choice has changed in favour of rotary pumps; 19%, 69% and 96% for E1, E2 and E3 respectively. Median duration of VAD support increased from 84 days (IQR 20–209) in E1 to 280 days (IQR 86–661) in E3 (p<0.01). Overall survival to 1 year after VAD implant rose from 52.9% (95%CI 40 to 64) in E1 to 65.6% (95%CI 54 to 75) in E3 (p=0.10). Of the 239 patients implanted, 83 (35%) have undergone HTx, 52 (22%) are alive on VAD support & 84 (35%) died on support. Twenty were explanted following myocardial recovery; 18 of these remain alive & 2 died. Survival after HTx for patients with or without a pre-HTx VAD was 81.4% (95%CI 71 to 88) & 90.3% (95%CI 88 to 92) respectively at 30-days (p<0.01) and 80.0% (95%CI 63 to 82) & 84.3% (95%CI 82 to 87) respectively at 1-year (p<0.01). 1-year survival conditional on 30-day survival was similar with & without a pre-HTx VAD (93% vs 91%, p=0.48).

Conclusion Heart transplant activity has declined and waiting times have become prolonged leading to an increased need for bridging to transplantation. There has been a shift from volume displacement VADs to rotary blood pumps and the duration of support has increased. Post VAD survival has improved. While bridging appears to increase mortality early after HTx, longer term survival is unaffected.

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CLINICAL AND HAEMODYNAMIC STATUS BEYOND 3 MONTHS OF MECHANICAL SUPPORT WITH THE HEARTWARE VENTRICULAR ASSIST DEVICE

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Introduction Limited data exist on the longer term clinical and haemodynamic impact of the HeartWare left ventricular assist device (HVAD®) when used as a bridge to heart transplantation. Patients who had a device longer than 3 months were reviewed. Methods 26 patients had a HVAD implanted from 07/2009 to 07/ 2010 (mean age 46.8 years, 18 male, 5174 total days of support). Baseline and follow-up NYHA functional class, peak VO2 (bicycle exercise), right heart haemodynamics, biochemistry and mortality outcome were compared using paired t test. Results: 22/26 (85%) patients survived beyond 3 months. 4 patients died before (mean survival 40 days, 2 stroke and 2 multi-organ failure) and 2 died after (mean survival 173 days, 1 stroke, 1 right heart failure) discharge from hospital. 2 patients were transplanted (at 3 and 241 days after implant) and 1 had recovery of LV function. Follow-up data is available for 14/20 survivors (mean 197 days from implant). Significant results are shown in the Abstract 83 table 1. There was no significant change in peak VO_2 (9.9±1.8 to 12.9±3.8, p=0.08),

Abstract 83 Table 1

to 105 ± 38 , p=0.19).

Parameter	Baseline	Follow-up	p Value
NYHA functional class	3.6 ± 0.4	2.1 ± 0.6	< 0.001
Mean PA pressure (mm Hg)	38±9	21 ± 8	< 0.001
Mean PW pressure (mm Hg)	25±5	10±5	< 0.001
Transpulmonary gradient (mm Hg)	12±5	9±3	0.02
Right atrial pressure (mm Hg)	11±6	5 ± 4	0.006
PA oxygen saturation (%)	51 ± 8	66±7	0.003
Cardiac Output (I/min)	$2.9 \!\pm\! 0.8$	4.3 ± 0.9	< 0.001
Sodium (mmol/l)	$134\!\pm\!4$	139±3	0.002

haemoglobin (12.7 \pm 1.7 to 12.1 \pm 1.2, p=0.3) or creatinine (122 \pm 41

Conclusions The HVAD® results in significant improvement in functional class, right heart haemodynamics, cardiac output and sodium levels beyond 3 months of therapy. Ongoing randomised clinical trials will establish the long-term outcome of this device.

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TREATMENT OF REFRACTORY RIGHT HEART FAILURE AFTER IMPLANTATION OF A LEFT VENTRICULAR ASSIST DEVICE. IS THE LEVITRONIX CENTRIMAG RIGHT HEART SUPPORT A SOLUTION?

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Introduction Right heart failure after left Ventricular Assist Device (LVAD) implantation is a severe complication, in extreme cases necessitating additional mechanical assist. We present our institutional experience with the Levitronix CentriMag used for right ventricular support commencing LVAD implantation with refractory right ventricular failure.

Material and Methods Between March 2001 and November 2010 109 patients underwent implantation of long term, total implantable, continuous flow LVADs: 60 HeartMate II, 25 Jarvik 2000 and 24 HeartWare. All patients requiring right ventricular support were included (n=24), for which the Levitronix CentriMag continuous flow, paracorporeal device was used. The analysis included patient demographics as well as overall duration of support and outcome parameters, including survival at 30, 90 days and 1 year.

Results 24 pts. underwent implantation, age 37.9±13.7 years, gender: M/F-15/9, underlying disease: dilated cardiomyopathy 22 (92%), peripartum cardiomyopathy 1(4%), viral myocarditis 1(4%). Median duration of support: 28 days (5–146). 3(12.5%) pts. underwent heart transplantation (HTx) on RV support, 14(58.5%) underwent RVAD explantation. Of these, 3 underwent successful HTx, 4 recovered LV function and underwent successful LVAD explantation, 3 remain on continuing LVAD support, 4 patients died after RVAD explantation (post explantation day 1, months 3 and 4 and at 2 years), 7(29%) patients died during RV support. Median ITU/hospital stay: 19.5 days (6–145)/78.5 days (10–219). 30-day/90-day/1-year survival: 79%/71%/60%. 15(62.5%) patients were discharged from hospital after treatment. Median survival after procedure: 473.5 days (10–1917).

Conclusion Levitronix CentriMag right ventricular support is an excellent option for post LVAD implantation treatment of refractory RV failure. It allows either bridging to transplantation or RV function improvement and provides an acceptable rate of survival.

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PREDICTION OF RESPONSE TO BIVENTRICULAR PACING FROM DYSSYNCHRONY INDICES: THE ABSOLUTE LIMIT ON PREDICTABILITY, AND ITS CLINICAL IMPLICATIONS

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Background It may be incorrect to believe that, with a good echocardiographic marker of mechanical dyssynchrony, response to biventricular pacing (BVP) should be predictable with a high r^2 value. Variability between repeat echocardiographic measurements, and between successive dyssynchrony measurements, may reduce r^2 . Both will mandatorily limit the achievable r^2 ; we determine this "contraction factor".

Method and Results We compared correlation coefficients of dyssynchrony indices with response markers, in externally monitored randomised controlled trials (EMRCTs) and highly skilled single centre studies (HSSCSs). Δ LVEF in CRT recipients comprises true CRT effect plus unpredictable spontaneous variability present in control patients (Abstract 85 figure 1, upper panel). The resultant depression in r^2 is calculated. HSSCSs overstate r^2 between

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