

Metabolic characterization of volume overload heart failure due to aorto-caval fistula in rats

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Abstract Metabolic interactions between adipose tissue and the heart may play an active role in progression of heart failure (HF). The aim of the study was to examine changes in myocardial and adipose tissue metabolism and gene expression in a rat HF model induced by chronic volume overload. HF was induced by volume overload from aorto-caval fistula (ACF) in 3-month-old male Wistar rats and animals were studied in the phase of decompensated HF (22nd week). HF rats showed marked eccentric cardiac hypertrophy, pulmonary congestion, increased LV end-diastolic pressure, and intraabdominal fat depletion.

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HF rats had preserved glucose tolerance, but increased circulating free fatty acids (FFA) and attenuated insulin response during oral glucose challenge. Isolated organ studies showed preserved responsiveness of adipose tissue lipolysis and lipogenesis to epinephrine and insulin in ACF. The heart of HF animals had markedly reduced triglyceride content (almost to half of controls), attenuated anti-oxidative reserve (GSH/GSSG), upregulated HF markers (ANP, periostin, thrombospondin-4), specific signaling pathways (Wnt, TGF- β), and downregulated enzymes of mitochondrial fatty acid oxidation, citric acid cycle, and respiratory chain. Adipose tissue transcription profiling showed upregulated receptor for gastric inhibitory polypeptide. In conclusion, ACF-induced HF model

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displays several deregulations of systemic metabolism. Despite elevation of systemic FFAs, myocardial triglycerides are low and insulin levels are attenuated, arguing against a role of lipotoxicity or insulin resistance in this model. Attenuated postprandial insulin response and relative lack of its antilipolytic effects may facilitate intraabdominal fat depletion observed in ACF-HF animals.

Keywords Heart failure · Lipid metabolism · Insulin · Triglycerides · Body composition · Free fatty acids

Introduction

Heart failure is often associated with derangements of systemic and myocardial fat metabolism that may have a role in the disease progression [1–5]. Fatty acids (FA) represent the predominant energetic substrate for the heart, covering 50–70% of myocardial ATP need [2]. The majority of FA supplied to the heart originates from adipose tissue. FA are released from adipocytes by lipolysis and transported in the bloodstream as free FA (FFA). After the entry into a cardiomyocyte, FA is either imported into mitochondria and oxidized or re-esterified into triglycerides (TG) and stored [6].

In chronic HF, excessive fat mobilization and weight loss is associated with poor prognosis [7]. Characteristics of adipose tissue in HF and the mechanisms of adverse effects of fat mobilization on cardiac function remain poorly understood. Circulating FFA are often elevated [8, 9] due to increased levels of prolipolytic hormones (catecholamines, TNF- α , and angiotensin-II) [10] that dominate over of antilipolytic factors, primarily insulin. Elevated circulating FFA can also contribute to insulin resistance [11] that is independently associated with adverse prognosis in HF [12].

Myocardial metabolism in HF is characterized by downregulation of enzymes of β -oxidation of FA and of other mitochondrial enzymes [3, 13, 14]. It has been suggested that imbalance between myocardial FFA delivery and utilization may lead to myocardial FA overload, TG over-accumulation, altered gene transcription due to lipid-responsive regulatory elements [3], and to deposition of toxic lipid intermediates [15] that can promote cardiac dysfunction. Evidence for cardiac “lipotoxicity” was found in obese rats [16], in transgenic animals [17] and in diabetic or obese end-stage HF patients [15]. The extent of myocardial lipid accumulation in HF due to hemodynamic overload, without concomitant diabetes or obesity, has never been addressed even in animal models.

Despite volume overload is clinically common cause or contributing factor of HF, it is far less studied than pressure overload or chronic myocardial infarction. Importantly,

mechanisms and consequences of cardiac adaptation may differ according to type of overload [18]. Chronic volume overload due to aorto-caval fistula (ACF) in rats is an established model of chronic HF [19–22]. The creation of ACF leads rapidly to development of compensatory cardiac hypertrophy to maintain elevated, but largely ineffective cardiac output. After long asymptomatic stage, HF signs gradually develop and overt HF is present in 80% of animals at 21st week post-fistula [22]. The ACF-HF model recapitulates many features of human advanced HF, including gradual onset, elevated cardiac filling pressures [21, 23–25], diminished “effective” cardiac output with splanchnic hypoperfusion [21], neurohumoral activation [23], and altered calcium handling with diminished cardiac energetic efficiency [26]. Moravec et al. [14, 27] had documented that myocardial long-chain FA oxidation is impaired in ACF rats [2], but substrate metabolism at the whole body level has not yet been further studied in this model. The aim of the study was to examine changes in myocardial and adipose tissue metabolism and gene expression in a rat HF model induced by chronic volume overload.

Materials and methods (for details see online Suppl. 1)

Chronic heart failure model

Volume overload HF was induced in male Wistar rats (300–350 g) by creating ACF using 1.2 mm needle from laparotomy in general anesthesia, as described previously [19, 22, 28]. The animals were kept on a 12/12-h light/dark cycle, fed normal salt/protein diet (0.45% NaCl, 19–21% protein, SEMED, and CR) and were weighted weekly until the end of the experiment in the 22nd week. The investigation conformed to the NIH Guide for the care and use of laboratory animals (NIH Publication No. 85-23, 1996), Animal protection laws of the Czech Republic (311/1997) and was approved by the ethics committee of IKEM.

Echocardiography and hemodynamics

Echocardiography was performed in general anesthesia with 10 MHz probe (Vivid System 5, GE, USA) 10th week post-ACF. End-systolic and end-diastolic LV volumes were derived by cubic equation [20] and stroke volume as their difference. Relative wall thickness was defined as sum of end-diastolic anterior wall + posterior wall LV thickness, divided by end-diastolic LV diameter. Hemodynamics was measured at the study end (22th week) with 2F micro-manometer catheter (Millar Instruments) inserted into the aorta and LV via carotid artery in general

anesthesia. The presence of ACF was verified from laparotomy and the animals were exsanguinated. The coronary tree of the excised heart was rapidly flushed with cardioplegic solution. The organs were weighted and normalized to body weight or tibial length.

Myocardial morphology and fat staining

Basal portions of the left ventricle were fixed in 4% paraformaldehyde, embedded into TissueTek, cut and stained with Sudan Black for lipids. Staining with anti- α actinin antibody together with Alexa488-conjugated wheat germ agglutinin (WGA) was performed to evaluate myocyte size and proportion of fibrosis. Images were acquired from Olympus BX51 microscope and Leica SPE confocal system and quantitatively analyzed using ImageJ (NIH) software.

Gene expression and GSEA pathway analysis in the myocardial and adipose tissue

Samples of LV tissue and epididymal fat ($n \geq 6$ in each group) were immediately harvested into RNA preserving solution (RNA-Later, AmbioGen, USA). Total RNA was isolated (RNeasy-MicroKit, Qiagen, USA), checked for integrity, amplified, and hybridized on Illumina RatRef-12v1 Expression BeadChip (Illumina, USA). The raw data were analyzed and processed using *beadarray* package of the *Bioconductor*, as described before [29]. Analysis of differential expression was performed with the *Limma* package [30] and annotated against RatRef_12_V1_0_R3_11222119_A.bgx manifest (Illumina, USA). Cut-off level for differential regulation was set to fold change >2 or <0.5 , with Storey $q < 0.01$. The data are MIAME-compliant and are deposited in the ArrayExpress database (accession #: E-MTAB-190). Gene set enrichment analysis (GSEA) was performed on the samples from LV tissue on gene pathways defined by the KEGG database (release 57.0) [31]. Only the pathways with false discovery rate < 0.05 are reported.

Quantitative real-time PCR

Reverse transcription was performed by QuantiTect[®] Reverse Transcription Kit (QIAGEN Inc., USA). The qRT-PCR was performed on LightCycler 2.0 System using LightCycler[®] 480 DNA SYBR Green I Master kit (Roche Diagnostics, Germany) and results were analyzed by LightCycler software. Crossing point values were further determined using the *R environment* (R Development Core Team 2007). Detailed description of the analysis and the list of amplicons/primers of target and housekeeping genes are provided in Supplementary file 1 and 3, respectively.

Biochemical analyses

Oral glucose tolerance tests (OGTT, 300 mg glucose/100 g BW after overnight fasting) were performed at the 21st week by sampling tail vein at 0, 30, 60, and 120 min. Serum glucose was measured by enzymatic assay (Pliva-Lachema, CR), FFA with colorimetry (Roche Diagnostics, Germany), and insulin with rat-specific ELISA (Mercodia, Sweden). Tissue TG were measured after N₂ pulverization, and chloroform/methanol extraction with enzymatic assay (Pliva-Lachema, CR) was also used for serum TG. The activity of SOD was analyzed by the reaction of nitroethrazolium blue reduction and nitrothymasane formation [32]. The reduced (GSH) and oxidized form of glutathione (GSSG) was determined by high performance liquid chromatography with fluorescent detection (Chromsystems, Germany). The level of thiobarbituric acid-reactive substances (TBARS) was determined by the reaction with thiobarbituric acid [33].

Metabolic assessment of isolated adipose tissue

Basal and epinephrine-stimulated epididymal adipose tissue lipolysis was examined during a 2-h in vitro fat tissue incubation in Krebs–Ringer buffer with or without epinephrine (0.25 mg ml⁻¹) as described previously [34]. Lipolysis was quantified as FFA release into the medium. Basal and insulin-stimulated lipogenesis was quantified by incorporation of ¹⁴C-U-glucose into neutral, chloroform-extracted lipids during 2 h incubation in Krebs–Ringer bicarbonate buffer with or without insulin (250 μ U ml⁻¹) as described previously [34].

Statistics

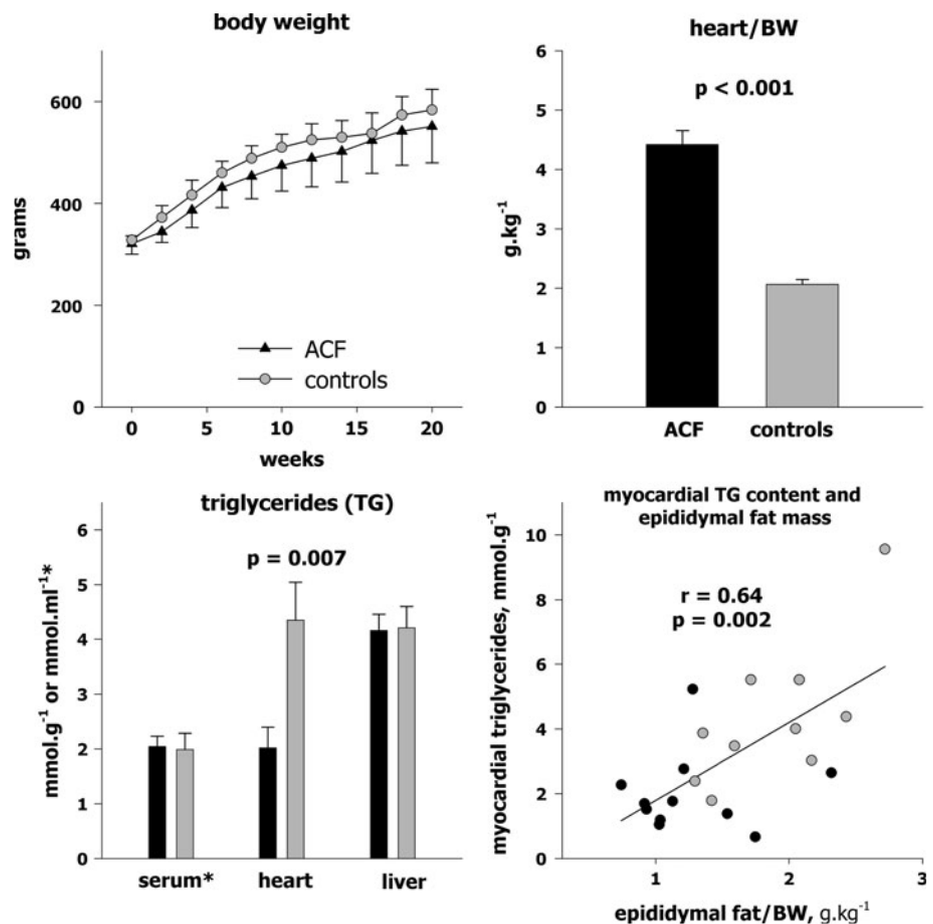
Animals that died during the experiment (18% total mortality including ACF operation) were excluded from the analyses. Sample size for echocardiography was 12–14/group, for other methods 5–8/group. Data are expressed as means \pm SD, in graphs as means \pm SE. Student's *t* test statistics was used for comparisons, and *P* value < 0.05 was considered significant. Gene expression differences were compared with false discovery rate (FDR) and described with Storey's *q* value.

Results

Echocardiography, hemodynamics, and organ weights

ACF animals had similar growth rate and final body size as controls (Fig. 1). Echocardiography (Table 1) confirmed cardiac enlargement, reduced relative wall thickness (0.33 ± 0.05 vs. 0.45 ± 0.05 , $P < 0.001$), mildly reduced

Fig. 1 *Upper panel* Body weight during the course of experiment, and the weight of the whole heart normalized to body weight. *Lower panel* Concentration of triglycerides (TG) in heart, or liver tissue and in serum, correlation between myocardial TG content and epididymal fat. *Gray* controls, *black* aorto-caval fistula (ACF)



fractional shortening, and markedly increased stroke volume and cardiac output (+196%) in ACF. By the end of experiment (22th week), 65% of ACF animals showed clinical HF signs (lethargy, jagged fur, and labored breathing). Hemodynamics (Table 1) showed similar systemic arterial pressure, LV dP/dt_{max} and maximal LV pressures, but markedly elevated LV end-diastolic pressure (+263%), longer systolic duration (+21%), or LV relaxation time constant τ (+33%), and slower relaxation rate dP/dt_{min} (+24%, $P = 0.07$) in ACF. ACF animals had a marked increase in heart (+148%), lung (+72%), and liver weights (+17%) (Table 2). Despite similar body weights, the ACF group had surprisingly lower epididymal fat pad weight (−29%) (Fig. 6). Subcutaneous (inguinal) fat pad and kidney weights were similar in both groups.

Myocardial morphology and biochemistry

LV mid-wall histology (Fig. 2) showed biventricular myocyte hypertrophy due to myocyte elongation [5], with no increase in morphologic measure of fibrosis (WGA-positive area: ACF: $25 \pm 0.8\%$, sham: $30 \pm 4\%$, $P = 0.39$). Sudan Black staining for lipids, performed to examine possible lipid

overload, showed no visible fat vacuoles and no difference in stain intensity (ACF: 205 ± 22 , sham 208 ± 23 intensity units, $P = 0.13$). Myocardial TG concentration was by 53% lower in ACF (2.2 ± 1.2 vs. $4.3 \pm 2.2 \mu\text{mol g}^{-1}$, $P = 0.006$, Fig. 1), but the liver TG content was similar (ACF: 4.2 ± 0.8 vs. $4.2 \pm 0.8 \mu\text{mol g}^{-1}$, $P = 0.9$). Myocardial TG correlated ($r = 0.64$, $P = 0.002$) with epididymal fat pad weight (Fig. 1). Parameters of redox state and oxidative stress were also analyzed in the myocardium. ACF group had a similar myocardial level of reduced glutathione (GSH: 4.4 ± 1.4 vs. $4.3 \pm 0.3 \mu\text{mol g}^{-1}$, $P = 0.8$), a trend toward more of oxidized glutathione (GSSG: 0.39 ± 0.11 vs. $0.32 \pm 0.03 \mu\text{mol g}^{-1}$, $P = 0.12$), and significantly lower redox reserve (GSH/GSSG ratio, Fig. 3) than controls. Superoxide dismutase (SOD) tended to be lower in ACF hearts ($P = 0.07$), and there was no difference in myocardial TBARS concentration ($P = 0.44$) as a marker of oxidative stress.

Myocardial gene expression

Exploratory principal component analysis (PCA) and hierarchical clustering (HC) revealed pronounced

Table 1 Echocardiography and invasive hemodynamics

	Sham	ACF	<i>P</i> value
Echocardiography			
Heart rate, s ⁻¹	368 ± 39	350 ± 32	0.2
Body weight, g	422 ± 30	427 ± 24	0.6
LV diastolic diameter, mm	7.3 ± 0.4	10.9 ± 0.9	<0.001
LV systolic diameter, mm	4.3 ± 0.5	6.9 ± 0.8	<0.001
Fractional shortening, %	41 ± 2.9	36 ± 3.5	0.002
LV anterior wall thickness in diastole, mm	1.6 ± 0.1	1.8 ± 0.2	0.05
LV posterior wall thickness in diastole, mm	1.6 ± 0.2	1.8 ± 0.2	0.03
RV diastolic diameter, mm	4.0 ± 0.9	5.8 ± 1	<0.001
LV end-diastolic volume, ml	0.38 ± 0.1	1.31 ± 0.3	<0.001
Stroke volume, ml	0.30 ± 0.1	0.96 ± 0.2	<0.001
Cardiac output, ml	110 ± 16	326 ± 64	<0.001
Invasive hemodynamics			
Aortic mean pressure, mmHg	99 ± 25	89 ± 22	0.5
Aortic pulse pressure, mmHg	45 ± 19	46 ± 10	0.9
Maximal LV pressure, mmHg	114 ± 21	120 ± 22	0.6
End-diastolic LV pressure, mmHg	4.1 ± 2.3	14.9 ± 4.6	0.0003
Heart rate, s ⁻¹	351 ± 53	338 ± 88	0.8
Systolic time of total cycle, %	48 ± 4	58 ± 7	0.009
dP/dt _{max} , mmHg s ⁻¹	7412 ± 2919	8170 ± 3568	0.7
dP/dt _{max} /IP, s ⁻¹	112 ± 27	122 ± 35	0.6
dP/dt _{min} , mmHg s ⁻¹	-7987 ± 2009	-6053 ± 1550	0.07
LV relaxation constant τ, ms	12 ± 2	16 ± 3	0.006

Values are means ± SD. Echocardiography performed 10th week after ACF, *N* = 12–14/group. LV left ventricle, RV right ventricle, ACF aortocaval fistula group. Hemodynamics performed at the end of study. *N* = 6–8/group, dP/dt_{max} maximal rate of LV pressure increase, IP instantaneous pressure, dP/dt_{min} maximal rate of LV pressure decrease

differences in gene expression between groups. Although the variation among ACF animals was large, the samples formed a well-defined and separated cluster. Of 23401 transcripts analyzed, 249 were differentially regulated (fold change >2 or <0.5, *q* < 0.01), 186 upregulated and 63 downregulated in ACF (heat maps in Supplementary file 2). GSEA analysis using KEGG database [31] was used to identify systematic changes of transcription in pathways (Table 4). The list of the most differentially expressed genes (*q* < 0.01, fold change > 3.8 or <0.38) in those pathways are presented in the Table 3. Fatty acid metabolism, oxidative phosphorylation, glycolysis, citric acid cycle, branched amino-acid degradation, and PPAR signaling pathway were the most downregulated pathways. In contrast, pathways of complement cascade, extracellular matrix–receptor interaction, cell adhesion, phagosome/lysosome, and antigen presentation and cell adhesion molecules were the most upregulated. Quantitative RT-PCR confirmed upregulation of heart failure markers (natriuretic peptide precursor A), EC-matrix components (periostin, thrombospondin-4) and components of regulatory WNT1 pathway (WISP2), along with downregulation

of enzymes of glycolysis (enolase 3) and FA oxidation (hydroxyacyl-CoA dehydrogenase) (Fig. 4).

Adipose tissue gene expression

Gene expression analysis was also performed in epididymal adipose tissue to identify changes induced by the presence of HF. PCA and HC revealed well-separated clusters of control and ACF rats. As ACF animals showed larger variation, only eight genes were differentially regulated at fold-change level > 2 or <0.5, *q* < 0.01 (all downregulated in ACF). The differences between groups were less pronounced than in myocardial tissue, so a GSEA was not performed. The list of the most differentially expressed genes (*q* < 0.01, fold change >1.5 or <0.5) is presented in Table 5. Full lists of differentially expressed genes and pathways (*q* or FDR < 0.05) are provided in online (Suppl. 2 and 3).

Oral glucose tolerance test

To assess the dynamics of metabolic substrates in the blood and its coordination by insulin, oral glucose tolerance test

Table 2 Organ weights

	Sham	ACF	<i>P</i> value
Body weight, g	487 ± 42	510 ± 37	0.3
Tibia length, mm	42 ± 2	43 ± 0.4	0.4
Heart weight, g	1.05 ± 0.08	2.61 ± 0.25	<0.0001
Heart weight/TL, mg mm ⁻¹	2.50 ± 0.11	6.14 ± 0.57	<0.0001
Lung weight, g	1.68 ± 0.27	2.88 ± 0.34	<0.0001
Lung weight/BW, g kg ⁻¹	3.44 ± 0.47	5.68 ± 0.88	0.0003
Liver weight, g	14.2 ± 1.1	16.6 ± 2.0	0.03
Liver weight/BW, g kg ⁻¹	29.3 ± 1.7	32.4 ± 2.2	0.02
Left kidney weight, g	1.26 ± 0.1	1.26 ± 0.08	0.9
Left kidney weight/BW, g kg ⁻¹	2.60 ± 0.2	2.48 ± 0.25	0.4
Right kidney weight, g	1.28 ± 0.1	1.29 ± 0.06	0.9
Right kidney weight/BW, g kg ⁻¹	2.63 ± 0.2	2.54 ± 0.23	0.4
Fat compartments			
Epididymal fat pad, g	9.75 ± 1.85	6.90 ± 2.72	0.01
Inguinal subcutaneous fat pad, g	1.05 ± 0.06	0.96 ± 0.69	0.8

Values are means ± SD. 22nd week after ACF—the end of the study. *N* = 6–8/group. *LV* left ventricle, *RV* right ventricle, *TL* tibial length, *BW* body weight

was performed. ACF and controls had similar baseline serum glucose (4.5 ± 0.6 vs. 4.6 ± 0.7 mmol l⁻¹, $P = 0.5$) and insulin (26 ± 20 vs. 37 ± 30 pmol ml⁻¹, $P = 0.25$), but increased serum FFA (1.57 ± 0.3 vs. 1.15 ± 0.2 mmol l⁻¹, $P = 0.001$, Fig. 5). Glycaemia at 30', 60', and 120' ($P = 0.12$; 0.11; and 0.40) and its AUC area (819 ± 101 vs. 761 ± 63 mmol s l⁻¹, $P = 0.2$) were similar between groups. Interestingly, serum insulin response during OGTT was markedly attenuated in HF animals—insulin at 60' and 120' were significantly lower in ACF (43 ± 27 vs. 105 ± 91 , $P = 0.007$; 58 ± 44 vs. 185 ± 151 pmol ml⁻¹, $P = 0.001$). FFA at 60' OGTT remained elevated in ACF by 63% ($P = 0.0007$).

Adipose tissue metabolism

To understand the mechanism for ACF-associated intraabdominal fat depletion and to assess responsiveness of fat tissue to humoral stimuli, epididymal adipose tissue was examined in vitro. Basal and epinephrine-stimulated of lipolysis was intact in ACF animals. Similarly, the lipogenesis was normal both at baseline and after insulin stimulation (Fig. 6), indicating preserved responsiveness of adipocytes.

Discussion

Presented study revealed several specific, previously not described metabolic abnormalities in rats with HF due to

ACF. The heart of HF animals had diminished redox reserve (GSH/GSSG ratio), upregulated HF markers, and downregulated mitochondrial metabolic pathways, including β -oxidation of FA. Despite circulating FFA were elevated, myocardial TG content was reduced in ACF group compared to sham. Body fat distribution was also affected by the presence of HF-ACF animals demonstrated selective depletion of intraabdominal adipose tissue, probably due to enhanced fat mobilization. The latter finding might be linked to abnormally attenuated insulin response to glucose challenge in ACF rats. The results argue against the role of myocardial lipid overload in this particular HF model and indicate that hypoinsulinaemia can contribute to intraabdominal fat tissue depletion observed in HF. The results of the study help to understand mechanisms leading to cardiac cachexia and suggest possible targets for an intervention.

Cardiac remodeling due to ACF

Creation of aorto-caval fistula led to volume overload with an increase of cardiac output by ~200% and to marked eccentric biventricular hypertrophy. Increased LV end-diastolic filling pressures and lung weights confirmed decompensated HF by the end of the study (22nd week post-ACF). Invasive hemodynamics showed relatively preserved LV contractility (dp/dt_{max}), although to fully exclude latent contractile dysfunction we would have to use truly load-independent contractile measures [35]. Diastolic function was already impaired, probably due to combined effect of eccentric chamber remodeling, increased myocardial mass, and upregulation of ECM proteins that may increase passive ventricular stiffness (lysyl oxidase, fibronectin, collagen VIII, and thrombospondin-4). The findings are in line with previous reports from ACF-HF model indicating that intense neurohumoral activation precedes the gross decline of myocardial performance [36]. Myocardial gene expression analysis did confirm robust changes typical for HF phenotype, with more than 1% of targeted transcripts being differentially expressed. The highest overexpression in ACF hearts (Table 3) was found for natriuretic peptide precursors, thrombospondin-4 [37], Wnt-pathway [18], TGF- β pathway components, periostin [38], and thyrotropin-releasing hormone [39] that were already identified as markers of stress-induced myocardial remodeling. Gene set enrichment analysis (GSEA) that detects concordant changes in transcription in functional pathways [31], indicated widespread upregulation of genes of extracellular matrix–receptor interaction, cell adhesion molecules (Table 4). This probably reflects increased matrix turnover due to stretch-activated transcription in ACF hearts, although the histology did not show excessive ventricular fibrosis, confirming previous reports [5, 18, 23, 24, 28]. The upregulation of complement cascade components, phagosome/

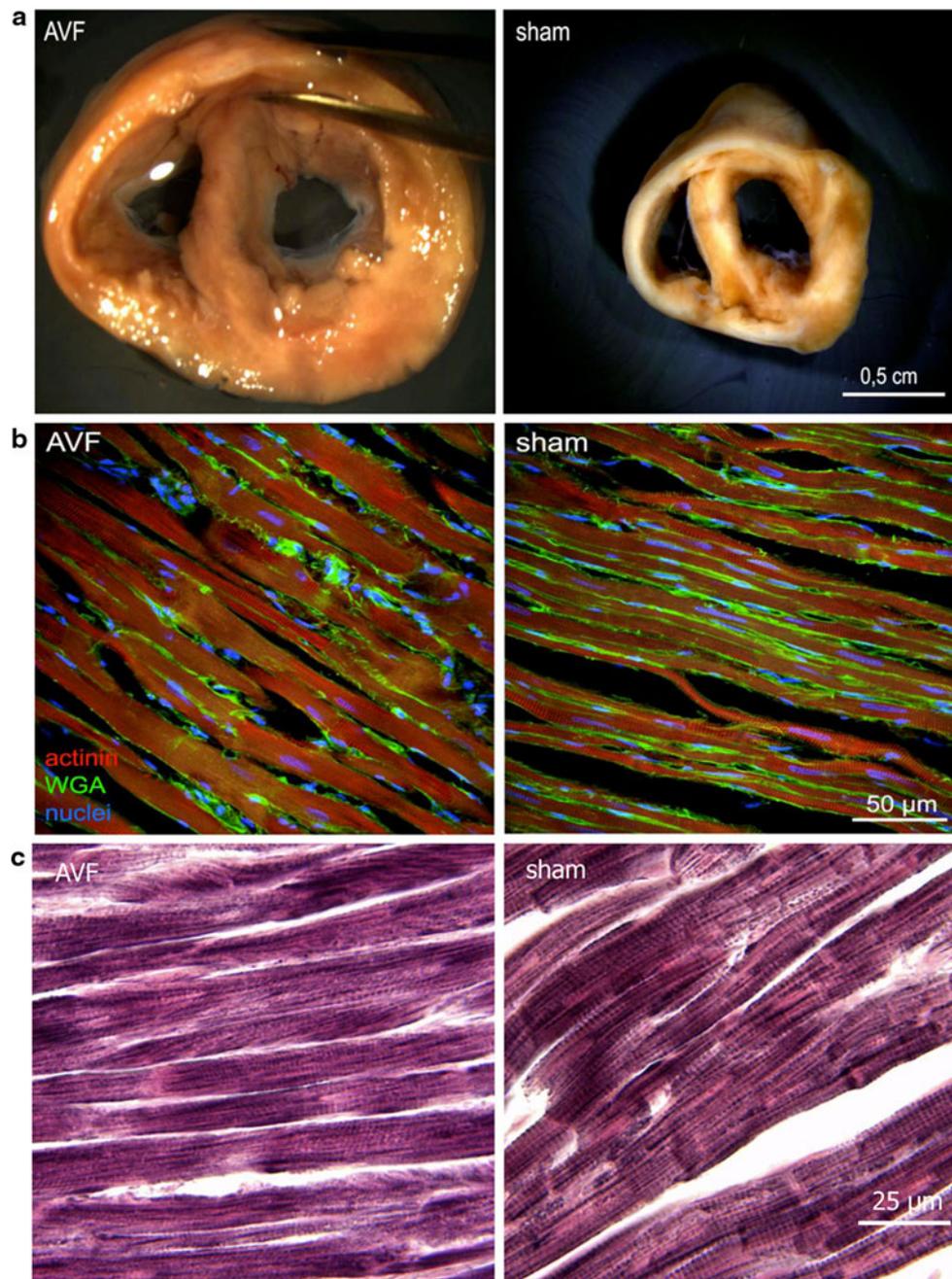


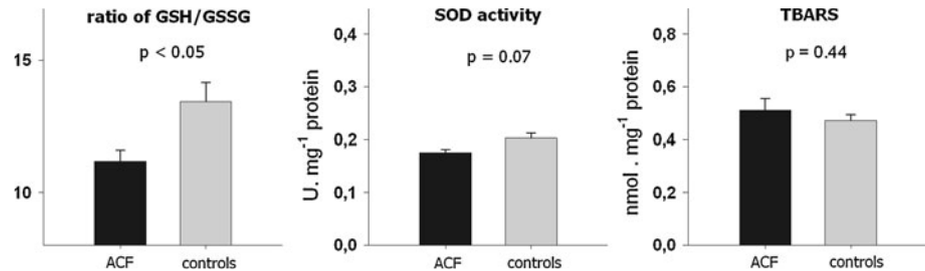
Fig. 2 Myocardial morphology. **a** Transversal section of the heart illustrates marked biventricular enlargement in the ACF animal (*left*) compared to control (*right*). **b** Confocal microscopic image of mid-section of the left ventricle. *Red* myocytes stained with α -actinin Ab, *Green* tissue fibrosis stained with Alexa488-conjugated wheat germ

agglutinin, *Blue* Hoechst nuclear counterstaining. Note wider and longer myocytes in ACF, but similar extent of fibrosis. **c** Sudan Black staining for fat. Note similar density of staining and absence of lipid inclusions in both groups. Figure in color is available in online version

lysosome, and cytokine–cytokine interaction pathways may reflect inflammatory cell infiltration, yet active synthesis of complement components in stressed cardiomyocytes is also possible [40]. The most significant change detected by GSEA was downregulation of pathways for FA oxidation, citric acid cycle, respiratory electron chain, and branched amino-acid catabolism (Table 4). Similar broad

downregulation of mRNAs of metabolic enzymes had been described also in other models of advanced HF [3] and it results from attenuated activity of PPAR α /RXR α /PGC-1 α regulatory complex [2]. Despite the functional impact of diminished transcription of these important genes is attenuated by redundancy in enzymatic activities [41], ACF hearts do show altered substrate use with selective decrease

Fig. 3 Markers of myocardial oxidative stress. *Left* The ratio of reduced to oxidized glutathione (*GSH/GSSG*), *middle* superoxide dismutase activity (*SOD*), *right* thiobarbituric acid-reactive substances (*TBARS*). *Gray* controls, *black* aorto-caval fistula (*ACF*)



in oxidation of long-chain FA [14, 27, 42] and the critical limit apparently is intramitochondrial import of fatty acyls [14].

Decreased intramyocardial TG content in ACF hearts

In ACF group, myocardial TG content was reduced almost to a half compared to control despite circulating FFA levels were elevated. This important finding suggest that myocardial TG content does not merely reflect prevailing serum FFA concentration as suggested [3, 43], but it is also affected by the presence of heart failure or perhaps by the cause of heart hypertrophy. Similarly to the study, 30% reduction in myocardial TG pool was described by Lewandowski's group in pressure-overloaded rat hearts [44]. The reduction of myocardial TG content was explained by an increase of "anaplerotic" influx of carbons into the citric acid cycle via malic enzyme [45] that competes with other NADPH-consuming reactions, like TG synthesis or glutathione reduction. This may occur in ACF as well, since we noticed a parallel reduction of GSH/GSSG ratio and TG content in ACF hearts. Low TG content might also be explained by limited intracellular FFA import, because several transport genes were downregulated in ACF hearts (CD36/FAT, FABP3, and FATP3, fold change: 0.7–0.8, $q = 0.02$ –0.004). On the other side, physiologic, exercise-induced cardiac hypertrophy is associated with an increase of myocardial TG content [46]. Myocardial TG content in rodents might have implications for cardiac reserve [4], as 10–30% of ATP production in heart comes from oxidation of intramyocardial TG [47]. Conversely, excessive myocardial TG accumulation is linked to "lipotoxicity" [15, 48]. Recent evidence suggests that not TG itself, but rather diacylglycerols, ceramides, or acylcarnitines are responsible for these toxic effects [49]. The observation of diminished myocardial TG content in pathological heart hypertrophy speaks against causal involvement of lipotoxicity in non-diabetic, non-obese HF.

Attenuated response of insulin to oral glucose challenge in ACF animals

Insulin has a key role in coordination of substrate metabolism [50], but whether ACF-induced HF has an impact on

insulin levels has never been addressed. Against the expectations, HF animals showed normal basal serum insulin, but attenuated increase of insulin after oral glucose loading. The reasons for attenuated postprandial dynamics of insulin in ACF are not clear. Diminished glucose absorption is ruled out, because post-load glycaemia was even higher in ACF. A steal effect of ACF is also improbable because the shunt is distal to the pancreas. Plausible explanation could be in insufficient insulin release due to HF-induced splanchnic vasoconstriction [21, 51] and/or due to protracted exposure of pancreatic β -cells to increased FFA that impairs insulin secretion [52]. Despite of postprandial hypoinsulinaemia, the whole body glucose tolerance (glucose-time AUC area) was maintained in ACF rats by an unknown mechanism. Extrapaneatic effects of incretins or increased insulin sensitivity of peripheral tissues could be involved and need further investigation. The findings suggest that drugs aiming to improve cardiac metabolism by improving insulin sensitivity may not be effective in ACF-induced volume overload HF. Because secretion of insulin after a meal suppresses adipose tissue lipolysis [53], relative postprandial *hypoinsulinaemia* in ACF may contribute to increased FFA release from adipose tissue and to body fat loss.

Depletion of intraabdominal adipose tissue in ACF

Animals with ACF had significantly reduced epididymal fat weight, although total body weight was similar, probably due to more pronounced fluid retention. Epididymal fat is highly vascularized and dynamic fat pool and represents 30–50% of confined intraabdominal depots in rats. Depletion of intraabdominal (retroperitoneal) fat was also reported in rat model of cardiac volume overload from aortic regurgitation [54]. In the study, adipose tissue of ACF animals showed several differentially expressed genes (Table 5), but the changes were less pronounced than in cardiac tissue. Interestingly, the most upregulated gene in HF adipose tissue was a receptor for gastric inhibitory polypeptide (GIP), a gut-produced hormone (incretin) with insulin-releasing and extrapancreatic glucoregulatory actions that also regulates fat deposition in adipocytes [55]. Recent genome-wide association study found that genetic

Table 3 Most differentially regulated genes in the heart (assigned to KEGG pathways)

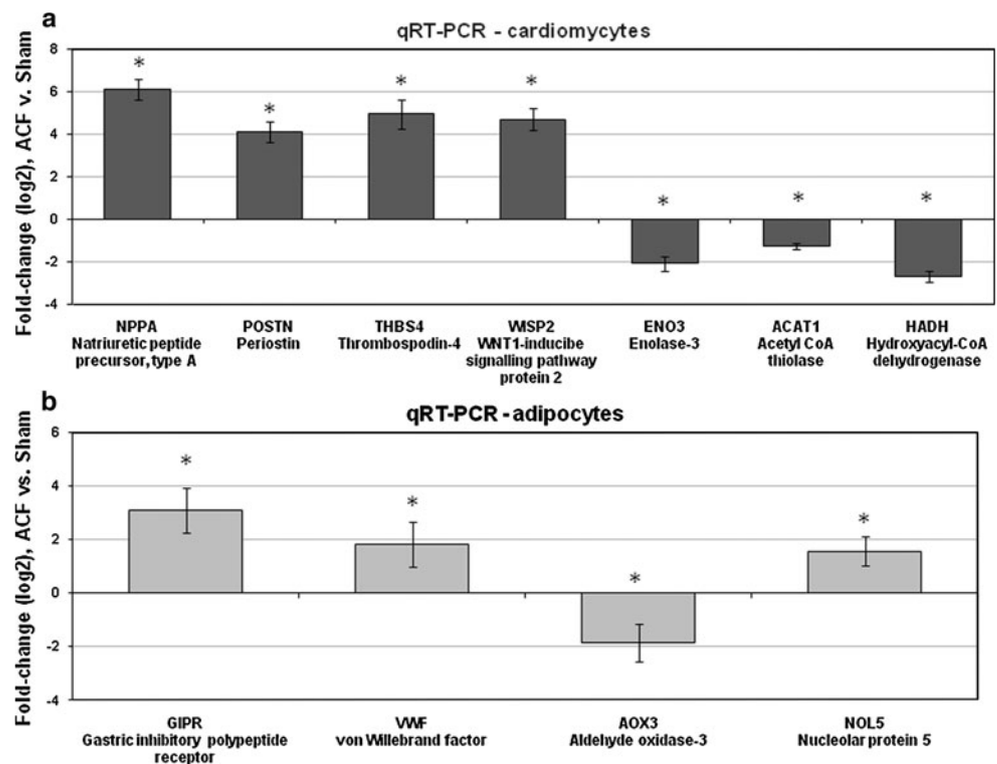
Gene description, name of pathway	Fold change	Adjusted <i>P</i>	Gene symbol	RefSeq ID
Phagosome (rno04145)				
Similar to Fcγ-(IgG) receptor II, α (predicted)	4.5	<0.00001	LOC498276	XM_573502.1
Complement and coagulation cascades (rno04610)				
Complement component 4, gene 2	5.6	0.00014	C4-2	NM_001002805.1
Complement component 1, qβ subcomponent	5	<0.00001	C1qb	NM_019262.1
Complement component 1, qC subcomponent	4.4	<0.00001	C1qc	NM_001008524.1
Lysosome (rno04142)				
CD68 antigen (predicted)	4.5	<0.00001	Cd68	XM_001079491.1
ECM–receptor interaction (rno04512)				
Thrombospondin 4	12.9	0.00001	Thbs4	XM_342172.3
Latent TGF-β binding protein 2	10.5	0.00001	Ltbp2	NM_021586.1
Secreted phosphoprotein 1	8.5	0.00015	Spp1	NM_012881.1
Procollagen, type VIII, α1 (predicted)	5	0.00014	Col8a1_pred.	XM_221536.4
Fibronectin 1	4.5	0.00032	Fn1	NM_019143.1
Lectin, galactose binding, soluble 3	4.4	0.00001	Lgals3	NM_031832.1
Connective tissue growth factor	3.9	0.00009	Ctgf	NM_022266.2
Cytokine–cytokine receptor interaction (rno04060)				
TGF β2	5.1	0.00001	Tgfb2	NM_031131.1
Platelet factor 4	4	<0.00001	Pf4	NM_001007729.1
TNF receptor superfamily, 11b (osteoprotegerin)	3.9	0.00009	Tnfrsf11b	NM_012870.2
Cell adhesion molecules (rno04514)				
Periostin (pred.)	16	0.00002	Postn_pred.	XM_342245.3
WNT1 inducible signaling pathway protein 2	4.5	0.00002	Wisp2	NM_031590.1
Limbic system-associated membrane protein	0.32	0.00017	Lsamp	NM_017242.1
Metabolic pathways (rno01100)				
Chitinase 3-like 1	5.5	0.00004	Chi3l1	NM_053560.1
Lysyl oxidase-like 1	4.2	0.00008	Lox1l	NM_001012125.1
Glutamic pyruvic transaminase 1, soluble	0.38	0.00002	Gpt1	NM_031039.1
Dehydrogenase/reductase member 7C (pred.)	0.25	0.00148	Dhrs7c_pred.	XM_001078936.1
Fatty acid metabolism (rno00071)				
Hydroxyacyl-Coenzyme A dehydrogenase	0.37	0.00079	Hadh	NM_057186.1
Peroxisome (rno04146)				
Epoxide hydrolase 2, cytoplasmic	0.3	0.001	Ephx2	NM_022936.1
Glycolysis/Gluconeogenesis (rno00010)				
Enolase 3	0.23	0.00003	Eno3	NM_012949.1
Other—membrane transport				
FXFD domain-containing ion transport regulator 3	0.38	0.00004	Fxyd3	NM_172317.1
Plasma membrane proteolipid	0.34	0.00001	Plip	NM_022533.1
Solute carrier family 22, member 3	0.31	0.00001	Slc22a3	NM_019230.1
ATPase, Na ⁺ /K ⁺ transporting, α2 polypeptide	0.29	0.00001	Atp1a2	NM_012505.1
Potassium channel, subfamily K, member 2	0.26	0.00008	Kcnk2	NM_172041.1
Other—regulatory				
Natriuretic peptide precursor type A	11.4	0.00225	Nppa	NM_012612.1
Thyrotropin-releasing hormone	7.2	0.0001	Trh	NM_013046.2
Nuclear protein 1	5.8	0.00001	Nupr1	NM_053611.1
TGF α	5.4	<0.00001	Tgfa	NM_012671.1
Hepcidin	4.6	0.00105	Hamp	NM_053469.1
Histone deacetylase 2	0.38	0.00005	Hdac2	XM_342149.3

Table 3 continued

Gene description, name of pathway	Fold change	Adjusted <i>P</i>	Gene symbol	RefSeq ID
Iroquois related homeobox 2	0.37	0.0045	Irx2	XM_001053004.1
Thyroid hormone receptor β	0.35	0.00385	Thrb	NM_012672.1
DNA-damage-inducible transcript 4-like	0.29	0.00004	Ddit4 1	NM_080399.1
Protein disulfide isomerase associated 2 (pred.)	0.23	0.00002	Pdia2_pred.	XM_213263.4
Other				
Tissue inhibitor of metalloproteinase 1	7.9	0.00001	Timp1	NM_053819.1
Protease inhibitor 16 (predicted)	4.7	0.00002	Pi16_pred.	XM_215351.4
Regenerating islet-derived 3 β	4.5	0.00337	Reg3b	NM_053289.1
Suprabasin (pred.)	4	0.00089	Sbsn_pred.	XM_214902.4
Membrane-spanning 4-domains, subfamily A	3.8	<0.00001	Ms4a4a_pred.	XM_001075321.1
Similar to RIKEN cDNA 1700028P14 (pred.)	3.8	0.00116	RGD1560242_pred.	XM_001078678.1
Tropomodulin 4 (predicted)	0.36	0.00008	Tmod4_pred.	XM_001054854.1
Ataxin 2 binding protein 1	0.33	0.00068	A2bp1	XM_001076609.1
Similar to microsomal glutathione S-transferase 3	0.31	0.00033	RGD1561381_pred.	XM_001073485.1

Presented are only transcripts with FC >3.8 or <0.33 and FDR <0.001, full list of differentially regulated transcripts is in online supplement 2 and 3. *KEGG* Kyoto Encyclopedia of Genes and Genomes [31]

Fig. 4 Validation of gene expression changes in cardiomyocytes (a) and adipocytes (b) using quantitative real-time polymerase chains reaction (qRT-PCR). Results are normalized to housekeeping genes (see methods in supplementary file 1) and expressed as fold change in ACF versus Sham animals. **P* < 0.01



variation GIP receptor significantly influences insulin and glucose responses to an oral glucose challenge in humans [56]. An involvement of GIP receptor in HF-induced changes of adipose tissue metabolism has never been reported and it requires further investigation. In vitro

studies of adipose tissue metabolism showed preserved lipolysis and lipogenesis in ACF hearts, with normal responsiveness to epinephrine or insulin stimulation. Increased catecholamines or angiotensin-II [10] can explain increased FFA release in ACF. Previously, it has

Fig. 5 Oral glucose tolerance test. *Upper panel* serum glucose and FFA. *Lower panel* insulin and area under the curve (AUC) of glucose–time plot. 0 min = administration of glucose. Note attenuated increase of insulin after glucose challenge in ACF group

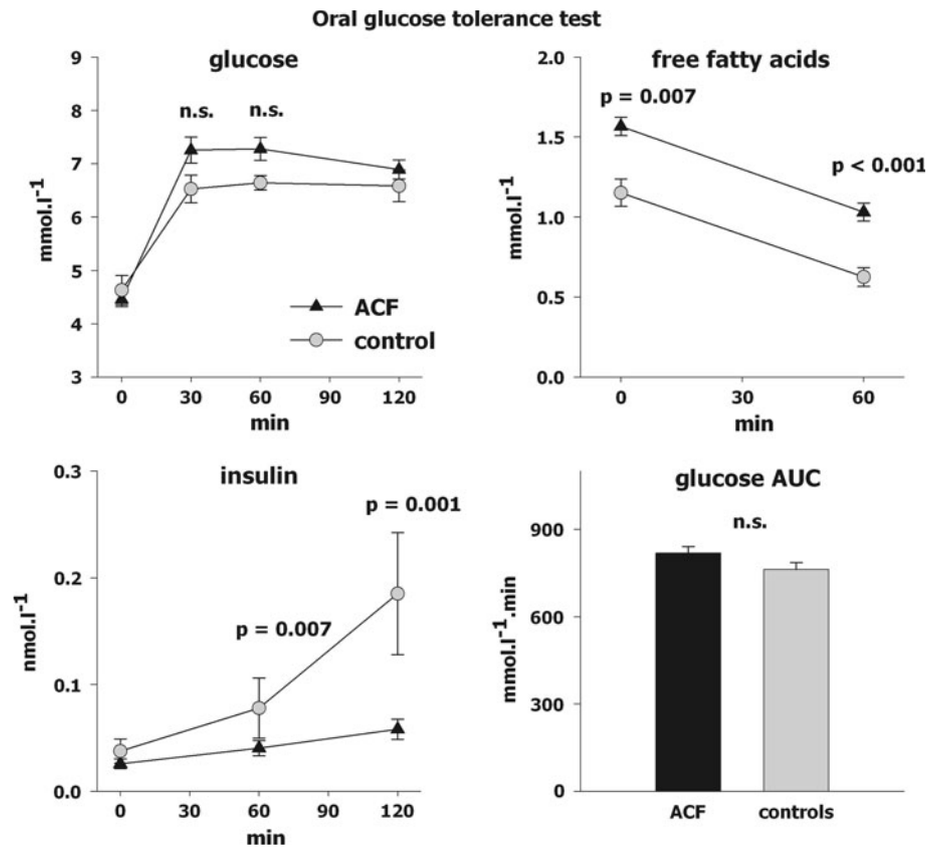
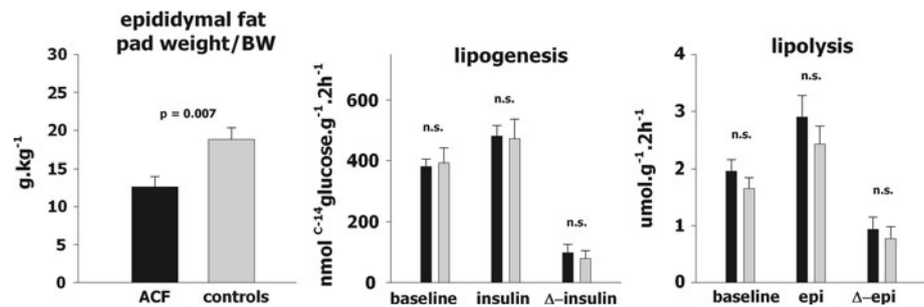


Fig. 6 Epididymal adipose tissue weight and metabolism. Epididymal fat weight, lipogenesis, and lipolysis in basal and epinephrine-stimulated state. Note intraabdominal–epididymal fat tissue depletion and preserved responsiveness of adipocytes to stimuli



been shown in rats that chronic angiotensin-II infusion leads to adipose tissue lipolysis and to weight loss and these effects are attenuated by angiotensin or β -adrenergic receptor blockade [10]. Postprandial hypoinsulinemia and attenuated antilipolytic effects can also contribute to excessive intraabdominal fat mobilization in HF due to ACF. Although HF in general is associated with elevated insulin levels and impaired whole body insulin sensitivity [9, 12], attenuated insulin secretion has been described in patients with decompensated advanced HF [51]. The study raises the question whether correction of hypoinsulinemia might favorably modulate fat depletion, or whether long-term inhibition of lipolysis could be beneficial in HF. Lipolysis in adipose tissue can be inhibited by GPR109A receptor agonists (nicotinic acid and acipimox) [57]. Acute

administration of acipimox acutely reduced cardiac work and oxygen efficiency [58], but long-term effects on cardiac function of in advanced HF have not yet been tested. Lipolysis can also be inhibited by angiotensin receptor antagonists or β -adrenergic receptor blockers [10] and their antilipolytic effects might contribute to favorable effects of these drugs on weight loss and prognosis of patients with HF.

The study has several limitations. First, we performed metabolic characterization of ACF state only in male rats. It has been shown previously that sex and estrogen status are important determinants of cardiac adaptation to volume overload, with less eccentric hypertrophy in females than in males [59, 60], but no study examined yet sex-related differences in myocardial metabolism in ACF. Second, cardiac and metabolic remodeling of the heart is

Table 4 Most differentially regulated pathways in the heart (KEGG pathway analysis)

Pathway description	KEGG ID	<i>P</i> value	FDR	Count	Size
Upregulated in ACF					
Phagosome	mo04145	<0.00001	<0.00001	26	113
Complement and coagulation cascades	mo04610	<0.00001	0.00003	13	39
Lysosome	mo04142	<0.00001	0.00008	17	72
ECM–receptor interaction	mo04512	0.00011	0.00194	10	37
Cytokine–cytokine receptor interaction	mo04060	0.00013	0.00204	17	94
Antigen processing and presentation	mo04612	0.00018	0.00252	12	54
Cell adhesion molecules (CAMs)	mo04514	0.00028	0.00353	14	73
Downregulated in ACF					
Metabolic pathways	mo01100	<0.00001	<0.00001	74	614
Fatty acid metabolism	mo00071	<0.00001	<0.00001	13	27
Citrate cycle (TCA cycle)	mo00020	<0.00001	<0.00001	11	20
Peroxisome	mo04146	<0.00001	<0.00001	15	50
Valine, leucine, and isoleucine degradation	mo00280	<0.00001	<0.00001	11	28
Oxidative phosphorylation	mo00190	0.00001	0.00014	13	65
Pyruvate metabolism	mo00620	0.00010	0.00091	7	24
Glycolysis/Gluconeogenesis	mo00010	0.00030	0.00247	9	46
PPAR signaling pathway	mo03320	0.00039	0.00277	8	38

ACF aorto-caval fistula, FDR false discovery rate, KEGG Kyoto Encyclopedia of Genes and Genomes [31]

Full list of pathways and elements (including heat maps) is in online supplement 2 and 3. *N* = 6/group

Table 5 Most differentially regulated genes in ACF adipose tissue

Gene description	Fold change	<i>q</i> value	Gene symbol	RefSeq_ID
Upregulated in ACF				
Gastric inhibitory polypeptide receptor	1.95	0.00501	Gipr	NM_012714.1
Similar to very large G-protein coupled receptor 1	1.90	0.00571	LOC362068	XM_001056150.1
Similar to CG8841-PA (predicted)	1.84	0.00354	RGD1311422_predicted	XM_001081678.1
Hypothetical protein RDA279	1.78	0.00332	Rda279	XM_001070821.1
Pleckstrin homology domain containing, F2 (pred.)	1.75	0.00401	Plekhf2_predicted	XM_342803.3
von Willebrand factor (Vwf) (predicted)	1.73	0.00665	Vwf	XM_001066203.1
Cyclin L2 (predicted)	1.66	0.00393	Ccnl2_predicted	XM_216597.3
Interferon regulatory factor 3	1.55	0.00401	Irf3	NM_001006969.1
Nucleolar protein 5	1.54	0.00921	Nol5	NM_021754.1
RT1 class I, CE7	1.53	0.00498	RT1-CE7	NM_001008845.1
Camello-like 1	1.52	0.00836	Cml1	NM_133558.1
Armadillo repeat containing 5	1.50	0.00772	Armrc5	NM_001009455.1
Laminin, beta 1 (predicted)	1.50	0.00424	Lamb1_predicted	XM_216679.4
Downregulated in ACF				
Aldehyde oxidase 3	0.33	0.00077	Aox3	NM_001008527.1
Similar to microsomal glutathione S-transferase 3 (pred.)	0.42	0.00958	RGD1561381_predicted	XM_001073485.1
Complement component 6	0.42	0.00401	C6	NM_176074.2
Cystatin E/M	0.42	0.00304	Cst6	NM_133566.1
RT1 class II, locus Db1	0.43	0.00077	RT1-Db1	NM_001008884.1
RT1 class II, locus Da	0.45	0.00681	RT1-Da	NM_001008847.1
Nephrosis 1 homolog (nephrin)	0.46	0.00536	Nphs1	NM_022628.1
Similar to hypothetical protein D630003M21	0.48	0.00698	RGD1563354_predicted	XM_001068317.1

Ranked by fold change. Full list of differentially regulated transcripts is in online supplement 3. *N* = 6–8/group

time-dependent, dynamic process, but we studied just one time-point—advanced HF stage. Further studies in different stages of HF development are warranted. Third, statistical methods that we used for gene expression analysis may not discern modest, but still potentially relevant changes in transcriptional regulation bellow the pre-specified cut-off of gene expression. Fourth, to fully quantify the impact of ACF on intrinsic LV contractility, we would have to subject animals to simultaneous invasive pressure–volume analysis of the heart and its lack is a limitation of the study.

In conclusion, animals with HF due to ACF displayed several specific alterations of FA metabolism and fat distribution. Circulating FFA were higher but cardiac TG content was reduced in HF group, arguing against the role of lipid overload. The observation of intraabdominal fat depletion and attenuated postprandial insulin dynamics may help to understand mechanisms of HF-induced fat loss and cachexia.

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