miniml and non-significant bias between the two modalities (bias= 0.01 m/s; p = 0.91 and -0.08 m/s; p = 0.91). This novel automated method demonstrated excellent reproducibility (Coefficient of variability 2.67% for peak E-wave mitral inflow velocity, Coefficient of variability 1.93% for peak A-wave mitral inflow velocity).

Conclusion We present a novel automated time-resolved transvalvular peak velocity assessment solution that can be used clinically for mitral inflow assessment and would circumvent the limitations of pulse-wave doppler echocardiography. Future studies are warranted to explore the diagnostic and prognostic advantages of our novel automated technique for mitral inflow assessment.

17 PARTICIPANTS WITH DIABETES MELLITUS HAVE PRESERVED METABOLIC FLEXIBILITY

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Background Measurement of the Phosphocreatine/Adenosine Triphosphate (ATP) ratio along with the Creatine Kinase (CK) rate constant (CK_{k}), allows calculation of the ATP delivery rate (CK flux). Metabolic flexibility may be impaired both in heart failure with reduced ejection fraction (HFrEF) and diabetes mellitus (DM). It is unknown to what extent flexibility can be influenced by artificially altering the substrate available for metabolism.

Purpose To examine cardiac function and energetics in diabetic participants with normal cardiac function and HFrEF, clamped on either fatty acid (FA) or glucose metabolism.

Methods Participants with non-insulin dependent diabetic mellitus (NIDDM) with both normal cardiac function (NHDM) and HFrEF (HFDM) were recruited and received intravenous separate visits, before undergoing multi-parametric cardiac MRI at 3 Tesla. Cardiac volume and function, PCR/ATP and CK_{k} were assessed. CK flux was calculated as CK_{k} x PCR/ATP x 5.7 μmol (g wet weight)^{-1} (assumed ATP concentration).

Results 15 NHDM participants (14 male, age 61.5 ± 7.3 years) and 9 HFDM participants (7 male, age 69.4 ± 7.8 years) were recruited. Left ventricular ejection fraction (LVEF) at rest was higher on IL compared to both baseline fasting and GI for NHDM (baseline 59.1±3.8%, GI 59.4±4.3%, IL 62.8±3.5%; p=0.01), with a non-significant trend for HFDM (baseline 37.3±7.6%, GI 36.8±9.2%, IL 38.8±8.0%, p=0.12). For both NHDM and HFDM there was no difference in PCR/ATP (NHDM: GI 1.98±0.31, IL 1.97±0.24, p=0.99; HFDM: GI 1.82±0.36, IL 2.01±0.32, p=0.09) or CK flux (NHDM: GI 2.6±1.1 μmol (g wet weight)^{-1} s^{-1}, IL 1.8±1.2 μmol (g wet weight)^{-1} s^{-1}, p=0.08; HFDM: GI 1.6±1.7 μmol (g wet weight)^{-1} s^{-1}, IL 2.3±1.1 μmol (g wet weight)^{-1} s^{-1}, p=0.39).

Conclusion Diabetic participants with HFrEF and normal cardiac function appear to have increased resting LVEF when clamped on FA as opposed to glucose metabolism, without a significant change in energetic status. This may imply that metabolic flexibility is relatively preserved in these groups.

18 SUBCLINICAL MYOCARDIAL INFLAMMATION IN ADULTS WITH TYPE 2 DIABETES: A CLINICAL STUDY USING MYOCARDIAL T2 MAPPING

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Background Chronic hyperglycaemia in Type 2 diabetes (T2D) results in a systemic low-grade inflammatory state. Inflammation is a key instigator in the development of heart failure in T2D. Cardiovascular magnetic resonance (CMR) T2 mapping is a technique which identifies myocardial oedema. The utility of T2 mapping to identify subclinical oedema as a marker of inflammation in T2D is unknown. We hypothesise that T2 times will be higher in subjects with T2D.

Methods CMR imaging on a 3-Tesla scanner was performed on 182 participants who were free of symptomatic cardiovascular disease. T2 images were acquired using the Siemens MyoMap sequence at the mid-ventricular short-axis slice. Twenty participants underwent a repeat CMR scan within two weeks to assess the test-retest reproducibility of T2. Intra-class correlation coefficient (ICC) and Bland-Altman plots were generated to assess reproducibility. T2 values were compared using T-test and Mann-Whitney test as appropriate. Clinical determinants of T2 in T2D were assessed using multivariable linear regression.

Results 124 T2D (mean age 64±7, 66% male) and 40 controls (mean age 61±8, 60% male) were analysed. T2 times exhibited excellent intra-observer (ICC 0.98–0.99), moderate inter-observer (ICC 0.48–0.99), and poor test-retest variability (ICC 0.33–0.90). T2 times were significantly lower in subjects with T2D compared to controls (39.0±2.2 ms versus 40.1±2.9 ms, P=0.013). Stratification by sex revealed significantly lower T2 in females with T2D (39.4±2.4 ms versus 41.7±3.1 ms, P=0.003), but not in males, when compared to controls. Following multivariable adjustment, T2 time was positively associated with a non-white ethnicity (β=0.245, P=0.007) and diabetic duration (β=0.197, P=0.03) and inversely associated with systolic blood pressure (β=−0.215, P=0.018).

Conclusions T2 mapping has moderate-excellent observer variability but poor test-retest reproducibility in a cohort T2D. Lower T2 times in T2D may reflect early myocardial fibrosis but does not provide evidence of subclinical myocardial oedema and therefore is not able to detect low-grade myocardial inflammation.

19 CARDIAC MAGNETIC RESONANCE TO IDENTIFY RAISED LEFT VENTRICULAR FILLING PRESSURE


Background Non-invasive imaging is routinely used to estimate left ventricular (LV) filling pressures (LVFP) in heart failure (HF), as an alternative to right heart catheterisation (RHC). Transthoracic echocardiography (TTE) estimates of LVFP are frequently deployed but produce largely dichotomised data limiting flexible clinical use and perform less well in patients with heart failure with preserved ejection fraction (HFrEF).