from published nomograms based on surface area. The initial TV Z-score was significantly smaller in those undergoing shunt than in those undergoing various RV outflow tract procedures (RVOT) with or without concomitant shunt (-2.2 \pm 0.3, -0.8 \pm 0.2 respectively, p = 0.0002, data shown as mean ± SEM). However the change in TV Z-score post-procedure was similar (shunt -0.7 ± 0.4 , RVOT -0.8 ± 0.2 , p = NS). Similarly, the RV inlet Z-score was smaller in the shunt group than the RVOT group (-1.6 \pm 0.1, -1.2 \pm 0.1 respectively, p = 0.0009). Again, the change in the Z-score was similar (shunt -0.06 ± 0.1 , RVOT -0.3 ± 0.1 , p = NS). Of those patients undergoing RVOT alone, neither type of procedure (catheter versus surgery), nor requirement for a shunt within 6 weeks of procedure allected the change in TV or RV inist Z-score. However, in patients with primary RVOT, change in TV and RV inlet Z-score was weakly correlated with TV and RV Z-scores pre-procedure. Larger increases in TV and RV inlet size post-procedure were associated with smaller initial TV and RV inlet sizes (r = 0.47, p = 0.01 and r = 0.44, p = 0.01 respectively). There was no such correlation in patients shunted alone. Following establishment of RV to PA continuity, growth of the TV and RV frequently does not exceed somatic growth but is more likely in those ventricles with small initial TV and RVs.

997-13 Postnatal Left Ventricular Growth Potential in Selected Infants With Non-Apex-Forming Left Ventricles

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To evaluate left ventricular (LV) growth potential, the echocardiograms of 5 term infants presenting with lesions including a marginally sized, but morphologically normal LV that was not apex-forming were reviewed. Four infants initially required a PDA to maintain cardiac output and received prostaglandins for 1.5 to 8 wks. Because of LV hypoplasia, 2 were listed for transplantation. Diagnoses were aortic stenosis in 4 (with coarctation in 3) and polysplenia in 1. Echo measurements at birth (3.6 ± 2 days) (mean \pm SEM) were compared to measurements at 1 mo (25.4 ± 2 days). Over this time period, weight (3.0 ± 0.1 vs 2.9 ± 0.1 kg) and BSA (0.2 ± 0.01 vs 0.2 ± 0.01 m²) did not change. However, there were significant increases in the diameters of the aortic annulus (ANN, mm) and aortic root (ROOT, mm), mitral valve area (MVA, cm²), LV end-diastolic volume (LVED, cc), LV end-systolic volume (LVES, cc), LV area (LVA, cm²), and the ratio of the long axis of the LV to the long axis of the heart (LAR).

	ANN	ROOT	MVA	LVED	LVES	LVA	LAR
							0.80 ± 0.1
1 Mo	5.7 ± 0.4*	7.3±0.6*	$0.7 \pm 0.1^{\circ}$	$5.6 \pm 0.8^{*}$	$2.2 \pm 0.5^{*}$	$3.8 \pm 0.4^{\circ}$	$0.91 \pm 0.1^{\circ}$

*indicates p < 0.05 when compared to birth.

Tricuspid valve diameter ($12 \pm 1 \text{ vs}$ $12 \pm 0.6 \text{ mm}$), long axis of the heart ($26 \pm 2 \text{ vs}$ $28 \pm 2 \text{ mm}$) and LVEF ($53 \pm 13 \text{ vs}$ $55 \pm 10\%$) did not change. Four of 5 infants had surgery with 1 death. At follow-up (1-4 yr), all 4 surviving infants have biventricular physiology. Echo at 290 ± 41 days showed normal LAR (0.97 ± 0.1). These data suggest that a morphologically normal, but non-apex-forming LV may have postnatal growth potential. Because LV size influences treatment options, a period of observation (maintaining ductal patency, if necessary) with serial echo measurements may be warranted in selected infants.

997-14 Progressive Aortic Valve Insufficiency Due to Kawasaki Disease

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Valvular involvements have been recognized as one of the cardiovascular sequelae of Kawasaki disease, and may become a fatal complication. In this regard, little is known about the clinical features of aortic insufficiency (AI) in this disease. From 1973 to 1994, 1588 patients were seen in our institution, of whom 4 (0.3%) cases complicated with audible AI. All patients were maie, whose onset of disease were ranged from 2 to 10 months. Aspirin alone or aspirin plus steroid were chosen as the acute stage treatment in 2 of each cases. All but 1 complicated with coronary artery aneurysms, and these lesions have not been resolved up to date. Heart murmur was noted at 1.4, 1.5, 7.0 and 15.0 years after the onset of disease, respectively. In 2 of 4 cases. Al was progressively worsened and valve replacement was required at 13 years after the onset of AI. On macroscopic view, the leaflets of excised valves were firmly thickened and rolled over. Calcilication and nodular change of leaflets, which frequently seen in rheumatic valvular disease, were not observed. Histologic examination revealed that massive proliferation of collagen fiber without acute inflammatory changes, which suggested that these abnormalities may be induced by the healing process of acute inflammatory changes and progress due to the long-standing hemodynamic stress. In other 2 cases, whose onset of AI were \geq 7 years after the onset of disease,

the degree of AI has not progressed during the follow-up. Although we could not address the long-term outcome of mild AI which occurred in the late convalescent stage, the more than moderate degree of AI is progressive and may require valve replacement. The oossibility of AI should be considered if new diastolic murmur appears during the follow-up, and careful long-term follow-up would be recommended.

998 Cardiac Pacing: Technical Advances and Experimental Observations

Wednesday, March 27, 1996, 9:00 a.m.–11:00 a.m. Orange County Convention Center, Hall E Presentation Hour: 9:00 a.m.–10:00 a.m.

998-69

Chronic Ventricular Pacing at Physiological Heart Rate Affects Local Myocardial Blood Flow and Local Wall Thickness

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Previous acute canine experiments indicated that asynchronous electrical activation reduces mechanical load and blood flow in early activated regions and increases them in late activated regions. We invostigated the shuctural adaptation and the distribution of blood flow in the left ventricular (LV) wall during chronic asynchronous activation, as induced by ventricular pacing. Under general anaesthesia 6 dogs received a pacemaker, with the stimulus electrode implanted 1 cm below the base of the LV free wall (LVFW). After full recovery the dogs were paced chronically (CVP) at physiological heart rate (A-V sequential, A-V interval 25 ms), for six months. On 2D-echocardio-grams the cross-section of the LV wall at the level of the stimulation electrode was divided into 6 sectors, from which mean local wall thickness was measured. Myocardial blood flow (microspheres) was determined in these sectors: during sinus rhythm (SR) and 15 min after ventricular pacing (VP), both at time of implantation and termination procedures. Wall thickness of the early activated LVFW decreased by $6.5 \pm 8.8\%$ and thickness of the late activated septum (s) increased by $24.0 \pm 12.2\%$ (p < 0.05) after CVP, compared with baseline. The ratio of LVFW and S (LVFW/S) thickness decreased from 1.17 \pm 0.11 (baseline) to 0.88 \pm 0.11 (CVP). The LVFW/S blood flow ratio was calculated as an index of blood flow distribution. This ratio was 0.81 ± 0.22 during SR and decreased to 0.62 ± 0.10 after 15 min of VP. After 6 months of continuous VP the LVFW/S ratio had returned to baseline values (0.80 ± 0.22) and increased to 1.19 \pm 0.22 (p < 0.05) 15 min after return to SR. We conclude that 1) myocardial structural adaptation to mechanical overload occurs locally, 2) local wall mass adapts so that myocardial blood flow per unit weight returns to initial (pre-pacing) values, so 3) changes of myocardial blood flow caused by VP predict changes in local wall mass.

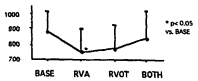


Effect of Pacing at Alternate and Combined Sites in the Right Ventricle on Systolic and Diastolic Function

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We hypothesized that pacing at 2 ventricular sites simultaneously would activate the myocardium more rapidly and improve function. Others have shown that right ventricular outflow tract (RVOT) pacing produces a better cardiac output (CO) than traditional RV apical (RVA) pacing. In 7 pts with EF < 40% (30 ± 8%) we measured QRS duration, CO, EF, dP/dt, -dP/dt, and Tau during baseline rhythm, DVI pacing at the RVA, the RVOT and both RV sites in random order. There were no significant differences in CO, EF, LV pressure, and Tau with different pacing sites. Significant differences were noted in peak dP/dT, -dP/dT, and QRS. With RVA pacing dP/dt was the shortest and the QRS the longest. As shown, dP/dT then tended to improve progressively with pacing in the RVOT and at 2 sites. dP/dt was inversely related to the changes in QRS durations.

DP/DT with Pacing Interventions



These data suggest that in pts with poor LV function, there are subtle

improvements in diastolic and systolic function with pacing in the RVOT and at combined sites in the RV compared to RVA pacing. Thus asynchrony with prolongation of QRS is an important determinant of LV function.

998-71 Sensing of Mechanical Atrial Systole — A Novel Technique for VDD Pacing

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Single-lead VDD pacemakers are limited by inconsistent atrial sensing. In addition, the latency between electrical and mechanical atrial systole may be variable, making optimization of AV delay difficult. The purpose of the current study was to assess the feasibility of sensing mechanical atrial systole using a catheter-based piezoslectric crystal (XTAL). A quadripolar electrode catheter was modified by adding an XTAL (7 × 2 mm) between poles 1 and 2. In 5 closed-chest dogs, the XTAL catheter was placed against the inferolateral wall of the right atrium. XTAL output (bandpass 1-25 Hz), right atrial (RA) pressure and RA electrograms (EGM) were recorded at baseline, during infusion of isoproterenol (1.5 µg/kg/min), during RA pacing, and after induction of complete AV block using radiofrequency catheter ablation. Results: Stable XTAL signals were recorded in all animals. Mean XTAL amplitude was 28 \pm 20 mV in sinus rhythm (SR) and 15 ± 5 mV during RA pacing. There was no detectable "farfield" signal resulting from ventricular systole during any recording. There was less variability in the latency between the peak of the XTAL signal and the peak of the A wave than between the RAEGM and the A wave (8.2 \pm 6.4 ms vs. 15.1 \pm 6.8 ms, p < 0.01). Complete AV block did not affect the morphology or timing of the XTAL signal. Conclusions: A catheter-based XTAL placed against the lateral RA produces signals that are large in amplitude and correlate more closely with the peak of the A wave than the endocardial EGM. Further study of XTAL-based single-lead VDD pacing systems is warranted.

998-72 Serial Fluoroscopic Evaluation of the Telectronics Accutix^{**} 330–801 "J"-Shaped Atrial Pacing Lead

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A recall cf the Telectronics Accufix[™] 330–801 atrial lead was issued in November 1994, because of fracture of the "J"-shaped retention wire. To assess the integrity of the retention wire, we have performed fluoroscopic screening on 209 patients with the 330–801 atrial lead in place. Each retention wire was classified as normal, indeterminate, fractured without protrusion, and fractured with protrusion. Cine films of all 135 patients originally classified as normal were re-evaluated after the original screening. Eighty patients have undergone two serial fluoroscopic evaluations separated by 158 ± 5.4 days

Nine retention wires originally classified as normal were re-classified as fracture without protrusion, and two were re-classified as fracture with protrusion when the original films were re-evaluated. Eight of the 11 leads reclassified were screened during our first two days of fluoroscopic evaluation of this lead. Of the 80 patients who underwent a second fluoroscopic screening, 70 were initially classified as normal (nine of which were later re-classified as described above), one fractured, and nine indeterminate. There were no new fractures detected in this group; of the indeterminate group, four were reclassified as fractured after rescreening, three were classified as normal, and two remained indeterminate. The fractured retention wires in this group did not appear to change during this period.

We conclude that there is a definite learning curve associated with fluoroscopic evaluation of this lead, and recommend that screening be performed by cardiologists experienced in fluoroscopic screening. Fractured retention wires remained stable in appearance during this observation time, and no new retention wire fractures were detected.

998-73 Assessment of Pacing Lead Curvature and Strain With Three Dimensional Reconstruction of Biplane Cineangiographic Images in Vivo

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Pacing lead integrity is vital to pacemaker function and patient safety. Understanding the point of maximum strain on implanted pacing leads will help in lead design to minimize the occurrence of lead fracture. However, there has been no study on pacing lead strain in vivo. The dynamic position of newly implanted pacing leads were imaged with biplane cineangiography. The three dimensional (3D) description of the leads in each frame was determined by a triangulation technique previously developed in quantilative arteriography for determining stenosis or calheter location in space. Both atrial and ventricular leads were studied in each of 8 patients. To estimate strain in the lead wire insulation, the 3D curvature was calculated from geometric data using a seven point smoothing operation. The components of curvature were calculated at 23 ± 7 (range 11–31) frames. The curvature magnitude was computed from the components. *Results:* High localized areas of curvature were note to 0 to 6 cm from the proximal electrode of the atrial lead, corresponding ... the curvature of the J-loop. Significant components of curvature occurred in the anterior-posterior direction. The peak curvature of the atrial lead was 2.5 ± 0.5/cm, indicating a minimum radius of 4.1 ± 0.9 mm. Contrarily, the locations of the ventricular lead curvature of the ventricular lead was 2.0 ± 0.6/cm, indicating a minimum radius of 5.6 ± 0.3 mm.

Conclusions: Using 3D reconstruction of lead position in space and a seven point smoothing operation for calculation of curvature, it is feasible to localize the areas of curvature and quantify the magnitude of curvature for pacing leads in vivo. The information derived from such analysis in large scale studies of acute and chronic leads should be important for improvement in the design of pacing leads to reduce the incidence of lead fracture.

998-74 Low Pacing Thresholds for Heart Wires Using a New Stimulation Technique

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OverLapping Biphasic (OLBI) stimulation of the heart is a new way of biphasic pacing by two pulses of different polarity. A significant threshold reduction has been reported in initial experiments.

Methods: External OLBI-stimulation was tested in 20 consecutive patients who underwent bypass or heart valve surgery before. Intraoperatively a bipolar heart wire (MHW 60-BP, BIOTRONIK) was fixed onto the right ventricle and a second (MHW 60-UP, BIOTRONIK) subcutaneous working as an indifferent electrode. Measurements of the threshold were performed up to the 3th postoperative day and were compared with thresholds of conventional unipolar and bipolar stimulation at the same time.

Results:

	Unipolar	Bipolar	OLBI
Day of surgery	1.5 ± 0.6 V	1.1 ± 0.4 V	0.5 ± 0.1 V
4th p.o. day	6.8 ± 2,7 V	4.3 ± 1.5 V	1.9 ± 0.7 V
8th p.o. day	>15 V**	8.8 ± 4.7 V**	3.5 ± 1.1 V

**In 4 patients no effective unipolar and bipolar stimulation was possible at day 8 but for all patients by OLBI stimulation.

Conclusion: 1) OLBI-stimulation is effective and shows lower thresholds compared with conventional unipolar and bipolar stimulation. 2) The increase of pacing threshold is less pronounced for OLBI-stimulation. 3) Therefore eflective cardiac stimulation is possible for a longer period and gives more safety, especially in therapy of patients with continued arrythmias postoperatively. 4) Using OLBI-stimulation irritation of the myocardial tissue caused by stimulation is reduced.

998-75 The Effect of Ventricular Pacing on Coronary Blood Flow

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Ventricular pacing is thought to produce the impairment of LV function by altering the sequence of ventricular activation and AV dyssynchrony. We hypothesized that ventricular pacing may produce deleterious effect on coronary blood flow (CBF). To test this hypothesis, we studied the effect of ventricular pacing on phasic coronary flow velocity (CFV), coronary arterial diameter (CAD) and coronary flow reserve (CFR) in 15 patients with normal coronary arteries. CFV, CAD and CFR in the left anterior descending coronary artery were measured using doppler flow wire during sinus rhythm, and during both atrial and ventricular pacing at a rate of 100 bpm in the same patient. Double product increased significantly during both pacing. CAD ouring ventricular (% increase: 16.4 ± 20.2%; P < 0.001) and atrial pacing (5.8 ± 10.1%; P < 0.001) significantly increased compared to that rluring sinus rhythm. APV during ventricular pacing (-18.9 \pm 12.9%) significantly decreased compared to that during sinus rhythm (P < 0.001) and during atrial pacing (5.0 \pm 19.9%; P < 0.005). CBF increased significantly during atrial pacing (34.9 ± 20.3%; P < 0.001), but not during ventricular pacing (24.5 ± 45.4%; P ≈ NS). Since the hyperemic flow after the administration of papaverine was not different among them, CFR during both atrial (3.8 \pm 1.3) and ventricular pacing (3.8 \pm 0.9) decreased compared to that during sinus rhythm (4.5 \pm 1.5). There was a significant positive correlation between CFR during sinus rhythm and the ratio of CBF during ventricular and atrial paring (R2 = 0.84, P < 0.001). Also