Acylated ghrelin treatment normalizes skeletal muscle mitochondrial oxidative capacity and AKT phosphorylation in rat chronic heart failure

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Abstract

Background Chronic heart failure (CHF) is associated with skeletal muscle abnormalities contributing to exercise intolerance, muscle loss, and negative impact on patient prognosis. A primary role has been proposed for mitochondrial dysfunction, which may be induced by systemic and tissue inflammation and further contribute to low insulin signalling. The acylated form of the gastric hormone ghrelin (AG) may improve mitochondrial oxidative capacity and insulin signalling in both healthy and diseased rodent models.

Methods We investigated the impact of AG continuous subcutaneous administration (AG) by osmotic minipump (50 nmol/kg/day for 28 days) compared with placebo (P) on skeletal muscle mitochondrial enzyme activities, mitochondrial biogenesis regulators transcriptional expression and insulin signalling in a rodent post-myocardial infarction CHF model.

Results No statistically significant differences (NS) were observed among the three group in cumulative food intake. Compared with sham-operated, P had low mitochondrial enzyme activities, mitochondrial biogenesis regulators transcripts, and insulin signalling activation at AKT level ($P < 0.05$), associated with activating nuclear translocation of pro-inflammatory transcription factor nuclear factor-κB. AG completely normalized all alterations ($P < 0.05$ vs P, $P = \text{NS}$ vs sham-operated). Direct AG activities were strongly supported by in vitro C2C12 myotubes experiments showing AG-dependent stimulation of mitochondrial enzyme activities. No changes in mitochondrial parameters and insulin signalling were observed in the liver in any group.

Conclusions Sustained peripheral AG treatment with preserved food intake normalizes a CHF-induced tissue-specific cluster of skeletal muscle mitochondrial dysfunction, pro-inflammatory changes, and reduced insulin signalling. AG is therefore a potential treatment for CHF-associated muscle catabolic alterations, with potential positive impact on patient outcome.

Keywords Ghrelin; Skeletal muscle; Mitochondria; Insulin signalling

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Introduction

Incidence and prevalence of chronic heart failure (CHF) are increasing worldwide, and CHF-associated morbidity and mortality are a major health problem. CHF commonly causes skeletal muscle (SM) metabolic and functional alterations that lead to exercise intolerance and ultimately muscle loss with strong negative impact on patient outcome. Altered energy metabolism with low mitochondrial oxidative capacity has been proposed to play a primary role in CHF-
induced SM changes. Cardiac and systemic pro-inflammatory signalling activation has been conversely hypothesized to contribute to low tissue mitochondrial function, and skeletal muscle inflammation and mitochondrial dysfunction could further impair insulin signalling thereby collectively enhancing muscle catabolism. Effective therapeutic strategies to counteract CHF-induced SM mitochondrial dysfunction and its potential metabolic consequences are therefore urgently needed.

The gastric hormone acylated ghrelin (AG) has orexigenic effects and complex metabolic activities that may include stimulation of SM mitochondrial oxidative capacity and insulin signalling in both healthy and diseased rodent models. Despite potential negative effects on circulating glucose reported in human and experimental models due to potential glucogenic activities, AG has been therefore extensively studied and increasingly suggested as potential anti-catabolic treatment under chronic disease conditions associated with loss of body and muscle mass. We and others have shown beneficial effects of AG on body weight, muscle mass, and systemic inflammation in cancer and renal cachexia models. In experimental chronic kidney disease, we specifically demonstrated that sustained peripheral AG administration enhances skeletal muscle mitochondrial function also independently of food intake. No information is, however, available on potential effects of AG administration on skeletal muscle mitochondrial function and insulin signalling in CHF. In addition, no studies investigated potential AG effects on skeletal muscle inflammation as potential mediator of AG metabolic activities in cachexia-inducing diseases in vivo.

In the current study, we therefore investigated the impact of continuous 28 day AG subcutaneous administration on SM mitochondrial enzyme activities, transcriptional expression of mitochondrial biogenesis regulators as well as insulin signalling and pro-inflammatory pathways in a rodent post-myocardial infarction (IMA) CHF model. Direct AG effects on mitochondrial function and enzyme activities were further verified in vitro in C2C12 myotubes, whereas animals were studied in vivo at AG doses reported to prevent global alterations in food intake through the whole experimental period. Because AG was previously reported to reduce systemic inflammation markers in different chronic disease models and systemic inflammation may be activated in CHF with potential negative impact on SM inflammatory pathways and mitochondrial and metabolic dysfunction, we further examined whether activating nuclear translocation of the master pro-inflammatory transcription factor nuclear factor-κB (NFκB) is altered in CHF and normalized by AG. The same parameters were finally investigated in the liver to determine potential tissue distribution and specificity of AG activities because differential acylated ghrelin effects were reported at hepatic level with potential to negatively affect plasma metabolic profile.

Materials and methods

Experimental protocol and measurements

Animal surgery and treatments

All procedures were approved by the competent ethics committee (Ladesamt fur Gesundheit und Soziales Berlin, Germany; permit number G 0116/05). The study was designed to determine the effects of ghrelin or ghrelin analogues on body weight and composition in chronic heart failure, and part of the study results have been published elsewhere. All rats were treated with buprenorphine for 2 days following surgery for analgesia; 24 h mortality was 31%. Surviving animals were caged individually and fed standard chow under a 12 h light cycle; all rats were treated with furosemide starting 14 days after surgery. After two additional weeks, IMA animals were further randomized (n = 18 each) to receive either placebo (IMA-P) or human ghrelin at 50 nmol/kg/day (IPSEN Pharmaceuticals; IMA-AG) via osmotic minipumps implanted subcutaneously on the back (Charles River, Sulzfeld, Germany); control animals received placebo via the same route (n = 14). After 14 day infusions, minipumps were replaced on Day 42 from surgery; animals were then sacrificed at Day 56 from surgery after identical additional 14 day infusions. Animal body weight and food intake were recorded twice weekly throughout the study period; body weight changes were analysed and compared among groups during the first week of each infusion period, that is, at Days 28–36 and 42–50 after initial surgery. Echocardiography was performed prior to sacrifice, using an 15 MHz probe and an Acuson Sequoia system (Siemens, Erlangen, Germany); animals with infarct size less than 25% of left ventricle were excluded from analyses (n = 5).

Cytochrome c oxidase and citrate synthase activities and RNA analysis

Mitochondrial cytochrome c oxidase and citrate synthase enzyme activities were measured spectrophotometrically from muscle homogenates as referenced. Total RNA was isolated from 40–50 mg of gastrocnemius muscle (Tri Reagent, MRC, Cincinnati, OH, USA). Transcript levels of regulators of mitochondrial biogenesis and lipid metabolism were measured by real-time polymerase chain reaction (7900 Sequence Detection System, Applied Biosystems, Applied Biosystems Italia, Monza, Italy) as referenced. Briefly, 1 mg of total RNA was reverse-transcribed (RNA Reverse Transcription KIT, Applied Biosystems) and amplified using primers (300 nM) and probes (50 nM) selected using the Primer Express Software (Applied Biosystems). Primers for PGC-1α

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amplification were as follows: forward: AGCAAGCTCTGATGCTCTGAAGGA; reverse: ACCGAAGTGAGTGCTTATGCAGT. The probe sequence for PGC-1α detection was TGCGGAAGCCCTTTGGTGA. The reference gene used was 28S rRNA. Target and housekeeping genes were amplified separately, and their final quantitation was achieved using a relative standard curve. Results for each gene were divided by the corresponding 28S rRNA abundance and expressed as arbitrary units.

Nuclear factor-κB nuclear content, AKT insulin signalling, and plasma biochemical profile

Nuclear protein extracts for measurement of nuclear NFκB were obtained as previously reported. Protein concentration in all samples was measured by spectrophotometer (BCA Protein Assay Reagent, Pierce, Rockford, IL, USA). For measurement of tissue p65 NFκB subunit a commercially available kit was used following manufacturer instructions (TransAM, Active Motif North America, Carlsbad, CA, USA). Total (T) and phosphorylated (P: activated) AKT muscle protein contents were determined using western blotting as previously described. The P/T ratio was calculated as a marker of tissue AKT activation, and results were expressed as a percentage of sham value. Plasma insulin concentration was measured by enzyme-linked immunosorbent assay (Ultrasensitive ELISA; DRG, Spring. Plasma glucose and triglycerides levels were determined by standard enzymatic colorimetric assays.

C2C12 myoblast experiments

C2C12 myoblasts were differentiated in myotubes as previously reported. After 4 day incubation periods with differentiation medium and 18 h starvation, they were treated with AG (0.1 or 1 μmol/L) for 48 h, followed by collection and processing. Adenosine triphosphate (ATP) production from cell preparations was measured by luciferin-luciferase luminometric assay and cytochrome c oxidase and citrate synthase activities were measured spectrophotometrically as referenced.

Statistical analysis

All values are expressed as mean ± SEM. Comparison of parameters among the three groups were performed using one-way analysis of variance followed by the Tukey post-hoc test. P values <0.05 were considered statistically significant. Weight changes within each group were compared using paired t-test.

Results

Body weight, food intake, and plasma metabolic profile

No statistically significant differences in body weight among groups were observed at any time, and body weight was comparable at time of surgery in all groups despite a less pronounced body weight gain in the last ghrelin infusion period (Table 1). A general overall decline in food intake was observed for all groups between the first and the second ghrelin infusion period, but cumulative food intake remained comparable among the three groups during the whole infusion periods (Table 2). Plasma insulin, glucose, and triglycerides concentrations were comparable in all study groups (Table 3).

Skeletal muscle mitochondrial enzyme activities and mitochondrial biogenesis transcript levels (Figure 1)

Myocardial infarction with placebo had lower skeletal muscle activities of both cytochrome c oxidase and citrate synthase (P < 0.05 vs control; Figure 1). This alteration was associated with lower transcript levels of master regulators of mitochondrial biogenesis and lipid oxidation PGC-1a and PPAR-gamma (Figure 2). Ghrelin treatment was in turn associated with higher activities of both enzymes compared with IMA-P.

Statistical analysis

All values are expressed as mean ± SEM. Comparison of parameters among the three groups were performed using one-way analysis of variance followed by the Tukey post-hoc test. P values <0.05 were considered statistically significant. Weight changes within each group were compared using paired t-test.
Transcript levels of PGC-1a were also higher after AG treatment than in IMA-P animals \( (P < 0.05 \text{ vs } \text{IMA-P}, P = \text{NS vs control}) \) whereas no effects of AG were observed on PPAR\(\gamma\) expression (Figure 2).

Effects of continuous subcutaneous administration of gastric hormone acylated ghrelin (50 nmol/kg/day for 28 days) by osmotic minipump in a rodent post-myocardial infarction chronic heart failure model on final glucose, insulin, and triglyceride plasma levels. No statistical significant differences were observed among groups. Mean ± SEM, \( n = 14\text{–}18/\text{group} \). IMA-AG, myocardial infarction with gastric hormone acylated ghrelin; IMA-P, myocardial infarction with placebo

\( (P < 0.05 \text{ vs IMA-P}, P = \text{NS vs control}; \text{Figure 1}) \). Transcript levels of PGC-1a were also higher after AG treatment than in IMA-P animals \( (P < 0.05 \text{ vs IMA-P}, P = \text{NS vs control}) \) whereas no effects of AG were observed on PPAR\(\gamma\) expression (Figure 2).

**Table 3** Animal characteristics

|                   | Time   | Sham \( n = 14 \) | IMA-P \( n = 18 \) | IMA-AG \( n = 18 \) |
|-------------------|--------|-------------------|-------------------|---------------------|
| Plasma glucose (mg/dL) | T56    | 112 ± 5           | 107 ± 9           | 121 ± 6             |
| Plasma insulin (μU/mL)  | T56    | 12.3 ± 2.1        | 12.8 ± 0.9        | 11.7 ± 1.3          |
| Plasma triglycerides (mg/dL) | T56    | 0.73 ± 0.08       | 0.61 ± 0.04       | 0.72 ± 0.06         |

Skeletal muscle nuclear factor-κB nuclear content and AKT phosphorylation (Figure 2)

Higher skeletal muscle nuclear NFkB content and lower AKT phosphorylation were observed in IMA-P rats compared with control animals. Ghrelin treatment was in turn associated with normalized nuclear NFkB and AKT phosphorylation (both \( P < 0.05 \text{ for IMA-AG vs IMA-P}; P = \text{NS for IMA-AG vs control} \)).

Gastric hormone acylated ghrelin activities *in vitro* in C2C12 myotubes (Figure 3)

In order to determine potential direct effects of AG in skeletal muscle tissue, we further investigated its mitochondrial activities in C2C12 myotubes. Consistent with direct mitochondrial effects in skeletal muscle tissue, AG incubation resulted in enhanced mitochondrial ATP production as well as cytochrome c oxidase and citrate synthase activities \( (P < 0.05) \).

Liver mitochondrial enzyme activities and mitochondrial biogenesis transcript levels (Figure 4)

At variance with gastrocnemius muscle, liver activities of both cytochrome c oxidase and citrate synthase were not affected

Figure 1  Skeletal muscle mitochondrial enzyme activities and mitochondrial biogenesis transcript levels. Effects of continuous subcutaneous administration of gastric hormone acylated ghrelin (50 nmol/kg/day for 28 days) by osmotic minipump in a rodent post-myocardial infarction chronic heart failure model on citrate synthase (A) and cytochrome c oxidase (B) activity and on PGC1\(\alpha\) (C) and PPAR\(\gamma\) (D) gene expression in skeletal muscle. a.u., arbitrary units, \( *P < 0.05 \text{ versus other groups, mean ± SEM, } n = 14\text{–}18/\text{group} \). IMA-P, myocardial infarction with placebo; IMA-AG, myocardial infarction with gastric hormone acylated ghrelin.
by IMA or AG treatment and were comparable in the three study groups. Transcript levels of master regulators of mitochondrial biogenesis and lipid oxidation were similarly comparable in control, IMA-P, and IMA-AG animals. Further supporting tissue-specific CHF and AG activities, no changes were observed in any groups also for NFκB nuclear translocation and AKT phosphorylation.

Discussion

The current study demonstrated that (a) sustained peripheral acylated ghrelin treatment in a rodent model of IMA-induced chronic heart failure completely normalized a cluster of skeletal muscle mitochondrial dysfunction and tissue pro-inflammatory and insulin-desensitizing changes; (b) acylated ghrelin appears to act at least in part directly at skeletal muscle level because its mitochondrial effects are fully reproduced in vitro in muscle cells; (c) effects of chronic heart failure per se as well as those of acylated ghrelin treatment on energy metabolism, inflammation, and insulin signalling are tissue-specific and do not occur at hepatic level.

Mitochondrial dysfunction, inflammation, and insulin resistance have emerged in recent years as potentially causally related abnormalities with strong negative impact on skeletal muscle metabolism and function in chronic heart failure.3,10–15,17 Comprehensive normalization of CHF-induced alterations therefore indicates acylated ghrelin as a novel potential treatment for skeletal muscle dysfunction and its negative prognostic impact in CHF patients.4,6,7 The current findings are notably in excellent agreement with previous reports of mitochondrial effects of acylated ghrelin with enhanced skeletal muscle oxidative enzyme activities in healthy rodents19 and normalized enzyme activities in experimental chronic kidney disease.21 Anti-inflammatory and insulin-sensitizing AG activities were conversely also reported in healthy and obese rodent models20,27,36 and in vitro in response to pharmacological induction of muscle cell damage.25 Causal links and vicious cycles likely contribute to associations between muscle mitochondrial dysfunction, inflammation, and low insulin signalling in CHF. In particular, systemic pro-inflammatory changes are reported to occur in CHF, and they could lead to both mitochondrial and insulin signalling abnormalities in skeletal muscle.13,14 Mitochondrial dysfunction is, however, in turn also a potential primary cause of tissue
inflammation and insulin resistance through enhanced reactive oxygen species generation by damaged organelles. A primary role of improved mitochondrial function in global tissue AG activities in the current model is strongly supported by AG-induced in vitro enhancement of mitochondrial ATP production and oxidative capacity of both Krebs cycle and respiratory chain enzymes. In the current study, we notably also observed a CHF-induced transcriptional suppression of master mitochondrial biogenesis and oxidative metabolism regulators; PGC1-α but not PPARγ expression was selectively normalized by AG treatment, and this finding is in agreement with a direct involvement of mitochondrial biogenesis in the observed stimulation of mitochondrial parameters. Further studies should directly confirm this hypothesis.

In the current study, we investigated acylated ghrelin effects in the established IMA-induced CHF model, and all studied animals showed evidence of impaired left-ventricular function. Under the current conditions, all groups had comparable body weight and food intake at different experimental stages. Consistent with CHF-induced catabolism, P-treated animals, however, had less pronounced growth and total weight gain during the second part of the acylated ghrelin infusion treatment (Day 42–56 from surgery; not shown) despite comparable food intake. Less pronounced growth was conversely not observed in AG-treated animals, consistent with potential muscle mass preservation through the observed muscle anti-catabolic effects. Skeletal muscle weight was not available in the current study, and further experiments should directly aim at dissecting potential differential contributions of lean and fat tissues in acylated ghrelin anabolic activities.

Assessment of potential tissue-specificity of CHF-induced metabolic alterations as well as differential acylated ghrelin effects in different tissues was an additional aim of the current study. At variance with skeletal muscle, CHF had no relevant impact on hepatic mitochondrial, inflammatory, and insulin signalling parameters. This finding is consistent with available reports of clinically relevant hepatic abnormalities following acute heart failure and hemodynamic changes while chronic alterations in heart function may only result in liver metabolic alterations at late disease stages. Lack of AG activities in liver tissue is also consistent with tissue-specific metabolic effects of AG on skeletal muscle and liver, with potentially negative effects on hepatic insulin signalling and stimulation of hepatic glucogenic pathways in healthy lean rodents. Under the current experimental conditions, lack of changes in hepatic metabolic parameters was also associated with no negative impact of AG on plasma glucose,

Figure 3 Gastric hormone acylated ghrelin (AG) activities in vitro in C2C12 myotubes. Effects of the incubation with increasing concentrations of AG or unacylated ghrelin versus control (Ct) on citrate synthase (A) and cytochrome c oxidase (B) activity and adenosine triphosphate (ATP) synthesis rate (C) in C2C12 myotubes. PPKM, pyruvate + palmitoyl-L-carnitine + α-ketoglutarate + malate; PCM, palmitoyl-L-carnitine + malate; GM, glutamate + malate; SR, succinate + rotenone. *P < 0.05 versus Ctrl, †P < 0.05 versus AG 0.1 μM, mean ± SEM, n = 14–18/group.
insulin, and triglycerides. These findings are in agreement with lack of effects in chronic kidney disease models, thereby collectively supporting the concept that AG administration under various chronic disease conditions is metabolically safe and does not alter plasma biochemical profile as well as pivotal liver metabolic pathways.

In conclusion, we demonstrated that sustained peripheral AG treatment normalizes a CHF-induced cluster of skeletal muscle mitochondrial dysfunction, pro-inflammatory changes, and reduced insulin signalling. Acylated ghrelin activities at skeletal muscle level appear to be at least partly direct and independent because mitochondrial effects are fully reproduced in vitro in muscle cells. The metabolic impact of chronic heart failure per se and acylated ghrelin treatment are tissue-specific and do not occur at hepatic level. These findings collectively indicate AG as a potential novel treatment for CHF-associated muscle catabolic abnormalities, with potential positive impact on skeletal muscle function and patient outcome.

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The authors certify that they comply with the ethical guidelines for authorship and publishing of the Journal of Cachexia, Sarcopenia and Muscle.

Conflict of interest

None declared.

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