1 TITLE: GLUCOCORTICOIDS REGULATE MITOCHONDRIAL 2 FATTY ACID OXIDATION IN FETAL CARDIOMYOCYTES 3 4 **RUNNING TITLE:** Glucocorticoid regulation of fatty acid oxidation 5 6 **AUTHORS:** Jessica R. Ivy<sup>1</sup>, Roderic N. Carter<sup>1</sup>, Jin-Feng Zhao<sup>2</sup>, 7 Charlotte Buckley<sup>1‡</sup>, Helena Urguijo<sup>1</sup>, Eva A. Rog-Zielinska<sup>1†</sup>, 8 Emma Panting<sup>1,3</sup>, Lenka Hrabalkova<sup>4</sup>, Cara Nicholson<sup>4</sup>, 9 Emma J. Agnew<sup>1#</sup>, Matthew W. Kemp<sup>5,6,7</sup>, Nicholas M. 10 Morton<sup>1</sup>, Sarah J. Stock, 8,4,6, Caitlin Wyrwoll<sup>3</sup>, Ian G. Ganley<sup>2</sup>, 11 Karen E. Chapman<sup>1,3\*</sup> 12 13 **AFFILIATIONS:** <sup>1</sup>University/BHF Centre for Cardiovascular Science, The 14 Queen's Medical Research Institute. The University of 15 Edinburgh, 47 Little France Crescent, Edinburgh, EH16 4TJ. 16 UK. <sup>2</sup>Medical Research Council Protein Phosphorylation and 17 Ubiquitylation Unit, University of Dundee, Dundee DD1 5EH. 18 3School of Human Sciences, The University of Western 19 Australia, Crawley, WA6009, Australia. 4The Centre for 20 Reproductive Health, The Queen's Medical Research 21 Institute, The University of Edinburgh, 47 Little France 22 Crescent, Edinburgh, EH16 4TJ, UK. 5Department of 23 Obstetrics and Gynaecology, Yong Loo Lin School of 24 Medicine, National University of Singapore, 1E Kent Ridge 25 Road, Singapore 119228, Republic of Singapore. 6Division of 26 Obstetrics and Gynaecology, The University of Western 27 Australia, Crawley, WA 6009, Australia. Centre for Perinatal 28 and Neonatal Medicine, Tohoku University Hospital, Sendai, 29 Japan. 8The Usher Institute, The University of Edinburgh, 47 30 Little France Crescent, Edinburgh, EH16 4UX, UK. Current 31 address: #EJA: Food Standards Scotland, Q Spur, Saughton 32 House, Broomhouse Dr, Edinburgh, EH11 3XD, †ER-Z: 33 Institute for Experimental Cardiovascular Medicine, 34 University Heart Center Freiburg · Bad Krozingen, and 35 Faculty of Medicine, University of Freiburg, Freiburg,

36 Germany, ‡CB: Strathclyde Institute of Pharmacy and 37 Biomedical Sciences, 161 Cathedral Street, Glasgow, G4 38 0RE 39 40 \*CORRESPONDENCE: Karen E. Chapman, University/BHF Centre for 41 Cardiovascular Science, The Queen's Medical Research 42 Institute, 47 Little France Crescent, Edinburgh, EH16 4TJ. 43 Tel: 44-131-242-6736 44 Fax: 44-131-242-6779; 45 Email: Karen.Chapman@ed.ac.uk 46 47 **KEY WORDS:** Glucocorticoid, cardiomyocytes, early-life programming, 48 heart, antenatal corticosteroids, preterm birth. 49 50 **CONTENTS CATEGORY:** Placenta, pregnancy and perinatal physiology 51 52 53 ABBREVIATIONS: GR - glucocorticoid receptor, E - embryonic day, P -54 postnatal day, PBS - phosphate buffered saline, SRB -55 sulforhodamine B, 2DG - 2 deoxyglucose, AR - antimycin 56 and rotenone, OCR - oxygen consumption rate, ECAR -57 extracellular acidification rate, FCCP - carbonyl cyanide-4-58 (trifluoromethoxy)phenylhydrazone, BSA - bovine serum 59 albumin, GFP - green fluorescent protein, DFP - deferiprone, 60 CPT-1 - carnitine palmitovltransferase-1, PGC-1 $\alpha$  - PPAR $\gamma$ 61 coactivator- $1\alpha$ , SD - standard deviation. 62 63 **FUNDING**: This work was supported by an MRC Project grant 64 (MR/P002811/1), a BHF Centre of Excellence award (RE/13/3/30183), BHF studentships (FS/13/52/30637 to EJA 65 66 and FS/08/065 to ER-Z), MRC funding to IGG 67 (MC\_UU\_00018/2), a Wellcome Trust Clinial Career Development Fellowship (209560/Z/17/Z to SJS), a grant 68 69 from the Western Australia Channel 7 Telethon Trust

70 (MWK), RNC was funded by a WT New Investigator Award 71 (100981/Z/13/Z) to NMM. 72 73 74 **ABSTRACT** 75 The late gestational rise in glucocorticoids contributes to the structural and functional 76 maturation of the perinatal heart. Here, we hypothesised that glucocorticoid action 77 contributes to the metabolic switch in perinatal cardiomyocytes from carbohydrate to fatty 78 acid oxidation. In primary mouse fetal cardiomyocytes, dexamethasone treatment induced 79 expression of genes involved in fatty acid oxidation and increased mitochondrial oxidation 80 of palmitate, dependent upon glucocorticoid receptor (GR). Dexamethasone did not, however, induce mitophagy or alter the morphology of the mitochondrial network. In 81 82 neonatal mice, dexamethasone treatment induced cardiac expression of fatty acid 83 oxidation genes in vivo. However, dexamethasone treatment of pregnant C57Bl/6 mice at 84 embryonic day (E)13.5 or E16.5 failed to induce fatty acid oxidation genes in fetal hearts 85 assessed 24 hours later. Instead, at E17.5, fatty acid oxidation genes were down-86 regulated by dexamethasone, as was GR itself. PGC-1α, required for glucocorticoid-87 induced maturation of primary mouse fetal cardiomyocytes in vitro, was down-regulated in 88 vivo in fetal hearts at E17.5, 24 hours after dexamethasone administration. Similarly, 89 following a course of antenatal corticosteroids in a sheep model of preterm birth, both GR 90 and PGC-1\alpha were down-regulated in fetal heart. These data suggest endogenous 91 glucocorticoids support the perinatal switch to fatty acid oxidation in cardiomyocytes 92 through changes in gene expression rather than gross changes in mitochondrial volume 93 or mitochondrial turnover. Moreover, our data suggest that treatment with exogenous 94 glucocorticoids may interfere with normal fetal heart maturation, possibly by down-95 regulating GR. This has implications for clinical use of antenatal corticosteroids when 96 preterm birth is considered a possibility. 97 98 INTRODUCTION 99 The dramatic increase in fetal glucocorticoid hormone concentration in late gestation is 100 essential to support the transition from intrauterine to extrauterine life (Hillman et al., 2012; 101 Rog-Zielinska et al., 2014). Administration of synthetic corticosteroids (betamethasone or 102 dexamethasone) to pregnant women at risk of preterm delivery is standard care in high 103 and middle-income countries, with the aim of maturing the fetus to reduce neonatal 104 morbidity and mortality (Kemp et al., 2016; Agnew et al., 2018). In addition to the well-

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known effects on lung maturation (Cole et al., 1995; Bird et al., 2015; Laresgoiti et al., 2016), glucocorticoids promote pro-survival adaptions in neonatal energy metabolism and in the cardiovascular system (Hillman et al., 2012). However, which of these effects are directly attributable to glucocorticoid activation of GR within tissues and which are mediated by other factors remains uncertain. Also unclear is whether antenatal administration of synthetic glucocorticoids mimics endogenous glucocorticoid action in the fetal cardiovascular system. Our previous data suggest antenatal dexamethasone treatment dysregulates cardiac function and down-regulates endogenous glucocorticoid action in the fetal heart (Agnew et al., 2019), potentially altering the normal trajectory of perinatal cardiac maturation. The normal increase in fetal glucocorticoids in late gestation supports neonatal blood pressure (Hillman et al., 2012) and is essential to structurally and functionally mature the fetal heart (Rog-Zielinska et al., 2013). In utero, our 'SMGRKO' mice, with Sm22-Cremediated GR deficiency in cardiomyocytes and vascular smooth muscle cells show impaired heart function, disrupted cardiac ultrastructure and fail to induce key genes required for cardiac contractile function, calcium handling and energy metabolism (Rog-Zielinska et al., 2013). Supporting direct effects of GR, glucocorticoid treatment of primary mouse fetal cardiomyocytes in vitro matures ultrastructure and increases contractile function, mitochondrial capacity (O<sub>2</sub> consumption rate, basally and after uncoupling of mitochondria) and markers of cardiomyocyte maturation (Rog-Zielinska et al., 2015). Similarly, in human embryonic stem cell (ESC)-derived cardiomyocytes treated with dexamethasone, contractile force is increased and systolic calcium transient decay is faster (Kosmidis et al., 2015). During the transition to a higher oxygen environment and a greater cardiac workload at birth, the cardiac preference for energy substrate switches. The fetal heart derives most of its ATP from glucose and lactate oxidation, with only a minor contribution from fatty acids. After birth, the increased demand for ATP is met primarily by oxidation of long chain fatty acids (Lopaschuk & Jaswal, 2010). This is associated with increased mitochondrial functional capacity. PGC-1α, a master transcriptional regulator of mitochondrial capacity. is expressed in the late gestation fetal heart and expression increases markedly after birth (Lehman et al., 2000). Mice with global knock-out of PGC-1 $\alpha$  show 50% mortality before weaning (Lin et al., 2004), suggesting it is important in the perinatal period. PGC-1 $\alpha$  is a glucocorticoid target gene and, in vivo in fetal heart, is induced 6 hours after glucocorticoid

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treatment (Rog-Zielinska et al., 2015). PGC-1 $\alpha$  is also induced in vitro in primary mouse fetal cardiomyocytes (Rog-Zielinska et al., 2015). Here, the GR-mediated increase in PGC-1α expression is crucial for the glucocorticoid-induced maturation of myofibril structure and increased mitochondrial O<sub>2</sub> consumption (Rog-Zielinska et al., 2015). Knock-down of PGC-1α abolished both. RNAseg analysis performed on primary mouse fetal cardiomyocytes harvested 2 hours after glucocorticoid addition in the presence of cycloheximide (to block secondary effects) identified a number of differentially expressed genes, likely to be primary targets of GR (Rog-Zielinska et al., 2015). As well as Ppargc1a (encoding PGC-1α), master regulators of mitochondrial fatty acid oxidation (Klf15, Lipin1, Cebpb, Ppara) were induced. This suggests that glucocorticoid action may promote the perinatal switch in cardiomyocytes from carbohydrate to fatty acid oxidation as the preferred substrate for ATP generation. Cellular differentiation is often associated with metabolic remodelling and the autophagic turnover of mitochondria by mitophagy (Rodger et al., 2018). Mitophagic removal of small fetal mitochondria in perinatal cardiomyocytes is reportedly a prerequisite for the formation of morphologically distinct adult mitochondria and maturation into cardiomyocytes optimised for fatty acid metabolism (Gong et al., 2015). The triggers for mitophagy in perinatal cardiomyocytes are currently unknown. *mito*-QC transgenic mice have a pHsensitive fluorescent mitochondrial signal that monitors mitophagy in vivo (McWilliams et al., 2016). These mice have revealed that mitophagy is occurring at E17.5 in the mouse fetal heart (McWilliams et al., 2016), a time co-incident with peak GR activation in heart (Rog-Zielinska et al., 2013). Furthermore, Bnip3, implicated in mitophagy, is a direct GR target gene in primary fetal cardiomyocytes (Rog-Zielinska et al., 2015), raising the possibility that glucocorticoids may be a trigger for mitophagy in perinatal cardiomyocytes. Here, we hypothesised that glucocorticoids increase fetal cardiomyocyte capacity for fatty acid oxidation. We also asked if any glucocorticoid-mediated increase in mitochondrial fatty acid oxidation capacity involves mitochondrial remodelling by mitophagy. MATERIALS AND METHODS **Animals** Experiments involving mice were approved by the University of Edinburgh Animal Welfare and Ethical Review Body and carried out in strict accordance with accepted standards of

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humane animal care under the auspices of the Animal (Scientific Procedures) Act UK 1986. Mice were maintained under controlled lighting and temperature. C57BL/6J/Ola/Hsd (C57Bl/6J) mice were purchased from Harlan, then bred in house.  $GR^{+/-}$  mice, heterozygous for a null mutation in the Nr3c1 gene encoding GR (Nr3c1gtESK92MRCHGU mice), have been previously described (Michailidou et al., 2008; Rog-Zielinska et al., 2013). GR+/- mice, congenic on the C57Bl/6J background (>12 generations) were intercrossed to give  $GR^{+/+}$ ,  $GR^{+/-}$  and  $GR^{-/-}$  fetal littermates. The morning of the day the vaginal plug was found was designated E0.5. Fetuses were collected at E17.5, hearts were dissected and rapidly frozen on dry ice. Genotyping of fetal tissue by PCR used LacZ primers for the Nr3c1gtESK92MRCHGU (5'-GAGTTGCGTGACTACCGG-3' and 5'-GTACCACAGCGGATGGTTCGG-3') and wild-type GR alleles as described (Michailidou et al., 2008), mito-QC mice, also on a C57Bl/6J background, have been described (McWilliams et al., 2016). mito-QC heterozygous fetuses were used for fetal cardiomyocyte cultures. Fetuses were collected and cardiomyocytes isolated as described below. For dexamethasone treatment, pregnant C57BI/6J females (time-mated with C57BI/6J males) were semi-randomised to experimental group (alternating groups as lifted from the cage) and injected (~0.1ml, intra-peritoneal) with dexamethasone (0.5mg/kg; Sigma-Aldrich, Poole, UK) or vehicle (5% ethanol) at either E13.5 or E16.5 and euthanised 24 hours later (E14.5 and E17.5, respectively). Fetuses were removed to ice-cold PBS. Hearts were excised and frozen on dry ice. Pregnant *GR*<sup>+/-</sup> dams were euthanised at E17.5 and fetal hearts removed and frozen as above. Neonatal C57Bl/6J mice were injected (intra-peritoneal) with dexamethasone (0.5mg/kg) or vehicle (5% ethanol) at postnatal day 1 (P1; day of birth being P0) and euthanised by decapitation 24 hours later. Hearts were removed and frozen as above. Tissues were identified by animal ID (blinding to genotype/treatment group) and stored at -80°C prior to analysis. Sheep protocols were approved by the animal ethics committee of The University of Western Australia (RA/3/100/1452). Date-mated merino ewes carrying singleton pregnancies were randomized to receive 2 injections (intra-muscular) spaced by 24 hours of either saline (control) or betamethasone acetate with betamethasone phosphate (Celestone Chronodose, Merck & Co., Inc, Kenilworth, NJ) 0.25mg/kg per injection. A third group received a single injection of betamethasone acetate (0.125mg/kg). Betamethasone acetate was a gift from Merck & Co. as a preparation of betamethasone acetate

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equivalent to that in Celestone. Merck & Co. did not participate in the design, execution, or analysis of the study. The 0.25mg/kg Celestone dose approximates the clinical dose of 12mg of betamethasone for a 50kg woman and was the same dose used for our previous studies (Kemp et al., 2018; Schmidt et al., 2019). To reduce the risk of preterm labour from antenatal corticosteroids, all animals, irrespective of subsequent treatment, received one intramuscular dose of 150mg medroxyprogesterone acetate (Depo-Provera, Pfizer, New York, NY), five days before corticosteroid treatment. No other doses of medroxyprogesterone acetate were administered, nor were other tocolytics administered. All animals were delivered at 122±1 days (term being ~147 days). Two days after their initial steroid or saline treatment, ewes received an intravenous injection of ketamine (10mg/kg) and midazolam (0.5mg/kg). A spinal injection of 3ml lignocaine (20mg/ml) was then administered and surgical delivery commenced. The lamb received an intramuscular injection of ketamine (10mg/kg) before placing a 4.5mm endotracheal tube by tracheostomy. Lambs were weighed, dried, and placed in an infant warmer (Fisher & Paykel Healthcare, New Zealand). Intermittent positive pressure ventilation was performed using Acutronic Fabian infant ventilators (Acutronic Medical System, Hirzel, Switzerland) and maintained for 30 minutes using the following settings: initial peak inspiratory pressure (PIP) of 40 cmH<sub>2</sub>O, positive end expiratory pressure (PEEP) of 5 cmH<sub>2</sub>O, respiratory rate of 50 breaths per minutes, inspiratory time of 0.5 seconds. Gas mix was 100% heated and humidified oxygen. PIP was titrated to achieve a tidal volume of 7ml/kg. Lambs were euthanised with an IV overdose of pentobarbitone sodium at 160mg/kg. Necropsy was performed immediately following a 30 minute ventilation procedure which commenced at delivery. At necropsy (within 40 minutes of delivery), left and right ventricles of the heart were dissected and samples snap frozen. **Fetal Cardiomyocyte Cultures:** Primary fetal cardiomyocyte cultures were prepared as described (Rog-Zielinska et al.. 2015). Briefly, hearts were rapidly dissected from 5-30 E14.5-E15.5 C57Bl/6J or mito-QC fetuses, placed in warm Tyrode's salt solution containing 0.1% sodium bicarbonate then rinsed in complete medium (DMEM supplemented with 100IU/ml penicillin, 100mg/ml streptomycin, 10% fetal bovine serum, 0.1% non-essential amino acids, Sigma-Aldrich, Poole, UK) before digestion at 37°C with gentle agitation for 10 minutes in 5ml enzyme buffer (PBS supplemented with 0.8% NaCl, 0.2% D-(+)-glucose, 0.02% KCl, 0.000575% NaH<sub>2</sub>PO<sub>4</sub> H<sub>2</sub>O<sub>1</sub> 0.1% NaHCO<sub>3</sub>, pH7.4) containing 0.03% type II collagenase (Worthington

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Biochemical Corp, Lakewood, New Jersey, USA), 0.125% porcine pancreatin (Sigma-Aldrich, Poole, UK). After 10 minutes, isolated cells and enzyme buffer were removed and the enzymatic reaction quenched by adding the same volume of complete medium. Fresh enzyme buffer was then added to the hearts. Approximately 8 digestions were performed on the hearts until their structure was lost. Isolated cells were centrifuged at 1000 rpm for 10 minutes at room temperature, the supernatant removed and pellets pooled. Pooled cells were centrifuged again and resuspended in 15ml isolation buffer (Ham's F12 supplemented with 100IU/ml penicillin, 100mg/ml streptomycin, 0.002% ascorbic acid, 1% fetal bovine serum, 0.1176% NaHCO<sub>3</sub>). To reduce the number of fibroblasts, cells were incubated in a tissue culture plate (ThermoFisher, UK) at 37°C, 5% CO<sub>2</sub> for 3 hours during which fibroblasts adhered to the plastic. Non-adherent cells were aspirated, centrifuged, resuspended in 1ml of complete medium and seeded at a density of 0.25 x 10<sup>6</sup>cells/ml for extra-cellular flux (ECF; Seahorse) assays, RNA analysis, or mitochondrial morphology. This protocol yields ≥98% cardiomyocytes (troponin T+ cells) (Rog-Zielinska *et al.*, 2015). Spontaneous beating of cardiomyocytes was observed within 12 hours. For ECF assays. cardiomyocytes were seeded onto 24 well gelatin (Sigma, UK) coated V7 Seahorse plates (Agilent Technologies LDA UK Ltd, Stockport, Cheshire, UK) in complete medium then treated with dexamethasone (1µM) or vehicle (0.01% ethanol) for 24 hours prior to ECF assay (below). We have previously shown that this dose of dexamethasone elicits maximal glucocorticoid responses in fetal cardiomyocytes (Rog-Zielinska et al., 2015). For RNA analysis, primary fetal cardiomyocytes were seeded in 12-well gelatin-coated tissue culture plates for 48 hours. Cells were treated with dexamethasone (1µM) or vehicle (0.01% ethanol) and lysed 6 or 24 hours later by adding 0.5ml TRIzol (Invitrogen, ThermoFisher, UK) following removal of medium. For measurement of mitochondrial morphology and mitophagy, cardiomyocytes were cultured on gelatin-coated glass chambered slides (Ibidi µ-Slide 4 well Glass bottom, Thistle Scientific LTD, Glasgow, UK) prior to staining. **Extra-cellular Flux (ECF) Assays:** ECF assays were carried out using a Seahorse XFe24 Bioanalyzer (Agilent Technologies LDA UK Ltd, Stockport, Cheshire, UK). All Seahorse reagents were purchased from Agilent Technologies LDA UK Ltd. All findings were reproduced in at least 2 experiments. For each experiment, treatment groups were randomised across the plate. Each well was assigned a number (based on the number of treatment groups) so that numbers were spread across the plate. Treatments were then randomised to numbers. On the day of the

assay, complete culture medium was gently aspirated from the cells and exchanged for Seahorse assay medium (with supplements defined below). The cells were gently rinsed 3 times with Seahorse assay medium, leaving a final volume of 525µl for the assay. During all assays, 3 measurements at 2.5 minute intervals, were recorded at baseline and after each drug addition. Normalisation of Extra-cellular Flux Assays using sulforhodamine B (SRB) Assay: ECF assays were normalised to protein measured using sulforhodamine B (SRB) dyebased protein assay (Skehan et al., 1990). Initial experiments confirmed the linearity of the SRB assay for use in quantifying primary cardiomyocyte protein levels (Supplementary Figure 1A). Following ECF assays, cells were fixed by addition of 50ul cold 50% trichloroacetic acid (Sigma-Aldrich, Poole, UK) per well and stored up to 1 week at 4°C. Cells were then washed 10 times with tap water and air-dried. 50µl SRB solution (0.4% w/v sulforhodamine B dye (Sigma-Aldrich, Poole, UK) in 1% acetic acid (Sigma-Aldrich, Poole, UK)) was added to dried cells and incubated for 30 minutes at room temperature. Cells were then washed 4 times with 1% acetic acid and air-dried. Cellbound dye was re-dissolved in 200 µl 10mM Tris pH 10.5. Absorbance was measured at a wavelength of 540nm. Glycolysis Assays: Assays to estimate the rate of glycolysis in fetal cardiomyocytes were performed in two different ways. In the first, complete culture medium was exchanged for pre-warmed Seahorse Assay medium supplemented with 10mM glucose (Sigma-Aldrich, Poole, UK), 1mM sodium pyruvate (Sigma-Aldrich, Poole, UK). After basal OCR/ECAR measurements, 2-deoxyglucose (2DG; Sigma-Aldrich, Poole, UK, 100 mM) was added to inhibit glycolysis, followed by antimycin and rotenone (AR, Sigma-Aldrich, Poole, UK,  $2\mu M$ ) to inhibit respiration. The second measure was a Glycolysis Stress Test, performed according to the manufacturer's protocol. Briefly, the cells were incubated in pre-warmed glucose-free Seahorse-XF Base medium for 1 hour in a (non-CO<sub>2</sub>) 37°C incubator prior to the assay. After 3 basal measurements, glucose (Sigma-Aldrich, Poole, UK, 10mM) was added to enable glycolysis, followed by oligomycin (Sigma-Aldrich, Poole, UK, 1.5µM) to inhibit respiratory ATP production and finally 2DG (Sigma-Aldrich, Poole, UK, 100 mM) to inhibit glycolysis.

#### **Fatty Acid Oxidation Assay:**

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To test the ability of primary fetal cardiomyocytes to utilize long chain fatty acids, cells were pre-treated with etomoxir (Sigma-Aldrich, Poole, UK) to inhibit the CPT-1 mitochondrial fatty-acid uptake transporter, prior to performing a standard Seahorse Mitochondrial Stress Test. Initial experiments with high concentrations of etomoxir (40-160μM) reduced OCR in the presence of BSA-palmitate (Supplementary Figure 1C). However, within this range (120µM), etomoxir also showed likely off-target inhibition of mitochondrial respiration (Supplementary Figure 1B). This is consistent with an emerging literature showing that etomoxir can inhibit complex I at commonly used high doses (Divakaruni et al., 2016; Yao et al., 2018). Accordingly, we used 6μM etomoxir for all further experiments, a dose which inhibits fatty acid oxidation and avoids the CPT-1independent effects associated with higher doses (Spurway et al., 1997). Briefly, culture medium was exchanged for Seahorse Assay medium supplemented with 5mM glucose (Sigma-Aldrich, Poole, UK), 0.5mM carnitine (Sigma-Aldrich, Poole, UK). Cells were pretreated with etomoxir (6µM in medium) or vehicle (medium) 15 minutes prior to the addition of BSA-Palmitate (100uM: Agilent Technologies LDA UK Ltd). The Seahorse Mitochondrial Stress test assay was started 15 minutes after the addition of BSA-Palmitate. Briefly, the test progressed as follows: basal respiration measurements, addition of oligomycin (1.5µM), addition of carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP; 1μM), addition of AR (2μM). Nonmitochondrial respiration was calculated as the minimum OCR remaining following AR treatment. The mean of the 3 baseline measurements was used for the Basal OCR. Basal respiration was calculated by subtracting non-mitochondrial respiration from Basal OCR. ATP production was calculated as the maximum change in OCR following the addition of oligomycin and maximum respiration was calculated as the maximum OCR measurement induced by FCCP corrected for non-mitochondrial respiration. Leak respiration was calculated as the average oligomycin-insensitive OCR corrected for non-mitochondrial respiration. **Measurements of Mitochondrial Morphology:** On the day of staining of primary fetal cardiomyocytes, complete culture medium was exchanged for serum-free medium. Mitotracker Deep Red (40nM; ThermoFisher, UK) was added and incubated for 30 minutes in cell culture conditions. Serum-free medium was then replaced. To image cardiomyocytes as a z stack, beating was stopped by adding 100mM nefidipine (Sigma-Aldrich, Poole, UK) immediately prior to imaging using an Andor Spinning Disk confocal microscope. The Andor Spinning disk system is based on an

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inverted Olympus IX83 microscope stand and a Yokogawa CSU-X1 spinning disk module. It is equipped with an Oko Labs environmental control chamber to maintain stable conditions for live cell imaging; 37°C, 5% CO<sub>2</sub> were used throughout. 488nm (BP525/25) and 561nm (LP568) laser lines (and emission filters) operated via an AOTF were used to acquire GFP and mCherry images, respectively. A plan super apochromat 100X 1.4NA oil immersion objective was used throughout, with z steps of 1µm taken. Images were acquired onto an Andor iXon Ultra EMCCD camera (512x512) using an EM gain of 200 with 50ms exposure. Mitochondrial volume was quantified using open-source Mitograph software (Viana et al., 2015; Harwig et al., 2018). This uses 3D reconstructions of labelled organelles to measure the morphology of individual mitochondria as well as characteristics of the mitochondrial network and provides the following outputs: mitochondrial volume, total length and average width. Post-image processing and MitoGraph analysis was performed as per the online protocols (available: http://rafelski.com/susanne/MitoGraph). Mitophagy Assay: Primary fetal cardiomyocyte cultures prepared from E14.5-15.5 mito-QC fetuses were seeded at a density of 0.25 x 10<sup>6</sup>cells/ml on gelatin-coated glass chambered slides (Ibidi μ-Slide 4 well Glass bottom, Thistle Scientific LTD, Glasgow, UK). After 48 hours, cells were treated with 1µM dexamethasone. 1mM deferiprone (DFP, an iron-chelator, used as a positive control) or vehicle (0.01% ethanol). After 24 hours dexamethasone/DFP/vehicle treated cells were imaged live and z stacks were generated 3, 8 and 45 hours after treatment, using the Andor Spinning Disk confocal live cell imaging system as above. Post-acquisition image analysis was performed blind to the treatment. For each cell, a z stack was acquired and red puncta, indicative of mitophagy, were counted. **RNA** extraction: Primary fetal cardiomyocytes seeded in 12 well plates were lysed in 500µl TRIzol (Invitrogen, ThermoFisher, UK) and aspirated into a 1.5ml tube. Chloroform (Sigma-Aldrich, Poole, UK; 100 µl) was added and samples vigorously shaken for 30 seconds. Samples were incubated for 2-3 minutes at room temperature then centrifuged 12000 rpm for 15 minutes at 4°C. The aqueous phase was transferred to a new tube containing 250 ul isopropanol (Sigma-Aldrich, Poole, UK) and centrifuged as above. The supernatant was discarded and the pellet washed with 500µl 70% ethanol twice. Samples were centrifuged

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at 10000 rpm for 10 minutes at 4°C, the supernatant removed and pellet air-dried at room temperature for 5-10 minutes before resuspension in 30µl RNAse free water. Mouse fetal hearts were individually homogenised using a stainless steel bead in 500 ul RLT buffer (RNeasy, Qiagen, Manchester, UK) and 1% β-mercaptoethanol with a TissueLyser II (Qiagen, Manchester, UK) at maximum speed for 2 minutes. 10μl Proteinase K (20mg/ml; Qiagen, Manchester, UK) and RNAse-free water were added (final volume of 900 µl). Samples were incubated at 56°C for 10 minutes, centrifuged at 12000 rpm for 5 minutes at room temperature and the supernatant transferred to a fresh tube containing 400µl 96% ethanol. The mixture was transferred to RNeasy spin tubes (Qiagen mini prep), processed according to the manufacturer's instructions and eluted in 20µl RNAse free water. This was incubated on the column for at least 5 minutes at room temperature before collection. The first eluate was reapplied to the column and incubated a further minute before the final elution. For sheep hearts, RNA extraction was similar except that samples were minced prior to lysis for 2 sessions each of 3 minutes. RNA quantity and integrity were determined using a Nanodrop (ThermoFisher, UK) spectrophotometer and gel electrophoresis, respectively. **Reverse Transcription and quantitative real time PCR:** 500ng mouse or 300ng sheep RNA was reverse transcribed using QuantiTect Reverse transcription kit (Qiagen, Manchester, UK) and qDNA wipe-out (to remove genomic DNA). according to the manufacturer's protocol. Included in each sample batch were "no template" and "no reverse transcriptase" controls. Resultant cDNA samples were stored at -20°C. Primers and probes used for qPCR are detailed in Supplementary Table 1. Assays for mouse qPCR were designed using the Roche Universal Probe Library and primers were purchased from Invitrogen (ThermoFisher, UK). A standard curve prepared from pooled cDNA samples was processed with samples on a Lightcycler 480 system (Roche Diagnostics, Burgess Hill, UK). For sheep, Tagman assays (ThermoFisher, UK) were used and performed using a 7900HT Fast Real Time PCR system (ThermoFisher, UK). Internal controls were Tbp (for mouse), and PGK1 and SDHA for sheep fetal heart; these did not differ across treatments. To normalise the spread, data were log<sub>10</sub> transformed.

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Mitochondrial DNA quantification: DNA was extracted from frozen samples using a DNeasy Blood & Tissue kit (Qiagen, Manchester, UK) according to the manufacturer's protocol. To quantify mitochondrial DNA relative to nuclear DNA, levels of mitochondrial-encoded genes (Co1, Co2, Nd2) were measured relative to an intronless nuclear-encoded gene (Cebpa) by qPCR (primer sequences in Supplementary Table 1). Statistics: Graphpad Prism 8 software was used for statistical analyses. All data are presented as mean ± standard deviation (SD). The number of biological replicates is provided in the figure legends together with the statistical tests used for analysis. All data were subject to Shapiro-Wilk normality testing prior to analysis. Parametric analyses, Student's t-tests and two-way ANOVA with *post hoc* Sidak's tests were used as stated in Figure Legends. **RESULTS** Dexamethasone increases mitochondrial respiration in primary fetal cardiomyocytes without affecting glycolysis Consistent with previous findings (Rog-Zielinska et al., 2015), without added fatty acid. basal respiration (oxygen consumption rate; OCR) prior to, and following, addition of 10mM glucose was increased 24 hours following dexamethasone treatment of primary fetal cardiomyocytes (Figure 1A, C). OCR was markedly decreased following addition of oligomycin, an inhibitor of ATP synthase (Figure 1A), suggesting high dependence on mitochondrial respiration for ATP production. As expected, addition of the glycolysis inhibitor, 2-deoxyglucose (2-DG) did not alter the OCR (Figure 1A). Glycolysis is an important energy source in early fetal cardiomyocytes (Porter et al., 2011). However, our murine primary fetal cardiomyocytes exhibited little dependence on glycolysis. The addition of 2DG did not alter the extra-cellular acidification rate (ECAR; Supplementary Figure 2). In a different approach, using glucose-deprived cells, although the addition of 10mM glucose increased ECAR, this was minimally impacted by 2DG (Figure 1B) and was unaffected by dexamethasone treatment (Figure 1E). This suggests primary fetal cardiomyocytes perform very little glycolysis, relying mainly on mitochondrial oxidation for ATP production.

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Dexamethasone increases palmitate oxidation via GR activation To investigate whether dexamethasone can increase capacity for long chain fatty acid oxidation in fetal cardiomyocytes, OCR was measured in the presence of palmitate. Long chain fatty acids are linked to carnitine and transported into the mitochondrial matrix by carnitine palmitoyltransferase-1 (CPT-1), which is inhibited by etomoxir. Following treatment of primary fetal cardiomyocytes with dexamethasone for 24 hours, in the presence of palmitate there was an increase in basal OCR (Figure 2A, B). Furthermore, in the presence of palmitate, dexamethasone-treated cardiomyocytes showed a larger change in OCR following oligomycin treatment (oligomycin-sensitive OCR; Figure 2C), indicative of increased mitochondrial ATP production. Addition of etomoxir attenuated the dexamethasone-induced increase in basal OCR and mitochondrial ATP production (oligomycin-sensitive OCR) (Figure 2A-C), suggesting that dexamethasone increased palmitate oxidation in fetal cardiomyocytes. Etomoxir itself had no effect on basal respiration or ATP production by fetal cardiomyocytes in the absence of palmitate (Supplementary Figure 1D. E). The dexamethasone-induced increase in fatty acid oxidation was dependent on GR, as pre-treatment of the cardiomyocytes with the GRantagonist, RU486 blocked the increase in basal respiration and ATP production (Figure 3A-C). Dexamethasone upregulates genes involved in long-chain fatty-acid oxidation in fetal cardiomyocytes The increase in ability to utilise palmitate as a fuel for mitochondrial respiration suggests glucocorticoids increase mitochondrial capacity for long chain fatty acid oxidation. Consistent with the rapid induction of PGC-1 $\alpha$  and other master regulators of lipid metabolism in dexamethasone-treated fetal cardiomyocytes (Rog-Zielinska et al., 2015) there was a marked induction of mRNAs encoding enzymes and transporters required for mitochondrial fatty acid oxidation in primary fetal cardiomyocytes 24 hours after treatment with dexamethasone (Figure 4). As well as the master transcriptional regulators, *Ppargc1a* and Lipin1, dexamethasone induced expression of Lcad and Mcad (encoding, respectively, long chain acyl dehydrogenase and medium chain acyl dehydrogenase), Cd36, encoding cluster of differentiation-36, also known as fatty acid translocase (a cellular importer of fatty acids), and Cpt1a and Cpt1b, encoding the alpha and beta subunits, respectively, of CPT-1 (Figure 4). Dexamethasone also increased expression of Ucp2, encoding uncoupling protein 2, an inner mitochondrial membrane protein that promotes mitochondrial fatty acid oxidation at the expense of mitochondrial catabolism of

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pyruvate (Pecqueur et al., 2008). At just 6 hours after addition of dexamethasone, although Fkbp5, a well-known glucocorticoid target was strongly up-regulated, levels of Nr3c1 mRNA, encoding GR, were down-regulated (Supplementary Figure 3), though they recovered by 24 hours (Figure 4). Pre-treatment with RU486 attenuated the dexamethasone-induced increase in *Ppargc1a*, *Lcad*, *Lipin1* and *Cd36* mRNAs (Supplementary Figure 4A-I). At E17.5, hearts of GR-/- fetal mice had reduced levels of Mcad mRNA and a trend for reduced levels of Ppargc1a mRNA (Supplementary Figure 4J). The latter is consistent with our previous finding of reduced *Ppargc1a* mRNA in hearts of E17.5 *GR*-√ fetal mice (Rog-Zielinska *et al.*, 2013). The failure to reach statistical significance here likely reflects smaller group sizes and over-night matings rather than the time-restricted matings adopted previously. Glucocorticoids do not cause mitochondrial remodelling in fetal cardiomyocytes Because there is a wave of mitophagy in vivo in the mouse fetal heart that coincides with the peak of fetal corticosterone levels (Rog-Zielinska et al., 2013; McWilliams et al., 2016), we hypothesised that glucocorticoids may stimulate the mitophagic replacement of fetal mitochondria by adult mitochondria optimised for fatty acid metabolism (Gong et al., 2015). mito-QC transgenic mice utilise a binary fluorescence system in which a ubiquitously expressed tandem mCherry-GFP tag is directed to mitochondria (McWilliams et al., 2016). Under steady-state conditions, the mitochondrial network fluoresces red and green (merged, yellow) in mito-QC mice. Upon delivery to lysosomes, the GFP fluorescence, but not that of mCherry, is quenched by the acidic microenvironment. Thus, mitochondria undergoing mitophagic removal appear as punctate mCherry-only foci. To investigate whether dexamethasone induces mitophagy, primary fetal cardiomyocytes from mito-QC mice were treated with dexamethasone, vehicle or a mitophagy-inducing agent, deferiprone (DFP) (Allen et al., 2013). As expected, DFP stimulated a robust mitophagy response (as visualised by an increase in the number of mCherry-only positive puncta) over the 45 hour time course (Figure 5A). In contrast, no significant increase in mitophagy could be seen in cardiomyocytes following a similar time course of dexamethasone treatment (Figure 5B). Thus, under these conditions, dexamethasone is not a potent inducer of mitophagy. Glucocorticoids have been associated with changes in mitochondrial number and function (Weber et al., 2002; Du et al., 2009; Lapp et al., 2018). Accordingly, we next investigated whether the dexamethasone-induced increase in fatty acid oxidation was associated with

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an increase in mitochondrial volume and/or number. Following dexamethasone treatment of primary fetal cardiomyocytes for 24 hours, mitochondria were labelled with Mitotracker Deep Red CM and the mitochondrial network imaged (Figure 5C, D). There were no differences in mitochondrial volume, length or width between dexamethasone and vehicle treated cardiomyocytes (Figure 5E-G). Similarly, measurements of the GFP-fluorescent mitochondrial network in fetal cardiomyocytes from mito-QC mice showed no differences in mitochondrial morphology as a result of dexamethasone treatment (Supplementary Figure 5A-C). Furthermore, mitochondrial DNA content, an indirect measurement of mitochondrial number, did not differ between hearts of E17.5 GR-/- mice and their control GR+/+ littermates (Supplementary Figure D-F). Thus, the glucocorticoid-mediated increase in fatty acid oxidation capacity most likely occurs independently of any change in mitochondrial number or morphology. In vivo, dexamethasone-induced changes in fatty acid oxidation genes in mouse hearts are developmental stage-dependent To investigate whether glucocorticoid administration in vivo can similarly induce cardiac fatty acid oxidation capacity, dexamethasone or vehicle was administered to pregnant dams at E13.5 or E16.5 or to neonatal mice at postnatal day (P)1. Hearts were examined 24 hours after injection. E14.5 fetal hearts appeared glucocorticoid-resistant: dexamethasone had no significant effect on any of the mRNAs examined, including the glucocorticoid target, Fkbp5, and Nr3c1 encoding GR itself (Figure 6A). However, at E17.5, dexamethasone downregulated cardiac Nr3c1 mRNA levels (Figure 6B). At the same time, levels of *Ppargc1a* and mRNA encoding enzymes and transporters for fatty acid oxidation (Mcad, Lcad, Lipin1, Cd36, Cpt1a, CPT1b) were strongly downregulated, despite a trend for a modest increase in Fkbp5 mRNA (Figure 6B). In complete contrast, at P2. Fkbp5 was strongly induced by dexamethasone, as was Ppargc1a and the fatty acid oxidation genes: Mcad, Lcad, Cd36, Cpt1b (Figure 6C). Moreover, levels of Nr3c1 mRNA were unchanged (Figure 6C). Thus, the effect of glucocorticoids upon cardiac capacity for fatty acid oxidation reflect the effect upon GR expression itself and its key target gene, Ppargc1a. To explore whether this regulation extends to a more translationally relevant model, we measured cardiac mRNA encoding GR and PGC-1 $\alpha$  in a sheep model of preterm birth following antenatal corticosteroid administration that mimics current clinical practice. Celestone is a mix of betamethasone phosphate and betamethasone acetate that is widely used as an antenatal corticosteroid in the USA, Europe (though not the UK), Australia and New Zealand. Preterm lambs delivered at 127

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days (term being ~147 days) 48 hours after initiating a course of Celestone (2 doses, administered 24 hours apart) showed reduced expression of NR3C1 mRNA in both left and right ventricles of the heart (Figure 7A, B). Administration of a single dose of betamethasone acetate (equivalent to just the betamethasone acetate component of Celestone) 24 hours before delivery also reduced NR3C1 mRNA levels, though this did not achieve significance in the right ventricle (Figure 7A, B). Levels of PPARGC1 mRNA were reduced in the right ventricle following Celestone administration, with a more modest effect (p>0.05) in the left ventricle. Betamethasone acetate alone caused a similar though non-significant reduction in PPARGC1 mRNA levels in both ventricles (Figure 7C, D). This suggests that antenatal corticosteroid administration may interfere with the normal maturation of the mid to late gestation heart by down-regulating GR. **DISCUSSION** Here, we find that mouse fetal cardiomyocytes use mainly mitochondrial metabolism to generate ATP when glucose is provided as substrate, with little reliance on glycolysis. This supports the view that metabolism has switched from anaerobic glycolysis to aerobic mitochondrial respiration by the end of the embryonic period at E14.5 in mice (reviewed (Porter et al., 2011)). Although glucocorticoids increase basal mitochondrial respiration (confirming previous findings (Rog-Zielinska et al., 2015)), they have no effect on glycolysis, ruling out a glucocorticoid-promoted switch from glycolysis to oxidative metabolism. Our data do not support a glucocorticoid-mediated increase in mitochondrial number or change in morphology to account for the increase in basal respiration. Instead, our data suggest glucocorticoid action in the fetal heart promotes mitochondrial ATP generating capacity, in line with their maturational effects. As well as increasing basal mitochondrial respiration with carbohydrate substrates. glucocorticoid treatment of fetal cardiomyocytes in vitro increases fatty acid oxidation. It has been suggested that mitophagy is required to replace the mitochondrial network in perinatal cardiomyocytes with mitochondria optimised for fatty acid oxidation (Gong et al., 2015). However, our data suggest that an increase in fatty acid oxidation in perinatal cardiomyocytes can occur without substantial mitophagy. Moreover, they are consistent with the notion that mitochondrial remodelling occurs in cardiomyocytes prior to our cell isolations at ~E15. Mitochondrial phenotype in the embryonic heart changes considerably between E9.5 and E13.5, compatible with mitochondrial remodelling by mitophagy being required for the switch in cardiomyocyte reliance from anaerobic glycolysis to aerobic

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respiration. By E13.5, the network is interconnected and spans the cell, more closely resembling that in late fetal cardiomyocytes (Porter et al., 2011). Our conclusions differ from a recent report that suggested that dexamethasone promotes mitophagy in mouse embryonic-stem cell-derived cardiomyocytes through Parkin (Zhou et al., 2020). However, in those experiments, detection of lysosomes (using lysotracker) was only possible in dexamethasone-treated cells (Zhou et al., 2020) making interpretation of the mitophagy findings difficult. Nevertheless, dexamethasone did not affect mitochondrial morphology in mouse embryonic-stem cell-derived cardiomyocytes (Zhou et al., 2020), consistent with our findings here. Our data reported here clearly show that although glucocorticoid treatment in primary mouse fetal cardiomyocytes increases fatty acid oxidation, it does not induce widespread mitophagy. Fatty acid oxidation dramatically increases around the end of the first postnatal week in mouse heart, and we found a marked induction of the pathway following glucocorticoid administration in vivo in neonatal mice (Lopaschuk & Jaswal, 2010). However, in preparation for postnatal life, fatty acid oxidation has already begun to occur in the late gestation fetal heart (Lopaschuk & Jaswal, 2010; Porter et al., 2011). In sheep, there is an increase in cardiac expression of genes related to fatty acid oxidation between late gestation and term that continues after birth (Richards et al., 2015). In silico transcription factor analysis suggests this is, at least in part, GR-mediated (Richards et al., 2015). The lower level of Mcad mRNA in hearts of GR knockout fetuses at E17.5, a stage when endogenous glucocorticoid levels have increased, supports a role for GR in the late gestation increase in cardiac expression of genes required for fatty acid oxidation. Fatty acid oxidation genes themselves are not primary targets of GR in mouse fetal cardiomyocytes (Rog-Zielinska et al., 2015) and are likely indirectly regulated by master regulators of fatty acid oxidation including CEBPB. PPAR $\alpha$  and PGC-1 $\alpha$ , which are primary GR targets (Rog-Zielinska et al., 2015). Thus, endogenous glucocorticoid action, via GR, may contribute to the normal rise in fatty acid oxidation capability as the fetus approaches term. This is likely to be important to meet the increase in cardiac energy demand after birth, consistent with the ergogenic effects of glucocorticoids (Addison, 1855; Morrison-Nozik et al., 2015) and the vital role of the late gestation increase in glucocorticoids to prepare for life after birth. Crucially, our data illustrate how exogenous glucocorticoid may interfere with the normal maturation of energy metabolism in the fetal heart. Although dexamethasone increases

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expression of fatty acid oxidation genes in neonatal mice, 24 hours after administration of dexamethasone at E16.5 (the peak of endogenous fetal glucocorticoid levels) the expression of genes related to fatty acid oxidation actually decreases in the fetal heart. The association with reduced Nr3c1 mRNA, encoding GR itself, suggests the decrease in the mitochondrial fatty acid oxidation pathway reflects down-regulation of glucocorticoid signalling per se in the fetal heart following dexamethasone treatment. We recently reported similar down-regulation of GR expression in fetal heart as well as reduced endogenous fetal corticosterone levels following dexamethasone treatment (via drinking water) between E12.5-E15.5 (Agnew et al., 2019). This was associated with a transient alteration in fetal diastolic heart function (Agnew et al., 2019). Whilst dexamethasone also down-regulates Nr3c1 mRNA in fetal cardiomyocytes in vitro, this is transient with recovery of Nr3c1 mRNA expression by 24 hours. In the neonatal heart, if Nr3c1 mRNA is transiently downregulated by dexamethasone, it has recovered within 24 hours. Dynamic and differential auto-regulation of GR, previously described for adult tissues (Kalinyak et al., 1987; Spencer et al., 1991; Freeman et al., 2004), may contribute to the complex and context dependent effects of perinatal glucocorticoid administration. Previous studies have examined the effect of antenatal glucocorticoids in rodents and reported contradictory findings (reviewed, (Rog-Zielinska et al., 2014)). Our data illustrate that exogenous glucocorticoids can potentially interfere with normal heart maturation. Indeed, maturation of the rat heart is delayed after prenatal treatment with dexamethasone (Torres et al., 1997). Timing may be critical. Although antenatal dexamethasone increased ATP content in the neonatal rat heart, the same treatment did not increase cardiac ATP content prior to birth, despite an increase in creatine kinase expression (Mizuno et al., 2010). GR expression was not examined in that study. Investigations in sheep have also highlighted adverse effects of antenatal glucocorticoid exposure on cardiac energy metabolism. Maternal hypercortisolaemia reduces fetal cardiac mitochondrial number and oxidative metabolism at term, associated with fetal ECG abnormalities, an inability to maintain fetal aortic pressure and heart rate during labour and a dramatic increase in perinatal death (Antolic et al., 2018). Again, whether GR expression was affected was not reported. Differential mRNA stability may explain some of the complex effects of glucocorticoids on downstream genes. PGC-1 $\alpha$  is essential for efficient and maximal fatty acid oxidation and ATP production in cardiomyocytes (Arany et al., 2005; Lehman et al., 2008). Ppargc1a mRNA has a short half-life, being less than 30 minutes in rat skeletal muscle extracts. This is further decreased with chronic muscle stimulation (Lai et al., 2010). Consistent with

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a short half-life, we have previously shown that blocking new protein synthesis with cycloheximide increases Ppargc1a mRNA levels in fetal cardiomyocytes (Rog-Zielinska et al., 2015), suggesting it is actively degraded. Glucocorticoid treatment rapidly increases levels of Ppargc1a mRNA in fetal heart in vivo and in fetal cardiomyocytes in vitro (Rog-Zielinska et al., 2015). Moreover, dexamethasone "super-induces" Ppargc1a in the presence of cycloheximide (Rog-Zielinska et al., 2015), again consistent with a rapid turnover of Ppargc1a mRNA. A very short half-life and a need for activated GR to continually enhance transcription of *Ppargc1a* mRNA may explain the close association between Nr3c1 and Ppargc1a mRNA in both fetal heart and mouse primary cardiomyocytes that we saw here. By contrast, in all of our experiments, with the exception of fetal heart at E14.5 (which appeared glucocorticoid resistant), Fkbp5 mRNA was upregulated by dexamethasone. Even in the E17.5 heart, there was a strong trend for increased Fkbp5 mRNA following dexamethasone, despite downregulation of GR and the fatty acid oxidation pathway at this time. Just 6 hours after addition of dexamethasone to fetal cardiomyocytes in vitro. Fkbp5 mRNA was already elevated, despite downregulation of GR. Thus, Fkbp5 mRNA appears a stable readout of early GR activation whereas Ppargc1a mRNA correlates with Nr3c1 mRNA at any particular time. Plausibly, antentatal corticosteroids may disrupt the normal maturation of energy metabolism in the human fetal heart, in part by down-regulating PGC-1α. In our sheep model that closely mirrors clinical practice, both NR3C1 and PPARGC1A were downregulated in fetal heart by antenatal corticosteroids. Whether this is transient, and both later recover to control levels merits further investigation. Nevertheless, this suggests that clinical administration of antenatal corticosteroids in mid- and late gestation may interfere with normal heart maturation. The E13.5 fetal heart appears resistant to dexamethasone. The reason for this is unclear, but it has implications for clinical practice. In humans, the reduction in fetal heart rate variability (a clinical marker of fetal hypoxia/poor outcomes) 2 to 3 days after maternal administration of glucocorticoids is greater in fetuses >30 weeks gestation than those <30 weeks (Mulder et al., 2009), consistent with gestation-stage dependent effects of antenatal glucocorticoids. The greater sensitivity to the haemodynamic effects of glucocorticoids coincides with an increase in fetal cortisol synthesis, from ~30 weeks gestation (Hillman et al., 2012). In mice, adrenal steroidogenesis initiates at E14.5 (Michelsohn & Anderson, 1992). This raises the possibility that the fetus is glucocorticoid resistant prior to the gestational increase in fetal glucocorticoid levels. Whether this is the

case or not merits future investigation, given wide-spread expression of GR in the mouse fetus prior to E14.5. It is interesting to note that although dexamethasone increases calcium handling and contraction force in human embryonic stem cell-derived cardiomyocytes (Kosmidis *et al.*, 2015), human induced pluripotent stem cell-derived cardiomyocytes (roughly corresponding to first trimester human fetal cardiomyocytes (van den Berg *et al.*, 2015)) do not respond to dexamethasone alone, possibly because they lack a competence factor (Birket *et al.*, 2015). The acquisition of competence to respond to glucocorticoids as well as the auto-regulation of GR itself may therefore be developmentally regulated and may differ between cell types. Understanding how this contributes to the maturational effects of glucocorticoids upon fetal organs and tissues will be vital to optimise antenatal corticosteroid therapy in the future, to limit possible harm and maximise benefit.

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#### DATA AVAILABILITY

- The data that support the findings of this study are available from the first and/or
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#### **COMPETING INTERESTS**

None of the authors have a competing financial or other conflict of interest.

#### **AUTHOR CONTRIBUTIONS**

- Conceptualisation JRI, EAR-Z, KEC; data curation JRI, KEC, SJS; data analysis JRI,
- LH, RNC, NMM; Funding KEC, IGG, SJS, MWK, NMM, CW; Investigation JRI, RNC, J-
- 729 FZ, CB, HU, EAR-Z, EP, LH, CN, SJS, MWK; Methodology JRI, RNC, J-FZ, CB, EAR-Z,
- 730 EJA, MWK; Project administration JRI, KEC; Resources IGG, MWK, NMM, KEC;
- Supervision JRI, KEC; Validation JRI, KEC; Writing original draft JRI, MWK, KEC;
- 732 Writing review and editing JRI, RNC, CB, EAR-Z, EJA, MWK, SJS, IGG, KEC. All

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Arany Z, He H, Lin J, Hoyer K, Handschin C, Toka O, Ahmad F, Matsui T, Chin S, Wu PH, Rybkin, II, Shelton JM, Manieri M, Cinti S, Schoen FJ, Bassel-Duby R, Rosenzweig A, Ingwall JS & Spiegelman BM. (2005). Transcriptional coactivator PGC-1 alpha controls the energy state and contractile function of cardiac muscle. Cell Metab 1, 259-271. Bird AD, McDougall AR, Seow B, Hooper SB & Cole TJ. (2015). Glucocorticoid regulation of lung development: Lessons learned from conditional GR knockout mice. Mol Endocrinol 29, 158-171. Birket MJ, Ribeiro MC, Kosmidis G, Ward D, Leitoguinho AR, van de Pol V, Dambrot C, Devalla HD, Davis RP, Mastroberardino PG, Atsma DE, Passier R & Mummery CL. (2015). Contractile defect caused by mutation in MYBPC3 revealed under conditions optimized for human PSC-cardiomyocyte function. Cell Reports 13, 733-745. Cole T, Blendy JA, Monaghan AP, Kriegelstein K, Schmid W, Fantuzzi G, Hummler E, Unsicker K & Schütz G. (1995). Targeted disruption of the glucocorticoid receptor blocks adrenergic chromaffin cell development and severely retards lung maturation. Genes Dev 9, 1608-1621. Divakaruni AS, Rogers GW, Andreyev AY & Murphy AN. (2016). The CPT inhibitor etomoxir has an off-target effect on the adenine nucleotide translocase and respiratory complex I. Biochem Biophys Acta; Bioenergetics 1857, e118. Du J, Wang Y, Hunter R, Wei Y, Blumenthal R, Falke C, Khairova R, Zhou R, Yuan P, Machado-Vieira R, McEwen BS & Manji HK. (2009). Dynamic regulation of mitochondrial function by glucocorticoids. Proc Natl Acad Sci U S A 106, 3543-3548. Freeman AI, Munn HL, Lyons V, Dammermann A, Seckl JR & Chapman KE. (2004). Glucocorticoid down-regulation of rat glucocorticoid receptor does not involve differential promoter regulation. *J Endocrinol* **183**, 365-374.

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Gong G, Song M, Csordas G, Kelly DP, Matkovich SJ & Dorn GW. (2015). Parkinmediated mitophagy directs perinatal cardiac metabolic maturation in mice. Science 350, aad2459. Harwig MC, Viana MP, Egner JM, Harwig JJ, Widlansky ME, Rafelski SM & Hill RB. (2018). Methods for imaging mammalian mitochondrial morphology: A prospective on MitoGraph. Anal Biochem 552, 81-99. Hillman NH, Kallapur SG & Jobe AH. (2012). Physiology of transition from intrauterine to extrauterine life. Clin Perinatol 39, 769-783. Kalinyak JE, Dorin RI, Hoffman AR & Perlman AJ. (1987). Tissue-specific regulation of glucocorticoid receptor mRNA by dexamethasone. J Biol Chem 262, 10441-10444. Kemp MW, Newnham JP, Challis JG, Jobe AH & Stock SJ. (2016). The clinical use of corticosteroids in pregnancy. Human Reproduction Update 22, 240-259. Kemp MW, Saito M, Usuda H, Watanabe S, Sato S, Hanita T, Kumagai Y, Molloy TJ, Clarke M, Eddershaw PJ, Musk GC, Schmidt A, Ireland D, Furfaro L, Payne MS, Newnham JP & Jobe AH. (2018). The efficacy of antenatal steroid therapy is dependent on the duration of low-concentration fetal exposure: evidence from a sheep model of pregnancy. Am J Obstet Gynecol 219, 301.e301-301.e316. Kosmidis G, Bellin M, Ribeiro MC, van Meer B, Ward-van Oostwaard D, Passier R, Tertoolen LG, Mummery CL & Casini S. (2015). Altered calcium handling and increased contraction force in human embryonic stem cell derived cardiomyocytes following short term dexamethasone exposure. Biochem Biophys Res Commun **467**, 998-1005. Lai RYJ, Ljubicic V, D'Souza D & Hood DA. (2010). Effect of chronic contractile activity on mRNA stability in skeletal muscle. Am J Physiol Cell Physiol 299, C155-C163. Lapp HE, Bartlett AA & Hunter R. (2018). Stress and glucocorticoid receptor regulation of mitochondrial gene expression. J Mol Endocrinol 62, R121-128.

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Laresgoiti U, Nikolic MZ, Rao C, Brady JL, Richardson RV, Batchen EJ, Chapman KE & Rawlins EL. (2016). Lung epithelial tip progenitors integrate Glucocorticoid and STAT3-mediated signals to control progeny fate. Development 143, 3686-3699. Lehman JJ, Barger PM, Kovacs A, Saffitz JE, Medeiros DM & Kelly DP. (2000). Peroxisome proliferator-activated receptor gamma coactivator-1 promotes cardiac mitochondrial biogenesis. J Clin Invest 106, 847-856. Lehman JJ, Boudina S, Banke NH, Sambandam N, Han X, Young DM, Leone TC, Gross RW, Lewandowski ED, Abel ED & Kelly DP. (2008). The transcriptional coactivator PGC-1alpha is essential for maximal and efficient cardiac mitochondrial fatty acid oxidation and lipid homeostasis. Am J Physiol Heart Circ Physiol 295, H185-196. Lin J, Wu PH, Tarr PT, Lindenberg KS, St-Pierre J, Zhang CY, Mootha VK, Jager S, Vianna CR. Reznick RM. Cui L. Manieri M. Donovan MX. Wu Z. Cooper MP. Fan MC, Rohas LM, Zavacki AM, Cinti S, Shulman GI, Lowell BB, Krainc D & Spiegelman BM. (2004). Defects in adaptive energy metabolism with CNS-linked hyperactivity in PGC-1alpha null mice. Cell 119, 121-135. Lopaschuk GD & Jaswal JS. (2010). Energy metabolic phenotype of the cardiomyocyte during development, differentiation, and postnatal maturation. J Cardiovasc Pharmacol 56, 130-140. McWilliams TG, Prescott AR, Allen GF, Tamjar J, Munson MJ, Thomson C, Muqit MM & Ganley IG. (2016). mito-QC illuminates mitophagy and mitochondrial architecture in vivo. J Cell Biol 214, 333-345. Michailidou Z, Carter RN, Marshall E, Sutherland HG, Brownstein DG, Owen E, Cockett K, Kelly V, Ramage L, Al-Dujaili EA, Ross M, Maraki I, Newton K, Holmes MC, Seckl JR, Morton NM, Kenyon CJ & Chapman KE. (2008). Glucocorticoid receptor haploinsufficiency causes hypertension and attenuates hypothalamic-pituitaryadrenal axis and blood pressure adaptions to high-fat diet. FASEB J 22, 3896-3907.

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Michelsohn AM & Anderson DJ. (1992). Changes in Competence Determine the Timing of 2 Sequential Glucocorticoid Effects On Sympathoadrenal Progenitors. Neuron 8, 589-604. Mizuno M, Takeba Y, Matsumoto N, Tsuzuki Y, Asoh K, Takagi M, Kobayashi S & Yamamoto H. (2010). Antenatal glucocorticoid therapy accelerates ATP production with creatine kinase increase in the growth-enhanced fetal rat heart. Circ J 74, 171-180. Morrison-Nozik A, Anand P, Zhu H, Duan Q, Sabeh M, Prosdocimo DA, Lemieux ME, Nordsborg N, Russell AP, MacRae CA, Gerber AN, Jain MK & Haldar SM. (2015). Glucocorticoids enhance muscle endurance and ameliorate Duchenne muscular dystrophy through a defined metabolic program. Proc Natl Acad Sci U S A 112, E6780-6789. Mulder EJ, de Heus R & Visser GH. (2009). Antenatal corticosteroid therapy: short-term effects on fetal behaviour and haemodynamics. Semin Fetal Neonatal Med 14, 151-156. Pecqueur C, Bui T, Gelly C, Hauchard J, Barbot C, Bouillaud F, Ricquier D, Miroux B & Thompson CB. (2008). Uncoupling protein-2 controls proliferation by promoting fatty acid oxidation and limiting glycolysis-derived pyruvate utilization. FASEB J 22, 9-18. Porter GA, Jr., Hom J, Hoffman D, Quintanilla R, de Mesy Bentley K & Sheu SS. (2011). Bioenergetics, mitochondria, and cardiac myocyte differentiation. Prog Pediatr Cardiol 31, 75-81. Richards EM, Rabaglino MB, Antolic A, Wood CE & Keller-Wood M. (2015). Patterns of gene expression in the sheep heart during the perinatal period revealed by transcriptomic modeling. *Physiol Genomics* **47**, 407-419. Rodger CE, McWilliams TG & Ganley IG. (2018). Mammalian mitophagy - from in vitro molecules to in vivo models. FEBS J 285, 1185-1202.

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Rog-Zielinska EA, Craig MA, Manning JR, Richardson RV, Gowans GJ, Dunbar DR, Gharbi K, Kenyon CJ, Holmes MC, Hardie DG, Smith GL & Chapman KE. (2015). Glucocorticoids promote structural and functional maturation of foetal cardiomyocytes: a role for PGC-1alpha. Cell Death Differ 22, 1106-1116. Rog-Zielinska EA, Richardson RV, Denvir MA & Chapman KE. (2014). Glucocorticoids and foetal heart maturation; implications for prematurity and foetal programming. J Mol Endocrinology 52, R125-135. Rog-Zielinska EA, Thomson A, Kenyon CJ, Brownstein DG, Moran CM, Szumska D, Michailidou Z, Richardson J, Owen E, Watt A, Morrison H, Forrester LM, Bhattacharya S, Holmes MC & Chapman KE. (2013). Glucocorticoid receptor is required for fetal heart maturation. Hum Mol Genet 22, 3269-3282. Schmidt AF, Jobe AH, Kannan PS, Bridges JP, Newnham JP, Saito M, Usuda H. Kumagai Y, Fee EL, Clarke M & Kemp MW. (2019). Oral antenatal corticosteroids evaluated in fetal sheep. Pediatric Res 86, 589-594. Skehan P, Storeng R, Scudiero D, Monks A, McMahon J, Vistica D, Warren JT, Bokesch H, Kenney S & Boyd MR. (1990). New colorimetric cytotoxicity assay for anticancer-drug screening. J Natl Cancer Inst 82, 1107-1112. Spencer RL, Miller AH, Stein M & McEwen BS. (1991). Corticosterone regulation of type-I and type-II adrenal steroid receptors in brain, pituitary, and immune tissue. Brain Res 549, 236-246. Spurway TD, Pogson CI, Sherratt HS & Agius L. (1997). Etomoxir, sodium 2-[6-(4chlorophenoxy)hexyl] oxirane-2-carboxylate, inhibits triacylglycerol depletion in hepatocytes and lipolysis in adipocytes. FEBS Lett 404, 111-114. Torres A, Belser WW, 3rd, Umeda PK & Tucker D. (1997). Indicators of delayed maturation of rat heart treated prenatally with dexamethasone. Pediatr Res 42, 139-144.

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van den Berg CW, Okawa S, Chuva de Sousa Lopes SM, van Iperen L, Passier R, Braam SR, Tertoolen LG, Del Sol A, Davis RP & Mummery CL. (2015). Transcriptome of human foetal heart compared with cardiomyocytes from pluripotent stem cells. Development 142, 3231-3238. Viana MP, Lim S & Rafelski SM. (2015). Quantifying mitochondrial content in living cells. Methods Cell Biol 125, 77-93. Weber K, Bruck P, Mikes Z, Kupper JH, Klingenspor M & Wiesner RJ. (2002). Glucocorticoid hormone stimulates mitochondrial biogenesis specifically in skeletal muscle. Endocrinology 143, 177-184. Yao CH, Liu GY, Wang R, Moon SH, Gross RW & Patti GJ. (2018). Identifying off-target effects of etomoxir reveals that carnitine palmitoyltransferase I is essential for cancer cell proliferation independent of beta-oxidation. PLoS Biol 16, e2003782. Zhou R, Li J, Zhang L, Cheng Y, Yan J, Sun Y, Wang J & Jiang H. (2020). Role of Parkinmediated mitophagy in glucocorticoid-induced cardiomyocyte maturation. Life Sci **255**, 117817. FIGURE LEGENDS Figure 1. Dexamethasone increased basal respiration in primary fetal cardiomyocytes Primary fetal cardiomyocytes were prepared by digesting pooled E14.5-15.5 C57BL/6J fetal hearts, then cultured for 48 hours in complete DMEM culture medium. Cardiomyocytes were treated with dexamethasone (Dex, 1µM, red) or vehicle (Veh, black). After 24 hours, medium was exchanged for Seahorse base medium and cardiomyocyte metabolism was analyzed by extra-cellular flux assay. After 3 basal measurements, glucose (10mM) was added, followed by oligomycin (Oligo, 1.5μM) and 2deoxyglucose (2DG, 100mM). Oxygen consumption rate (OCR, A) and extra-cellular acidification rate (ECAR, B) were measured three times over 7.5 minutes following each addition. (C) Basal OCR was calculated as the mean of the 3 basal measurements. (D) ATP production was estimated as the maximum change in OCR following the addition of oligomycin (oligomycin-sensitive OCR). (E) Glycolysis (2DG-sensitive ECAR) was

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measured as the maximum change in ECAR following addition of 2DG. Representative data from an experiment using cardiomyocytes pooled from tens of fetuses across 9-10 wells: data are mean ± SD, ns=not significant, \*\*p<0.01 \*p<0.05 (t-tests) (B), or two-way ANOVA followed by *post hoc* Sidak's tests (A). Figure 2. Dexamethasone increases fatty acid oxidation Primary fetal cardiomyocytes, prepared by digesting pooled E14.5-15.5 C57BL/6J fetal hearts, were cultured for 48 hours in complete DMEM culture medium then treated with dexamethasone (Dex, 1µM, red) or vehicle (Veh, black). After 24 hours, medium was exchanged for Seahorse assay medium supplemented with 5mM glucose, 1mM pyruvate and 0.5mM carnitine. Cells were treated with etomoxir (Eto, 6µM) or vehicle (Control) 15 minutes prior to the addition of BSA-Palmitate (100µM). After a further 15 minutes incubation, cardiomyocyte metabolism was analyzed by extra-cellular flux assay. After 3 basal measurements, oligomycin was added (Oligo, 1.5μM) followed by carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP, 1μM) then antimycin and rotenone (AR, 2μM). Oxygen consumption rate (OCR, A) was measured three times over 7.5 minutes following each drug addition. Non-mitochondrial respiration was estimated as the mean OCR remaining after AR addition. (B) Basal respiration was estimated as the mean of the 3 basal OCR measurements corrected for non-mitochondrial respiration. (C) ATP production was estimated as the maximum change in OCR following the addition of oligomycin. (D) Maximum respiration was estimated as the maximum OCR (following FCCP) corrected for non-mitochondrial respiration. (E) Leak respiration was estimated as the mean oligomycin-insensitive OCR corrected for non-mitochondrial respiration. Representative data are from an experiment performed on cardiomyocytes pooled from several fetuses across 4-5 wells, mean ± SD, ns=not significant, \*p<0.05, \*\*\*p<0.001 for comparisons between Veh/Dex, # p<0.05 for comparisons between control and Eto by two-way ANOVA followed by post hoc Sidak's tests. Three samples were excluded due to a technical failure (leakage of AR from the port during basal measurements). Figure 3. GR mediates the dexamethasone-induced increase in fatty acid oxidation Primary fetal cardiomyocytes, prepared by digesting pooled E14.5-E15.5 C57BL/6J fetal hearts were cultured for 48 hours before treatment with RU486 (1µM) or vehicle (control) 30 minutes prior to addition of dexamethasone (Dex. 1µM, red) or vehicle (Veh. black). After 24 hours, medium was exchanged for Seahorse assay medium supplemented with 5mM glucose, 1mM pyruvate and 0.5mM carnitine. Cells were treated with etomoxir (Eto,

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6μM) or vehicle (Control) 15 minutes prior to addition of BSA-Palmitate (100μM). After 15 minutes incubation with BSA-Palmitate, cardiomyocytes were subjected to extra-cellular flux assay. After 3 basal measurements, oligomycin was added (Oligo, 1.5µM) followed by carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP, 1µM) and antimycin and rotenone (AR, 2µM). Oxygen consumption rate (OCR, A) was measured three times over 7.5 minutes following each drug addition. Non-mitochondrial respiration was estimated as the average OCR remaining after AR addition. (B) Basal respiration was estimated as the mean of the 3 basal OCR measurements corrected for non-mitochondrial respiration. (C) ATP production was estimated as the maximum change in OCR following addition of oligomycin. (D) Maximum respiration was estimated as the maximum OCR measurement corrected for non-mitochondrial respiration. (E) Leak respiration was estimated as the average oligomycin-insensitive OCR corrected for non-mitochondrial respiration. Data are from cardiomyocytes pooled from tens of fetuses (a total of 12 wells across 3 different pools) and are mean ± SD, ns=not significant, \*\*p<0.01, \*\*\*\*p<0.001 (two-way ANOVA with post hoc Sidak's tests). Figure 4. Dexamethasone increases the expression of genes involved in mitochondrial fatty acid oxidation Primary fetal cardiomyocytes, prepared by digesting pooled E14.5-E15.5 C57BL/6J fetal hearts were cultured for 48 hours before treatment with dexamethasone (Dex. 1µM, red) or vehicle (Veh, black) for 24 hours. Cardiomyocyes were lysed in TRIzol and RNA isolated for analysis by qRT-PCR relative to Tbp, used as internal control. Data are from n=4 independent pools of cardiomyocytes, prepared on different days. Data are mean ± SD and were analysed by two-way ANOVA followed by post hoc Sidak's tests; \*p<0.05, \*\*p<0.01, \*\*\*p<0.001, \*\*\*\*p<0.0001. Figure 5. Dexamethasone does not alter mitochondrial morphology or induce mitophagy in primary fetal cardiomyocytes Primary cardiomyocytes were isolated from E14.5-E15.5 mito-QC (A, B) or C57Bl/6J (C-G) pooled fetal hearts, then cultured for 48 hours. (A, B) Cardiomyocytes were then treated with dexamethasone (Dex. 1µM), vehicle (Veh) or a mitophagy inducing agent. deferiprone (DFP, 1mM) and imaged live, 3, 8 and 45 hours later. z stacks of individual cells were obtained using a spinning disk confocal microscope (100X magnification). (A) Maximum z projection images are presented: white arrows indicate examples of puncta in magnified panels. Scale bars=10µm and 5µm for main and high magnification panels,

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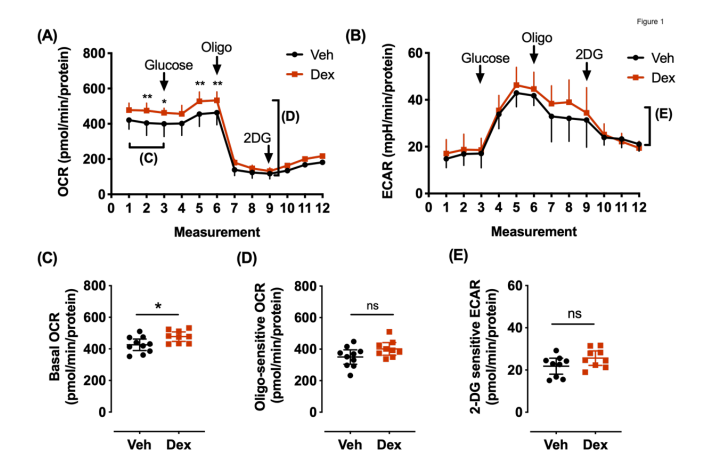
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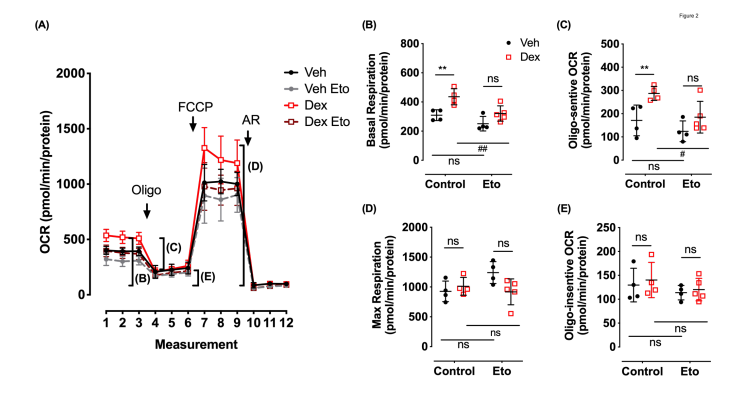
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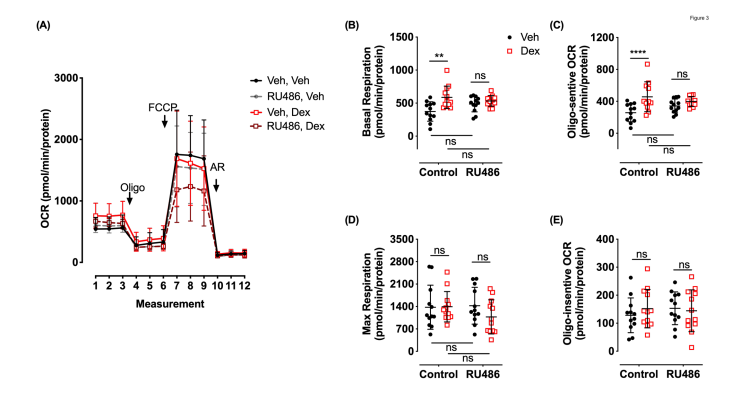
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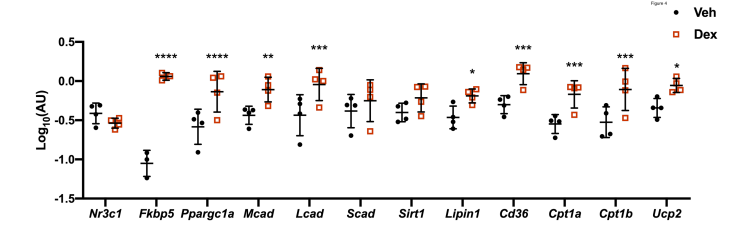
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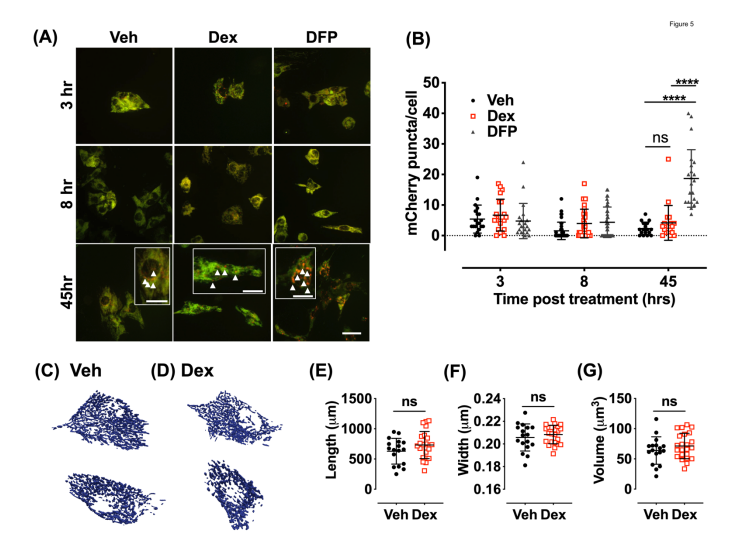
respectively. Puncta were counted manually through z stacks (B). Data are mean ± SD, n=17-23 individual cells. \*\*\*\*p<0.0001, ns=not significant, two-way ANOVA with post hoc Sidak's tests. (C-G) Cardiomyocytes were cultured for 48 hours then treated with dexamethasone (Dex. 1µM) or vehicle (Veh) for 24 hours. Mitochondria were labeled with MitoTracker Deep red CM and cardiomyocyte beating stopped with nefidipine (100mM). z stacks of individual cells were obtained using a spinning disk confocal microscope (100X magnification). Mitochondrial morphology was assessed using MitoGraph software (23, 24) and 3D renderings are shown for vehicle (C) and dexamethasone (D) treated cardiomyocytes. Parameters included (E) length. (F) width and (G) total volume. Data are mean  $\pm$  SD, n=17-40 individual cells. ns=not significant (t-tests). Figure 6. Dexamethasone regulates cardiac mitochondrial fatty acid oxidation in perinatal mice in vivo Pregnant C57Bl/6J dams were injected (i.p.) with 0.5mg/kg dexamethasone or vehicle at E13.5 (A) or E16.5 (B). C57Bl/6J neonates (C) were injected on P1 with dexamethasone (0.5mg/kg) or vehicle. After 24 hours hearts were excised and analysed by qRT-PCR for genes involved in fatty acid oxidation. Data are from n=8-10 individual animals from at least 5 litters/group. Data are mean ± SD and were analysed by two-way ANOVA followed by post hoc Sidak's tests; \*p<0.05, \*\*p<0.01, \*\*\*p<0.001, \*\*\*p<0.0001. Figure 7. Cardiac GR and PGC-1α expression are reduced following treatment with antenatal corticosteroids in a sheep model of pre-term birth Lambs were delivered preterm at 122±1 days (term being ~147 days), 48 hours after initiating a course of celestone (Cel: 2 doses, 24 hours apart) or 24 hours after a single dose of betamethasone acetate (BmA). (A, B) mRNA encoding GR (NR3C1) and (C, D) PGC-1α (PPARGC1A) were measured by qPCR in the left ventricle (LV) and right ventricle (RV). Data are mean ± SD, ns=not significant, \*p<0.05, \*\*p<0.01, analysed by one way ANOVA with post hoc Tukey's tests. For RV: Veh n=10, BmA n=10, Cel n=9. For LV: Veh n=11, BmA n=10, Cel n=9.





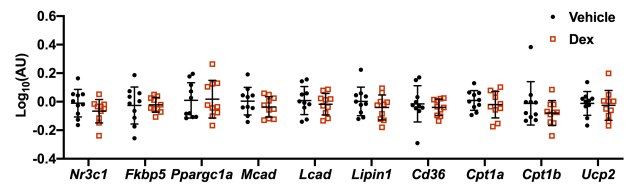




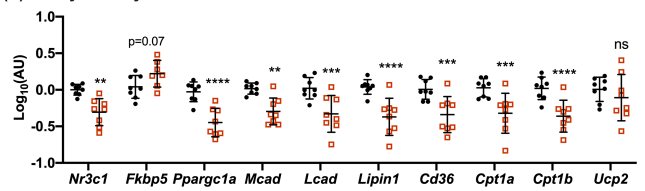








#### (B) Embryonic day 17.5



## (C) Post natal day 2

