84A ABSTRACTS

LEFT VENTRICULAR DUAL-ENERGY SUBTRACTION ANGIOGRAPHY IN PATIENTS: A MOTION IMMUNE DIGITAL SUBTRACTION TECHNIQUE

Michs S. Van Lysel, Ph.D., Dale G. Senior, M.D., Vinod K. Gupta, M.D., David J. Ende, M.D., David J. Albright, B.S., William P. Miller, M.D., F.A.C.C., Section of Cardiology, University of Wisconsin, Madison, WI

Left ventricular (LV) digital subtraction angiography (DSA) using the standard temporal subtraction technique is limited by misregistration artifacts caused by cardiac, respiratory and gross patient motion. Motion artifacts are particularly severe during temporal subtraction DSA exercise tress studies. Dual-energy DSA imaging, in which 30 Hz x-ray pulses alternate between 70 and 120 kVp, is insensitive to patient motion. In this initial patient study, the image quality of dual-energy DSA was assessed and compared with standard cine ventriculography. Eight patients, ranging in weight from 54-100 kg and in cardiac index from 2.4-4.3 L/min/m² (mean 3.5±0.7), underwent 30° RAO direct LV injection cine ventriculography. Fifteen minutes later 40 ml of Hypaque-75 was injected ventricular contour by dual-energy DSA images of the left ventricle were obtained. Subjective visualization of the left ventricular contour by dual-energy DSA was good in all cases, and image quality did not deteriorate with patient motion. To assess the ability to visually detect left ventricular endocardial borders by dual-energy DSA, LV end-systolic and end-diastolic volumes as determined by the area-length method using direct LV injection cine (C) and dual-energy DSA (DE) were compared. The regression line was DE=0.93 C + 4.6 ml. The r value was 0.96. The range of volumes was 15-216 ml. We conclude that dualenergy DSA from a pulmonary artery injection of contrast produces high spatial resolution images of the left ventricle, even in the presence of patient motion. The motion immunity of dual-energy DSA may allow for the essessment of left ventricular function during exercise in patients using a venous injection of contrast.

Tuesday, March 20, 1990 10:30AM-12:00NOON, Room 23 Cardiac Transplantation: Donor Allograft Physiology and Adaptive Responses

EVIDENCE FOR SYMPATHETIC REINNERVATION AFTER CARDIAC TRANSPLANTATION IN HUMANS Robert F. Wilson M.D., Betsy V. Christensen B.S., Ada Simon Ph.D., Maria-Teresa Olivari M.D., and Carl W. White M.D., F.A.C.C. University of Minnesota, Minnesota, MN

Ph.D., Mana-Teresa Olivari M.D., and Carl W. White M.D., F.A.C.C. University of Minnesota, Minneapolis, MN Cardiac reinnervation after orthotopic transplantation (CT) occurs in animal models but human studies suggest that reinnervation does not take place. To more specifically determine if sympathetic reinnervation occurs in transplanted human hearts, we measured myocardial norepinephrine (NE) release [Δ[NE] in blood simultaneously obtained from the ascending aorta (Ao) and coronary sinus(CS)] at rest, after tyramine (an agent that causes degranulation of infact sympathetic nerve terminals, 0.55μg/kg, IV), and after sustained isometric handgrip (a physiologic stimulus) in 3 patients <2 months after CT, 17 patients ≥1 year after CT, and 4 normal patients. The reproducibility of Δ[NE]_{CS-Ao} measurements (radioenzymatic method) was assessed by comparing repeat measurements (n=12, mean absolute difference ±SD=39±41pg/ml), Evidence of reinnervation was defined as an intervention-induced increase in Δ[NE]_{CS-Ao} by >3SD control (i.e.> 123pg/ml).

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Results':		basal	peak+	ΔΑΡ(mmHg)	Δheart				
intervention	n after CT	A[NE]CS-No	Δ[NE]CS-Ao	sys `		rate(bpm)				
tyramine	<2mo	27±21	12±22	27±6	16±4	1±2				
	≥lyr	48±15	439±78°†	19±3	9±2	4±1				
	normal	42±41	1477±147°	31±9	9±7	9±7.				
hangrip	<2mo (n=2)		29±20	24±4	13±4	4±3				
	≥1 yr (n=6)		263±97*	18±4	22±3	2±3				
	normal (n=3))	217±130*	22±5	10±7	9±7				

AmenatSEM, + after intervention, *p<.05 vs. \$2mo after CT, †p<.05 vs.normal

13 of 17 patients studied ≥1yr after CT had a significant transmyocardial release of NE after tyramine and all of those patients had a significant NE release during isometric hangrip (Δ[NE]cs-Ao=340±113pg/ml). Conclusion: Early after CT the myocardium does not release NE, suggesting denervation. Late after CT, the majority of patients can release substantial, but subnormal, quantities of NE in response to tyramine and physiologic stimuli, suggesting that limited sympathetic reinnervation occurs in the majority of transplanted human hearts.

CARDIAC DONOR HEART DYSFUNCTION: EVIDENCE FOR CATECHOLAMINE-MEDIATED MYOCARDIAL INJURY

Robert J. Wiechmann, M.D., Ted Eastburn, M.D., June Murray, R.N., J. David Port, B.S., John B. O'Connell, M.D., F.A.C.C., Dale G. Renlund, M.D., F.A.C.C., Ray E. Hershberger, M.D., Michael R. Bristow, M.D., Ph.D., F.A.C.C. University of Utah, Salt Lake City, UT

B R fmol/mg Tissue Cat (ng/g) CR mg ISO Ca2+ <u>\$₁</u> <u>\$₂</u> *92.2 18.6 NE EPI 862 *1230 *880 64 540 (n=7) ±134 ±265 ±6.6 ±3.9 ±249 ±16 ±132 Normals 1848 1218 71.2 18.6 658 48 457 (n=14) ±238 ±200 ±5.0 ±1.8 ±95 ±7.0 ±79 CONCLUSIONS: 1) Organ donors with unexplained DHD have a marked decrease in isoproterenol stimulated muscle contraction despite normal-increased &, R levels. 2) This uncoupling of DHD &-adrenergic R is consistent with exposure to high levels of adrenergic neurotransmitter, which may be the cause of DHD.

INCREASED EXPRESSION OF $\&plantsize{B}_2$ -ADRENERGIC RECEPTORS IN SURGICALLY DENERVATED, PREVIOUSLY TRANSPLANTED HUMAN VENTRICULAR MYOCARDIUM

J. David Port, B.S., Lisa Skerl, John B. O'Connell, M.D., F.A.C.C., Dale G. Renlund, M.D., F.A.C.C., Patti Larrabee, B.S., Michael R. Bristow, M.D., Ph.D., F.A.C.C. University of Utah, Salt Lake City, UT

We have previously reported that the surgically denervated, transplanted human heart (TX) exhibits presynaptic supersensitivity to catecholamines but normal total ß receptor density. We now report measurement of ß receptor subtypes, tissue norepinephrine (NE) and adenylate cyclase (AC) in material prepared from right and left ventricular myocardium (VM) removed from transplant recipients with normal global LV function who were retransplanted for graft atherosclerosis (±SEM): *p<0.05

Grou	P Rece	ptor	Density,	A	AC, pmol					
$\langle u \rangle$	(n) fmol/mg			cA	cAMP/min/mg					
	Total	B B	1 B ₂	ISO	NaF	Forsk				
TX	93.6	62	.2 31.0*	40.4*	38.7	219	47×			
(10)	±12.4	±9	.0 ±5.1	±3.1	±4.6	±22	±19			
NL†	94.7	75	.2 18.6	30.5	35.6	238	817			
(68)	±3.6	±3	.3 ±1.4	±1.6	±2.0	±12	±92			
normal innervated VM from organ donor controls;										
ISO-isoproterenol; Forsk-forskolin; NE-norepinephrine										
CONCLUSION: TX contains an increased density of B2										
adrenergic receptors. Since the AC response to ISO is										
mediated predominantly by \$2 receptors in human VM, the										
increased \$2 population appears to be coupled to a										
functional response. Thus TX is not only supersensitive										
to circulating epinephrine by virtue of denervation and										
loss of uptake ₁ , but the B_2 "epinephrine receptor" is										
also increased in density in TX VM.										
also increased in density in IX vm.										