluation and review of investigations, a kidney biopsy was advised to identify the cause. The kidney biopsy showed mesangial expansion with endocapillary hypercellularity. According to the Oxford classification, the biopsy was graded as M1 E1 S0 T0 C0, confirming a diagnosis of IgA nephropathy. These findings indicated active inflammation in the kidneys without evidence of chronic or irreversible damage. The patient was counselled about IgA nephropathy, its variable course, and available treatment options. He was started on supportive treatment, including renin–angiotensin system blockade with telmisartan 40 mg twice daily, to control blood pressure and reduce proteinuria. He was also advised on lifestyle measures such as salt restriction and regular blood pressure monitoring. As proteinuria persisted despite supportive care and the biopsy showed active disease (M1E1) with preserved kidney function, treatment with controlled-release budesonide was initiated. He was advised to continue treatment for 9 months. The patient was informed in detail about the reason for treatment (numerous option- anti-proteinuric plus immunosuppression), expected benefits, and possible side effects. Close follow-up was planned, and he was asked to return in 2–3 weeks for reassessment of blood pressure, kidney function, proteinuria, and treatment tolerance.</p>	<ul style="list-style-type: none"> The role of CR-Budesonide in management of IgA nephropathy Evidence for the role of budesonide in IgA nephropathy Discussion on the Indian data on CR-Budesonide in IgA nephropathy 	<p>Moderator: Prof Amit Gupta Panelist: Dr Anuja porwal Dr Alok Kumar Dr Reetesh Sharma Dr RK Yadav Prof Sanjeev Gulati, Dr Vijay Kumar Sinha</p>	
11:30-12:00 PM	Management of Vasculitis-ANCA	By experts in the field	Prof Aman Sharma	Dr Alok Kumar Pandey Dr Jayant Hota Prof Ritambhra Nada Dr SNS Yadav
12:00-12:30 PM	Management of Multiple Myeloma what nephrologist should know	By experts in the field	Prof Pankaj Malhotra	Prof Dhananjai Aggarwal Dr DK Sinha Prof Ritambhra Nada Dr Sandeep Mahajan
12:30-1:30 PM	Climate change is real—but our (global and not India) approach is hypocritical	Debate	Dr (Maj. Gen.) Ranjith Nair and Dr Urmila Anandh	
1:30-2:15 PM (Lunch Symposium)	Newer Potassium Binders in Hyperkalemia management- Are they worth the debate in Indian Scenario	<ul style="list-style-type: none"> Role of sodium zirconium cyclosilicate in management of acute hyperkalemia Dosing protocol for its use for treatment of hyperkalemia 	Dr Ajit Singh Narula Dr Deepak Diwan Dr Raka Kaushal Dr Salil Jain Dr Sanjiv Jasuja Prof Vijay Kher	

		<ul style="list-style-type: none"> • Role as a maintenance agent for hyperkalemia 		
2:15-2:30 PM	Break			
2:30-3:15 PM	A challenging case of complement-mediated TMA	<ul style="list-style-type: none"> • To discuss the role of eculizumab in CM-TMA • To discuss role of PLEX and eculizumab in anti-CFH-TMA • Interpretation of genetic testing in patients with CM-TMA • Functional testing for complement activity • Procurement of eculizumab using the rare-disease programme 	Moderator: Prof Aditi Sinha Dr Arun Prabhakar Dr Gaurav Sharma Dr Rajesh Jhorwat Dr Sharon Kandari Dr Shubham Shukla Dr Srinivasa Vardhan	Dr M.L Patel
3:15-4:00 PM	Two lectures by ICMR scientist on how to write proposals		Dr Harpreet Singh Dr Sudipta Roy	
4:00-4:15 PM	Tea Break			
4:15-5:15 PM	<p>A 73-year-old gentleman, a former small-business owner, presented with progressive lethargy, reduced exercise tolerance, anorexia, and generalized itching for nearly one year. His background medical history is notable for Type 2 diabetes mellitus for over 25 years, complicated by advanced diabetic retinopathy with significant visual impairment and diabetic autonomic neuropathy. Over the last 6–9 months, he has developed recurrent postural dizziness, early morning falls, and episodic presyncope, particularly on dialysis days. He is a known case of ischemic heart disease, having sustained an inferior wall myocardial infarction 6 years ago, followed by coronary artery bypass grafting. He remains on antiplatelet therapy and beta-blockers, with limited tolerance to aggressive volume shifts. In addition, he has a history of recurrent urinary tract infections, including one episode of emphysematous cystitis 18 months ago and another episode of urosepsis requiring ICU admission in the past year. He was diagnosed with secondary membranous nephropathy nearly 7 years ago in the setting of long-standing diabetes and chronic infections. Given his age, comorbidity burden, and repeated infective episodes, immunosuppressive therapy was deferred, and he was managed conservatively. Despite optimal supportive care, he progressed to end-stage kidney disease.</p> <p>Dialysis and Vascular Access History The patient was initiated on maintenance haemodialysis 11 months ago through a right internal jugular tunnelled dialysis catheter due to poor superficial veins and delayed referral. A left brachio-cephalic arteriovenous fistula was subsequently created. Although the fistula matured clinically, it was complicated by high venous pressures and suboptimal dialysis adequacy. Doppler evaluation revealed outflow stenosis, for which he underwent percutaneous transluminal angioplasty on two occasions. Currently, the fistula supports a blood flow of approximately 230–260 mL/min, with intermittent access alarms during dialysis sessions.</p> <p>Infectious and Nutritional Concerns The dialysis course has been complicated by one episode of catheter-related bloodstream infection</p>	<p>The patient now presents with worsening pruritus, generalized weakness, bone pain, and declining functional independence. Family members express concern regarding his quality of life and frequent hospital visits. The treating team is faced with multiple management challenges:</p> <ul style="list-style-type: none"> • What is the dialysis optimising strategies in this case • Improving dialysis adequacy with HDF in the setting of limited AVF flow and intradialytic hypotension • What are the factors predicting a better outcome with HDF • Indian data on the quality of life with HDF • How do you manage skin discoloration in a patient with ESKD • How do you manage intractable pruritus in a 	Dr Aakanksha Sharma Prof Dinesh Khuller Dr Munish Chauhan Prof PP Varma Dr Ritika Bansal Dr Saurabh Nayak Dr Simran Kaur	

	<p>and repeated lower urinary tract infections with prior cultures growing extended-spectrum beta-lactamase-producing organisms. He has experienced progressive weight loss and poor nutritional intake, with intermittent low-grade inflammatory markers, raising concern for chronic inflammation and protein-energy wasting.</p> <p>Current Clinical Assessment On examination, the patient appears elderly and frail, with reduced muscle bulk and slow gait. His skin shows diffuse hyperpigmentation with a greyish-brown hue, especially over the face, neck, and extensor surfaces. There is marked xerosis, excoriations, and pruritic nodules consistent with chronic uremic pruritus. He also reports altered taste sensation and restless sleep. Blood pressure measurements demonstrate significant orthostatic hypotension, with systolic blood pressure falling from 148 mmHg supine to 112 mmHg on standing. Dialysis sessions are frequently interrupted due to intradialytic hypotension, limiting ultrafiltration and resulting in persistent volume and uremic symptoms.</p> <p>Laboratory Investigations Recent laboratory parameters are as follows:</p> <ul style="list-style-type: none"> • Haemoglobin: 9.1 g/dL, on erythropoiesis-stimulating agents • Blood urea nitrogen: 164 mg/dL • Serum creatinine: 9.8 mg/dL • Corrected serum calcium: 8.6 mg/dL • Serum phosphorus: 6.4 mg/dL • Intact parathyroid hormone (iPTH): 842 pg/mL • Serum albumin: 3.1 g/dL • Ferritin: elevated with borderline transferrin saturation <p>These findings indicate suboptimal dialysis adequacy, advanced CKD-mineral and bone disorder, anemia of chronic kidney disease with inflammation, and protein-energy wasting.</p>	<p>patient with ESKD</p> <ul style="list-style-type: none"> • Managing severe secondary hyperparathyroidism in an elderly patient with ischemic heart disease, what are the targets and indications for parathyroid surgery • Addressing recurrent infections and minimizing further vascular access-related complications • Optimizing anaemia and nutritional status amid chronic inflammation <p>10 minutes: Role of central dialysis fluid delivery system in a haemodialysis facility</p>		
5:15-5:45 PM	Esaxerenone- an antihypertensive and anti proteinuric therapy		Prof Amit Gupta Prof P P Varma Dr Raka Kaushal Dr Sanjiv Saxena Prof Vijay Kher	
06:00 PM	GBM			
07:30 PM	Inauguration and Dinner			

12th April 2026

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Time	Topic		Speakers	Chairpersons
8:00-8:30 AM	A challenging case- Peritoneal Dialysis	Discussion on clinical challenges (not logistical or financial) while managing patients on peritoneal dialysis- 4 to 5 clinical scenarios	Prof Anil Bhalla Prof Himanshu Mahapatra Prof (Brig) Pavitra Manu Dogra Dr Vikas Makkar	
8:30-9:15 AM	<p>Case 1 A 32-year-old male with biopsy-proven IgA nephropathy progressed to end-stage kidney disease (ESKD) and is planned for a living donor kidney transplantation, with his mother as the donor. His sensitizing history includes transfusion of four units of packed red blood cells (PRBCs) approximately eight months prior, at the time of initial diagnosis and management of his primary disease. His current immunological workup reveals:</p> <ul style="list-style-type: none"> • ABO compatible • Negative complement-dependent cytotoxicity (CDC) crossmatch • Negative flow cytometry crossmatch • Single antigen bead (SAB) assay negative for donor-specific antibodies (DSA) against HLA-A, HLA-B, HLA-DR, and HLA-DQ 	<ul style="list-style-type: none"> • How would you risk-stratify patient immunologically? Would you consider this low-, intermediate-, or high-risk? • What would be your approach to induction immunosuppressive therapy in the two clinical scenarios 	Dr Deepak Pathania Dr Manish Jain Dr Manish Singla Dr Manoj Singhal Dr Suman Lata Prof Vinay Malhotra Prof Vikas Makkar	

	<ul style="list-style-type: none"> Panel reactive antibody (PRA) low/absent <p>He has no prior transplant history and no other known sensitizing events. His comorbidities include well-controlled hypertension, with no diabetes or cardiovascular disease.</p> <p>Case 2 A 33-year-old woman with ESKD secondary to analgesic nephropathy has been on maintenance haemodialysis for three years. She is scheduled to undergo a living donor kidney transplantation, with her husband as the donor (spousal transplant). Her sensitizing history includes:</p> <ul style="list-style-type: none"> Two prior pregnancies One spontaneous abortion Two PRBC transfusions <p>Immunological evaluation shows:</p> <ul style="list-style-type: none"> ABO compatible Negative CDC crossmatch Flow crossmatch pending/weakly positive (if relevant to discuss) SAB assay demonstrating anti-HLA antibodies, including: <ul style="list-style-type: none"> HLA-A*02:02 with MFI 2400 HLA-B*03:01 with MFI 2100 <p>Calculated PRA is moderately elevated. No prior transplant history. She has secondary hyperparathyroidism and mild left ventricular hypertrophy but is otherwise clinically stable.</p>	<ul style="list-style-type: none"> What is the Indian experience with use of Induction agents: Thymoglobulin Evidence in favour of low-dose ATG in kidney transplant Role of CD3 monitoring in patients receiving thymoglobulin 		
9:15-10:00 AM	Debate (AI will take away our prescription job)		Dr Arun Kumar Dr Gagandeep Chhabra	Dr Sanjiv Saxena Prof Amit Gupta Dr Santosh Kumar
10:00-10:15 AM	Tea Break			
10:15-10:45 AM	Prof KK Malhotra Oration: Prof Narayan Prasad, SGPGIMS, Lucknow		Prof Narayan Prasad	Prof Ashwini Gupta (President ISN) Prof Sanjay D Cruz (President NZ-ISN)
10:45-11:15 AM	Presidential oration		Prof Sanjay D Cruz	Prof Dipankar Bhowmik Dr K N Singh
11:15-11:30 AM	ATG-Grafalon in Kidney Transplant		Dr Manish Malik	-
11:30-12:10-PM	<p>Case 1 A 54-year-old gentleman with type 2 diabetes mellitus, long-standing hypertension, and heart failure with reduced ejection fraction (HFrEF, EF 30%) presents with worsening shortness of breath, orthopnea, and bilateral pedal edema. On examination, he has elevated jugular venous pressure, bilateral basal crackles, and pitting edema up to the knees. Over the past 72 hours, his urine output has declined. His serum creatinine has increased from a baseline of 1.1 mg/dL to 2.3 mg/dL. Serum potassium is 4.9 mEq/L, sodium 132 mEq/L, and NT-proBNP is elevated. He is currently on:</p> <ul style="list-style-type: none"> Torsemide 10 mg once daily Sacubitril/valsartan Metoprolol succinate Spironolactone 25 mg daily Empagliflozin <p>Chest X-ray shows pulmonary congestion. Blood pressure is 104/68 mmHg.</p>	<p>Case 1 How would you escalate diuretic therapy in this setting?</p> <ul style="list-style-type: none"> Oral dose escalation vs. intravenous loop diuretics Bolus vs. continuous infusion Dose conversion strategy <p>How would you assess for diuretic resistance? When would you introduce sequential nephron blockade (e.g., thiazide-type diuretic such as metolazone)? What parameters will you monitor to assess response?</p> <ul style="list-style-type: none"> Urine output (target per hour/day) 	Prof Anupama Kaul (M) Prof Himanshu Varma Prof Muthu Kumar Dr Nishant Nadda Dr Sahil Garg Prof Shivendra Singh	Dr Ankur Chaudhary Prof. K L Gupta Dr Mohit Khirbat Dr Sourya Sourabh Mohakuda

	<p>Case 2 A 21-year-old woman is admitted with generalized swelling and biopsy-proven minimal change disease (MCD). She presented with nephrotic syndrome characterized by:</p> <ul style="list-style-type: none"> • Proteinuria 8 g/day • Serum albumin 1.6 g/dL • Total cholesterol 320 mg/dL • Serum creatinine 0.9 mg/dL at admission <p>Two days after the renal biopsy, she develops worsening edema and reduction in urine output to 400 mL/day. She remains normotensive. There is no gross hematuria.</p> <p>Color Doppler ultrasound of the renal vessels is normal, with no evidence of renal vein thrombosis or arterial compromise.</p> <p>Current medications include:</p> <ul style="list-style-type: none"> • Prednisolone (initiated after biopsy) • Pantoprazole • No diuretics yet <p>She has tense ascites, periorbital edema, bilateral pleural effusions on ultrasound, and difficulty ambulating due to anasarca.</p>	<ul style="list-style-type: none"> • Daily weight • Net fluid balance • Spot urine sodium • Renal function and electrolytes <p>How would you decide whether to increase or decrease the diuretic dose? At what point would you consider ultrafiltration? How would you differentiate cardiorenal syndrome from over-diuresis?</p> <p>Case 2</p> <ul style="list-style-type: none"> • What are the possible causes of acute oliguria in this setting? • How will you assess intravascular volume status in nephrotic syndrome? • How will you assess intravascular volume status in nephrotic syndrome? • How will you assess intravascular volume status in nephrotic syndrome? • What complications should you watch for during aggressive diuresis? • Would you initiate anticoagulation prophylaxis in this patient? If yes, what are your criteria? • How will you monitor response to therapy? 		
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12:10-12:40 PM	Vaccine in kidney diseases	To discuss the vaccine schedule in patient with kidney disease with special focus on Herpes Zoster vaccination and PCV20	Prof RK Sharma Dr Raghuvendra Singh Dr Rajiv Kumar Bhatia Dr Manju Aggarwal Dr Vishal Saxena Dr Vivek Ruhela	
12:45 PM	Valedictory function, Lunch and Departure			