• 1528 $\frac{\beta_2}{MiCROGLOBULIN}$ $(\beta_2\mu)$ IN RENAL TRANSPLANT RECIPIENTS. Manoj Narayanan, Lifida J. Peterson and Alan M.Robson. Dept. of Peds, Washington Univ. School of Medicine and St. Louis Children's Hospital, St. Louis, MO. The handling of $\beta_2\mu$ was studied in 19 renal transplant reciptents, aged 2 to 18 yr, to find an improved method to identify rejection and separate it from other causes of renal failure. Serum B u concentrations measured daily usually increased 24 to 48 b

The handling of $\beta_2 \mu$ was studied in 19 renal transplant recipients, aged 2 to 18 yr, to find an improved method to identify rejection and separate it from other causes of renal failure. Serum $\beta_2 \mu$ concentrations, measured daily, usually increased 24 to 48 h before serum creatinine (S₂) increased in patients undergoing transplant rejection. In addition, being poorly dialyzable, serum $\beta_2 \mu$ was superior to S₂ in demonstrating changes in GFR in transplant recipients requiring dialysis post transplant.

 $\beta_2\mu$ is reabsorbed almost completely from tubular fluid in the normal proximal tubule making urinary excretion of $\beta_2\mu$ a good indicator of proximal tubule function. Fractional excretion (FE) of $\beta_2\mu$ was followed daily in transplant recipients. Most values were elevated, especially in the first 4 days after transplantation, a time when acute tubular necrosis (ATN) is expected. Values also rose in several patients 3 to 7 days after rejection had been established. This is consistent with the thesis that acute rejection may be complicated by ATN which may persist after rejection is reversed by steroid pulses. The study demonstrates that daily measurement of serum $\beta_2\mu$ is a valuable adjunct in the management of renal transplant recipients, also that measurement of FE of $\beta_2\mu$ has the potential of distinguishing rejection from ATN in transplant recipients, helping to determine the need for treatment with steroid pulses. Measurement of FE is superior to measurement of urinary concentration of $\beta_2\mu$ for this purpose.

MINIMAL CHANGE NEPHROTIC SYNDROME (MCNS): AN AUTO-IMMUNE DISEASE? Terry Phillips, Leticia U Tina, Pedro A Jose, Zoe L Papadopoulou & Philip L Calcagno. Depts of Peds & Path. Georgetown Univ. Med. Ctr., Wash. D.C. The presence of low-level tissue directed antibodies (AAb's)

The presence of low-level tissue directed antibodies (Abb's) and immune complexes (IC) were determined in 12 MCNS and 5 mesangial proliferative glomerulonephritis (MesPGN) patients with biopsy proven disease. The AAb's were detected by immunofluorescence (IF) against human cadaveric tissues and the IC detected by 3 different methods and isolated by polyethylene glycol precipitation/sedimentation. The isolated antibodies from the IC were also tested by IF for AAb activity against the same tissue antigen battery. The presence of such AAb activity was demonstrated in 8/12 MCNS and in all of the MesPGN sera. In the MCNS group 7/8 of the AAb positive sera also contained IC with corresponding AAb activity (2/7 directed against kidney basement meuscle antigens but no other tissue antigen. Of these 3/5 also demonstrated the presence of IC which contained AAb reactive with the same kidney antigen.

reactive with the same kidney antigen. The presence of such AAb's and their formation of IC indicates that these antibodies may have a functional role in the initial immunological insult to the kidney.

1531 CONTINUOUS AMBULATORY PERITONEAL DIALYSIS (CAPD) IN CHILDREN. <u>Donald E. Potter, Tarran K. McDaid, Kathy</u> California, San Francisco.

We have evaluated CAPD in 9 children, 9-15 years of age for periods of 1.5-14 months (mean 6.9 months). All but one had previously been treated with hemodialysis(HD)or intermittent peritoneal dialysis. Training was accomplished in 1-16 days. A mean volume of 41 ml/kg was exchanged 4 times a day, 7 days a week. The mean peritoneal urea clearance was 3.5 ml/min and steady-State BUN and creatinine levels were 68 and 9.3 mg/dl, respectively. Other biochemical levels were:

Na	ĸ	CI	co_2	Ca	P	Cholesterol	Triglyceride
138	4.6	99	25	9.4	5.0	254	268
The r	nean h	ematoc	rit lev	vel was	; 21% a	and the transf	usion require-
ment was	s 0.13	units	per mo	onth. I	here v	vas a striking	improvement
in anemia in children transferring from HD to CAPD. Ultrafiltra-							
tion and maintenance of dry weight was easily achieved and the							
mean blood pressure was 110/74. Dialysate protein loss was 0.17							
g/kg/day, dietary protein intake 2.1 g/kg/day, and serum albumin							
3.6 g/dl. Glucose absorption from dialysate was 2.5 g/kg/day.							
Calorie intake from diet plus dialysis was 58 kcal/kg/day. The							
mean growth rate of 3 children was 2.7 cm/year. Peritonitis was							
a major problem with one episode every 4.3 months. Three child-							
ren discontinued CAPD because of infection. Despite peritonitis							
and catheter complications children were enthusiastic about CAPD							
and preferred it to HD. CAPD has subjective and objective bene-							
fits in children and is an important new form of therapy.							

1532 RENIN PROFILING IN CHILDHOOD HYPERTENSION(HBP). James W.Prebis, Alan B.Gruskin, Martin S.Polinsky, H.Jorge Baluarte. St.Christopher's Hos.for Children, Section of Nephrology, Dept. Ped., Temple Univ. Med.Sch., Phila., PA.

Renin profiling(comparison of plasma renin activity(PRA)to urinary sodium excretion) is usually not a routine segment of the diagnostic evaluation of (HBP) children and adolescents. PRA and urinary sodium excretion were performed in a nonstimulated manner in 28 normal children,12 males and 16 females,11-19 years old and in 68 HBP pediatric patients; 43 had essential HBP and 25 an un-derlying renal etiology. The mean PRA±1SD in the control populaderiving femal settings. The mean restriction ranged from 62-356 mEq/ $1.73m^2/24hr$. No correlation existed between sodium excretion and PRA. The PRA of the essential HBP was 5.4t3.3 while that of the renal HBP was 15.8±7.7. The mean PRA of the renal HBP was significantly higher than that of the normals (p<0.001) and the essential HBP(p<0.001). 21/25(84%) of renal HBP had high renin HBP when hyperreninemia was defined as a PRA >2S.D. above the control mean, while 7/43(16%) of the essential HBP had high renin HBP. This study demonstrates that renin profiling is of diagnostic value in childhood HBP. The presence of a high PRA HBP indicates the liklihood of a renal etiology, but as in adult studies about 15% of pediatric patients with essential HBP also have high PRA. Renin profiling in an unstimulated state as performed here cannot accurately identify children with low and/or suppressed PRA. Stimulated renin measurements may identify renal HBP with normal PRA as well as differentiate high PRA essential HBP from renal Supported by RR75 and HL23511-01. HBP.

URINARY N-ACETYL GLUCOSAMINIDASE (NAG) AS AN INDICAT- **1533** OR OF GENTAMICIN (G) NEPHROTOXICITY IN PREMATURE IN-FANTS. <u>Rajchgot, P., MacLeod, S., Klein, J., Chabot,</u> J. and <u>Radde, I</u>. (Sponsored by A. Sass-Kortsak). Dept. of Paediatrics, Hospital for Sick Children, Toronto. Urinary concentrations of NAG reflect nephrotoxicity and are

generally expressed per mg creatinine (Cr). We studied the time course of G-induced renal effects in 22 neonates of 32-36 wks gestational age. Gp A (n=13) received G; Gp B (n=9) otherwise comparable but not requiring G served as controls. Timed urine collections started on day of first G treatment (Gp A) or on admission (Gp B). NAG, β_2 -microglobulin (β_2 -M) and Cr were assayed.
 NAG excretion
 (units/min) (mtsEM)
 Day 1
 Day 3
 Day 4-7

 Gp A (while receiving G)
 84.2±15.3
 109±17.6
 91.2±18.1

 Gp B (*p<0.01)</td>
 27.4±6.1*
 36.2±8.9*
 35.0±8.7*
Day 4-7 91.2±18.1 NAG increased within 11-25 h of starting G and decreased to control 2-3 days after cessation of G. The pattern with β_2 -M was similar. The rapid decline of NAG following cessation of G suggests that <u>de novo</u> exposure to the drug is more important in in-ducing enzymuria than is persistence of G in renal tubular cells. Urinary Cr was not different between Gps (p>0.2), but rose with postnatal age (PNA):0.015±0.002 mg/min (PNA 1 day); 0.017±0.002 (PNA 3 days); 0.028 \pm 0.004 (PNA 4-7 days). Serial values of NAG/mg Cr were also greater in Gp A (p<0.01). Although age-related increase in U Cr was not a problem with serial evaluations as described here, in analysis of subtle differences in comparative drug studies, timed output of NAG may be a more reliable indicator of nephrotoxicity when studies are not controlled for age.