Acylated and unacylated ghrelin impair skeletal muscle atrophy in mice

Paolo E. Porporato,1 Nicoletta Filigheddu,1 Simone Reano,1 Michele Ferrara,1 Elia Angelino,1 Viola F. Gnocchi,1 Flavia Prodam,2 Giulia Ronchi,3 Sharmila Fagoonee,4 Michele Fornaro,3 Federica Chianale,1 Gianluca Baldanzi,1 Nicola Surico,1 Fabiola Singaglia,1 Isabelle Perroteau,5 Roy G. Smith,6 Yuxiang Sun,7 Stefano Geuna,3 and Andrea Graziani1

1Department of Translational Medicine, Interdisciplinary Research Center of Autoimmune Diseases (IRCAD), and Biotechnology Center for Applied Medical Research (BRMA), Università del Piemonte Orientale “Amedeo Avogadro” — Alessandria, Novara, Vercelli, Italy. 2Department of Health Sciences, Università del Piemonte Orientale “Amedeo Avogadro” — Alessandria, Novara, Vercelli, Italy. 3Neuroscience Institute “Cavaleri Ottolenghi” (NICO) and Department of Clinical and Biological Sciences, University of Torino, Orbassano (TO), Italy. 4Molecular Biotechnology Center and Department of Genetics, Biology and Biochemistry, and 5Department of Life Sciences and Systems Biology, University of Torino, Torino, Italy. 6Department of Metabolism and Aging, The Scripps Research Institute, Scripps, Florida, USA. 7USDA ARS Children’s Nutrition Research Center, Departments of Pediatrics and Molecular and Cellular Biology, Baylor College of Medicine, Houston, Texas, USA.

Cachexia is a wasting syndrome associated with cancer, AIDS, multiple sclerosis, and several other disease states. It is characterized by weight loss, fatigue, loss of appetite, and skeletal muscle atrophy and is associated with poor patient prognosis, making it an important treatment target. Ghrelin is a peptide hormone that stimulates growth hormone (GH) release and positive energy balance through binding to the receptor GHSR-1a. Only acylated ghrelin (AG), but not the unacylated form (UnAG), can bind GHSR-1a; however, UnAG and AG share several GHSR-1a–independent biological activities. Here we investigated whether UnAG and AG could protect against skeletal muscle atrophy in a GHSR-1a–independent manner. We found that both AG and UnAG inhibited dexamethasone-induced skeletal muscle atrophy and atrogene expression through PI3Kβ-, mTORC2-, and p38-mediated pathways in myotubes. Upregulation of circulating UnAG in mice impaired skeletal muscle atrophy induced by either fasting or denervation without stimulating muscle hypertrophy and GHSR-1a–mediated activation of the GH/IGF-1 axis. In Ghsr–deficient mice, both AG and UnAG induced phosphorylation of Akt in skeletal muscle and impaired fasting-induced atrophy. These results demonstrate that AG and UnAG act on a common, unidentified receptor to block skeletal muscle atrophy in a GH-independent manner.

Introduction
Skeletal muscle atrophy involves massive loss of muscle structural proteins, which leads to muscle weight decrease and progressive loss of muscle function. Skeletal muscle atrophy is induced by muscle denervation and disuse, and it is also the key component of cachexia, a catabolic, debilitating response to several diseases. Cachectic patients not only sustain a decreased quality of life, but also face a worse prognosis of the underlying pathology, making cachexia an important target for treatment (1). Ghrelin is a circulating peptide hormone, octanoylated on Ser3, that is mainly produced by the stomach, which, by acting on the hypothalamus and the pituitary, induces GH secretion and stimulates food intake and adiposity through binding to its receptor, GHSR-1a (2–5). In addition to its endocrine activities, ghrelin protects cardiac function after heart damage (6, 7). In vitro, ghrelin inhibits the apoptosis of cardiomyocytes and other cell types by activating PI3K/Akt and ERK-1/2 pathways (8–10). Acylated ghrelin (AG) and unacylated ghrelin (UnAG) are generated from the same precursor, which can be acylated by the specific intracellular ghrelin-O-acyltransferase G0AT (11, 12). UnAG, which is far more abundant in plasma than AG, does not bind to GHSR-1a, lacks any GH-releasing activity (13), and has been considered for many years to be the inactive product of ghrelin catabolism. However, UnAG shares with AG common high-affinity binding sites on several cell types lacking GHSR-1a, including myocardial and skeletal myocytes, where they stimulate survival and differentiation, respectively (8, 9, 14–16). Furthermore, UnAG regulates gene expression in fat, muscle, and liver independently of GHSR-1a (17).

In both human patients and experimental models, AG ameliorates cachexia induced by several pathological conditions (6, 7, 13, 18–21). Although AG may inhibit cachexia by stimulating food intake, positive energy balance, and release of GH and IGF-1, the mechanisms underlying its anticachectic activity have not been fully elucidated. Since we have previously shown that AG and UnAG, independently of GHSR-1a, inhibit apoptosis of cardiomyocytes by activating PI3K/Akt (8), a major antiatrophic signaling pathway (22, 23), and stimulate C2C12 skeletal myoblast differentiation (16), we investigated whether AG and UnAG could protect skeletal muscle from atrophy. Here, we provided evidence in vitro and in vivo that AG and UnAG, independently of GHSR-1a and activation of the GH/IGF-1 axis, trigger an antiatrophic signaling pathway by acting directly on the skeletal muscle, thereby protecting it from experimentally induced atrophy.

Results
AG and UnAG prevent dexamethasone-induced atrophy in C2C12-derived myotubes via mTORC2. C2C12 myotubes are a widely used model to study in vitro skeletal muscle atrophy induced by the synthetic glucocorticoid dexamethasone (24–26). Muscle atrophy was measured
AG and UnAG protect C2C12 myotubes from dexamethasone-induced atrophy without induction of protein synthesis or hypertrophy. (A) Myotube diameters were measured after 24-hour treatment in differentiation medium (DM) with 10 nM AG, 10 nM UnAG, and/or 1 μM dexamethasone (DEXA). In every experiment, 10 ng/ml IGF-1 was used as positive control for antiahropheic/hypertrophic activity. (B) and (C) Atrogin-1 and MuRF1 expression analysis upon dexamethasone treatment with or without AG and UnAG. (D) Treatment with 100 nM wortmannin (W) or 20 ng/ml rapamycin (R) reverted the antiahropheic activity of AG and UnAG on myotube diameter. Control myotubes in differentiation medium were treated with DMSO, a vehicle for both wortmannin and rapamycin. (E) and (F) Phosphorylation of Akt$^{S473}$ and FoxO3a$^{S256}$, detected by Western blotting, upon treatment for 20 minutes with 1 μM AG or UnAG. Shown are representative blots and quantification of 3 independent experiments. (G–I) IGF-1, but not AG and UnAG, induced protein synthesis, as determined by phosphorylation of S6K$^{T389}$ and S6$^{S235/236}$, a ribosomal protein whose phosphorylation mediates protein synthesis (29). AG and UnAG did not induce phosphorylation of S6K$^{T389}$ and S6$^{S235/236}$ (Figure 1, G and H), nor protein synthesis (as measured by $[^{3}H]$-leucine incorporation; Figure 1I) or myotube hypertrophy (Figure 1A). Conversely, IGF-1 induced S6K$^{T389}$ and S6$^{S235/236}$ phosphorylation, $[^{3}H]$-leucine incorporation, and myotube diameter increase, as expected.

By silencing raptor and rictor, specific components of mTORC1 and mTORC2, respectively (Figure 1), we observed that down-regulation of rictor abrogated the protective effect of both peptides on dexamethasone-induced muscle atrophy, measured as myotube diameter, while it did not affect the antiahropheic activity of IGF-1 (Figure 1K). Conversely, raptor silencing impaired IGF-1 antiahropheic activity without affecting that of AG/UnAG. These results indicate that mTORC2 pathway mediates AG/UnAG antiahropheic activity in C2C12 myotubes, without involving mTORC1-mediated protein synthesis.

To identify the signaling pathways differently activated by AG/UnAG and IGF-1, we investigated the role of p38 kinase, whose activation by AG/UnAG mediates C2C12 myoblast differentiation (16). In C2C12 myotubes, AG/UnAG, as well as IGF-1, induced phosphorylation of p38$^{Y180,T182}$ (Figure 2A), and its pharmacological inhibition impaired the antiahropheic activity of AG/UnAG, but not of IGF-1 (Figure 2B).

Activation of p38 has been reported to downregulate Atrogin-1, thereby contributing to the protection of skeletal muscle from atrophy (33). On the other hand, p38 mediates induction of Atrogin-1 by TNF-α and oxidative stress and of MuRF1 by serum starvation (34–37). Inhibition of p38 with SB203580 reduced dexamethasone-induced expression of both Atrogin-1 and MuRF1; nevertheless, induction of Atrogin-1, but not MuRF1, was still significant (Figure 2, C and D). In the presence of SB203580, AG and UnAG, but not IGF-1, failed to further reduce the residual induction of Atrogin-1, which indicates that p38 mediates AG/UnAG signaling in regulating Atrogin-1 expression.

To further characterize AG/UnAG antiahropheic activity, we treated C2C12 myotubes with NF449, a compound uncoupling Gα, from GPCRs, which inhibits antiapoptotic activity of AG and UnAG in pancreatic β cells (9, 38). NF449 completely abrogated Akt$^{Y473}$ phosphorylation and antiahropheic activity of AG/UnAG without affecting IGF-1 activities (Figure 2, E and F), which supports the hypothesis that AG and UnAG act through a GPCR, as previously suggested (9).

P38 $\alpha$ and $\beta$ isoforms mediate Akt activation upon stimulation of tyrosine kinase receptors and GPCRs, respectively (39, 40). We dissected the contribution of P13K$\alpha$ and P13K$\beta$ to IGF-1 and AG/UnAG antiahropheic activity using isoform-specific P13K inhibitors. Whereas inhibition of P13K$\alpha$ by PIK-75 abolished IGF-1 antiahropheic activity, it did not affect AG/UnAG protection. Conversely, inhibition of P13K$\beta$ by TGX-221 impaired AG/UnAG antiahropheic activity while not affecting IGF-1 protection (Figure 2G). The involvement of P13K$\beta$ in AG/UnAG antiahropheic activity was further supported by the finding that TGX-221 prevented AG/UnAG from reducing dexamethasone-induced Atrogin-1 expression (Figure 2H). Together, these data strongly suggest that AG/UnAG acts through GPCR-dependent signaling pathways involving a P13K isoform distinct from that of IGF-1.

Glucocorticoids induce muscle mass reduction by also upregulating the expression of myostatin, a TGF-β family member that acts as a negative regulator of muscle mass. Myostatin reduces the...
size of human skeletal muscle cell–derived myotubes by reducing mTOR/Akt/p70S6K signaling, while simultaneous treatment with IGF-1 restores myotube size, Akt phosphorylation, and protein synthesis (41, 42). In C2C12 myotubes, dexamethasone treatment actually induced the expression of myostatin, which was significantly reduced by IGF-1. However, AG/UnAG had no effect on myostatin expression (Supplemental Figure 1C), providing further evidence that ghrelin and IGF-1 inhibit muscle atrophy through distinct, partially overlapping, mechanisms.

**Figure 2**

AG and UnAG antiautrophic signaling is mediated by p38 and acts through a GPCR-dependent signaling pathway involving PI3Kβ. (A) Phosphorylation of p38[Y180/182], detected by Western blotting, after 20-minute treatment with 1 μM AG or UnAG. Shown are representative blots and quantification of 3 independent experiments. (B) Treatment with the p38 inhibitor SB203580 (5 μM) reverted the antiautrophic activity of AG and UnAG on myotube diameter upon treatment with dexamethasone. (C and D) Atrogin-1 and MuRF1 expression analysis upon dexamethasone treatment with or without AG and UnAG in the presence or absence of 5 μM SB203580. (E) AG and UnAG phosphorylation of AktS473 was abolished upon treatment with 10 μM NF449, a Gαs subunit–selective G protein antagonist. Shown are representative blots and quantification of 3 independent experiments. (F) Treatment with 10 μM NF449 reverted the antiautrophic activity of AG and UnAG on myotube diameter upon dexamethasone treatment. (G) Treatment with 25 nM PIK-75, an inhibitor of PI3Kα, abolished the antiautrophic effect of IGF-1 on myotube diameter upon dexamethasone treatment, without affecting AG and UnAG activity. The antiautrophic effect was abrogated by treatment with 200 nM TGX-221, an inhibitor of PI3Kβ. (H) Atrogin-1 expression analysis upon dexamethasone treatment with AG, UnAG, and IGF-1 in the presence or absence of 200 nM TGX-221. In experiments with SB203580, NF449, PIK-75, and TGX-221, control myotubes in differentiation medium were treated with DMSO, a vehicle for all these compounds. *P < 0.05, **P < 0.01 vs. DM control; *P < 0.05, **P < 0.01 vs. DEXA treatment.
mice did not feature any change in cir-
and adiposity, Myh6/Ghrl
inability of UnAG to activate GHSR-1a and to promote GH release
muscle (Supplemental Figure 2A). Moreover, consistent with the
Myh6/Ghrl
mice, without leakage in the skeletal
myocardium of
Ghrl
Tg mice (43–45).

- overexpressing
a 50-fold increase of circulating UnAG, without affecting AG lev-
ings were calculated as the mean of right and left hindlimbs. \( n = 7 \) per group (fed); \( 4 \) per group (fasted 48 hours); \( 5 \) per group (denervated). Data are mean ± SEM. *\( p < 0.05 \) vs. WT.

Table 1
Phenotypical characterization of Myh6/Ghrl mice

<table>
<thead>
<tr>
<th></th>
<th>WT</th>
<th>Myh6/Ghrl</th>
</tr>
</thead>
<tbody>
<tr>
<td>UnAG (pg/ml)</td>
<td>445.4 ± 155</td>
<td>25,000.5 ± 360(^a)</td>
</tr>
<tr>
<td>AG, fed (pg/ml)</td>
<td>41.7 ± 1.6</td>
<td>39.3 ± 1.5</td>
</tr>
<tr>
<td>AG, fasted (pg/ml)</td>
<td>75.7 ± 8.8</td>
<td>68.2 ± 9.5</td>
</tr>
<tr>
<td>IGF-1, fed (ng/ml)</td>
<td>748.5 ± 56</td>
<td>765.5 ± 120</td>
</tr>
<tr>
<td>IGF-1, fasted (ng/ml)</td>
<td>398 ± 93</td>
<td>328 ± 37</td>
</tr>
<tr>
<td>Insulin (pg/ml)</td>
<td>571 ± 58</td>
<td>631 ± 129</td>
</tr>
<tr>
<td>Tibial length (mm)</td>
<td>19.65 ± 0.11</td>
<td>19.62 ± 0.22</td>
</tr>
<tr>
<td>Nasoanal length (mm)</td>
<td>91.59 ± 0.51</td>
<td>90.61 ± 0.95</td>
</tr>
<tr>
<td>BMI, fed (g/cm²)</td>
<td>3.32 ± 0.12</td>
<td>3.33 ± 0.08</td>
</tr>
<tr>
<td>BMI, fasted (g/cm²)</td>
<td>2.93 ± 0.06</td>
<td>2.92 ± 0.09</td>
</tr>
<tr>
<td>Gastrocnemius weight, fed (mg)</td>
<td>134.86 ± 4.6</td>
<td>137.2 ± 5.62</td>
</tr>
<tr>
<td>Gastrocnemius weight, fasted (mg)</td>
<td>118 ± 4</td>
<td>124 ± 3.1</td>
</tr>
<tr>
<td>Gastrocnemius weight/tibial length, fed (mg/mm)</td>
<td>6.86 ± 0.22</td>
<td>6.99 ± 0.25</td>
</tr>
<tr>
<td>Gastrocnemius weight/tibial length, fasted (mg/mm)</td>
<td>5.89 ± 0.14</td>
<td>6.36 ± 0.15(^b)</td>
</tr>
<tr>
<td>Heart weight, fed (mg)</td>
<td>117.5 ± 11.8</td>
<td>122 ± 6.8</td>
</tr>
<tr>
<td>Heart weight, fasted (mg)</td>
<td>103.0 ± 7.1</td>
<td>105 ± 3.3</td>
</tr>
<tr>
<td>Heart weight/nasoanal length, fed (mg/mm)</td>
<td>1.22 ± 0.12</td>
<td>1.28 ± 0.08</td>
</tr>
<tr>
<td>Heart weight/nasoanal length, fasted (mg/mm)</td>
<td>1.16 ± 0.07</td>
<td>1.16 ± 0.03</td>
</tr>
<tr>
<td>Daily food intake (g)</td>
<td>4.66 ± 0.17</td>
<td>4.73 ± 0.07</td>
</tr>
<tr>
<td>Daily food intake, denervated (g)</td>
<td>4.60 ± 0.17</td>
<td>4.60 ± 0.18</td>
</tr>
</tbody>
</table>

To investigate whether UnAG might protect from muscle wasting, we induced skeletal muscle atrophy by food deprivation. After 48 hours of fasting, gastrocnemius weight was decreased by approximately 14% in WT mice, and by approximately 9% in Myh6/Ghrl mice, compared with fed animals (Figure 3A), which indicates that increased circulating UnAG results in 30% protection from fasting-induced loss in gastrocnemius weight. Accordingly, gastrocnemii CSA was reduced by 29% in WT mice and by 19% in Myh6/Ghrl mice compared with fed animals (Figure 3B), indicative of 34% protection. Similarly, extensor digitorum longus (EDL) muscle weight and mean fiber area of Myh6/Ghrl mice was reduced to a lesser extent than in WT animals (Figure 3, D and E). This protection was reflected by shift in CSA distributions of gastrocnemii and EDL — toward fibers with wider area — in Myh6/Ghrl compared with WT mice under fasting conditions (Figure 3, C and F).

After 48 hours of fasting, Atrogin-1 and MuRF1 expression in gastrocnemii of WT animals dramatically increased. In Myh6/Ghrl mice, the induction of Atrogin-1 was significantly reduced by one-third, while MuRF1 was only slightly, not significantly, decreased (Figure 3, G and H).

Plasma levels of glycerol and FFAs did not change in fasted Myh6/Ghrl and WT mice (Supplemental Figure 2, G and H), which indicates that fasting did not significantly affect either glycerol or FFA concentrations, consistent with previous reports in the FVB mouse background (48, 49). Moreover, hepatic phosphoenolpyruvate carboxykinase (PEPCK) expression was induced to the same extent in fasted Myh6/Ghrl and WT littermates (Supplemental Figure 2I). Together, these data suggest that muscle wasting–resistant properties of Myh6/Ghrl mice do not depend on effects of UnAG on energy balance.

Furthermore, Myh6/Ghrl mice were protected from denervation-induced muscle atrophy, an experimental procedure that does not affect animal daily food intake (Table 1). At 7 and 14 days after denervation, gastrocnemius weight of WT animals was reduced by 21% and 27%, respectively, while the loss of muscle weight in Myh6/Ghrl animals was significantly lower (Figure 4A). Consistently, gastrocnemii mean fiber CSA of WT animals was remarkably reduced at both 7 and 14 days after denervation, whereas CSA in Myh6/Ghrl animals was reduced to a lesser extent (Figure 4B). At 7 days after denervation, Myh6/Ghrl mice featured a mild shift of gastrocnemii CSA distribution that became impressive after 14 days (Figure 4, C and D). A strong inhibition of atrophy at 7 days after denervation was also evident in EDL (Figure 4, E and F) and tibialis anterior (TA) muscles (Figure 4, G and H).

Moreover, in gastrocnemii of Myh6/Ghrl mice, the induction of Atrogin-1 was reduced by 40% (Figure 4I). Conversely, MuRF1 was only slightly, not significantly, reduced (Figure 4J), consistent with the fasting-induced atrophy data. Together, these observations indicated that constitutive high levels of UnAG impair experimentally induced atrophy in vivo, likely through a mechanism independent of GHSR-1a and activation of the GH/IGF-1 axis.

UnAG pharmacological treatment induces antiaatrophic signaling in muscle and inhibits fasting- and denervation-induced atrophy. Acute

a 50-fold increase of circulating UnAG, without affecting AG lev-
elations (Table 1), as previously observed in other Ghr1-overexpressing Tg mice (43–45). Ghr1 mRNA overexpression was restricted to the myocardium of Myh6/Ghrl mice, without leakage in the skeletal muscle (Supplemental Figure 2A). Moreover, consistent with the inability of UnAG to activate GHSR-1a and to promote GH release and adiposity, Myh6/Ghrl mice did not feature any change in circulating IGF-1 concentration, tibial and nasoanal length, BMI, or food intake compared with their WT littermates. In addition, fasting decreased IGF-1 and increased ghrelin circulating concentrations to the same extent in WT and Myh6/Ghrl mice (Table 1). These data strongly indicate that the upregulation of circulating UnAG in Myh6/Ghrl mice does not activate GHSR-1a in the pituitary and hypothalamus, stimulate the GH/IGF-1 axis, or affect endogenous ghrelin regulation. Moreover, tissue expression of IGF-1, which in skeletal muscle may act locally in a paracrine/autocrine manner (46), was not altered in Myh6/Ghrl mice, either in fed or in fasted animals (Supplemental Figure 2B).

Although AG and UnAG differently regulate insulin release and sensitivity (47), basal insulin level, glucose uptake, and insulin sensiti-

Notably, compared with WT animals, fed Myh6/Ghrl mice did not feature any difference in heart and gastrocnemius muscle weight (Table 1), fiber cross-sectional area (CSA) distribution, or hindlimb force (as measured by grasping test; Supplemental Figure 2, E and F), which indicates that high levels of circulating UnAG do not induce skeletal muscle hypertrophy in vivo, consistent with the inability of UnAG to induce hypertrophy in C2C12-derived myotubes.

- dependent of GHSR-1a and activation of the GH/IGF-1 axis.

- indicated that constitutive high levels of UnAG impair experimen-
tally induced atrophy in vivo, likely through a mechanism inde-
pendent of GHSR-1a and activation of the GH/IGF-1 axis.

UnAG pharmacological treatment induces antiaatrophic signaling in muscle and inhibits fasting- and denervation-induced atrophy. Acute
administration of exogenous UnAG at 100 µg/kg, a dose previously used for in vivo studies (6), induced phosphorylation of AktS473, FoxO3aT32, and p38/ERK1/2 in WT gastrocnemii (Figure 5, A–C), which indicates that, in vivo, UnAG activates the same antiaatrophy signaling pathway as it does in C2C12 myotubes.

Repeated administration (every 12 hours) of UnAG protected mice from skeletal muscle atrophy induced by either fasting or denervation (Figure 5, D–I). UnAG treatment preserved gastrocnemii from weight and mean fiber CSA loss (Figure 5, D and E). Accordingly, frequency distribution of gastrocnemii CSA of fasted mice injected with UnAG showed a dramatic shift toward bigger fiber areas compared with saline-injected mice (Figure 5F).

Similarly, UnAG treatment of denervated mice resulted in a 25% protection from gastrocnemii muscle weight loss and a significantly lower decrease of mean fiber CSA, although the CSA distribution of UnAG-injected mice showed only a very mild shift compared with saline-injected animals (Figure 5, G–I). Although the plasma concentration of UnAG after injection dropped to basal levels in about 2–4 hours (Supplemental Figure 3A), these data indicate that repeated acute stimulation is sufficient to protect from experimentally induced skeletal muscle atrophy without affecting muscular IGF-1 expression (Supplemental Figure 3B).

AG and UnAG induce antiaatrophy signaling and impair muscle atrophy in Ghsr–/– mice. The findings reported above, along with previous data on common binding sites for AG/UnAG in C2C12 lacking Ghsr (16), strongly suggest that AG and UnAG stimulate antiaatrophy signaling in skeletal muscle through activation of a receptor distinct from GHSR-1a. To verify this hypothesis, we assayed AG/UnAG antiaatrophy signaling and activity in Ghsr–/– mice, in which AG fails to activate the GH/IGF-1 axis or stimulate appetite (50). Injection of either AG or UnAG induced AktS473 phosphorylation in gastrocnemii of Ghsr–/– mice (Figure 6A). Consistently, treatment of Ghsr–/– mice with 100 µg/kg AG or UnAG twice daily reduced gastrocnemii weight loss induced by 48-hour fasting by 30% compared with saline-treated animals (Figure 6B). Moreover, the mean CSA loss of AG- and UnAG-injected mice strongly decreased compared with saline-injected animals, and CSA distribution shifted toward bigger areas (Figure 6, C and D).

In summary, these findings demonstrated that both AG and UnAG activate a direct antiaatrophy signaling pathway in skeletal muscle and protect from experimentally induced muscle atrophy, independently of the AG receptor GHSR-1a.

**Discussion**

Several studies have shown that AG protects from cachexia and prevents muscle proteolysis in vivo, supposedly through stimulation of appetite and activation of the GH/IGF-1 axis mediated by AG binding to GHSR-1a (6, 7, 18–21). However, here we provided in vitro and in vivo evidence that AG and UnAG exert antiaatrophy activity by acting directly on the skeletal muscle, even in Ghsr–/– mice.

Upregulation of circulating UnAG, which does not bind GHSR-1a and does not activate the GH/IGF-1 axis, counteracted muscle atrophy induced by either fasting or denervation. Consistently, UnAG has been reported to reduce burn-induced skeletal muscle proteolysis and local TNF-α upregulation (51).

We achieved upregulation of circulating UnAG either by myocardial Ghrl overexpression in Myhb/Ghrl mice or by repeated administration. The antiaatrophy activity of UnAG cannot be mediated by its conversion to AG in the plasma, since acylation occurs only intracellularly on the ghrelin precursor by the ghrelin-specific acyltransferase GOAT (11, 12). The negligible myocardial expression of GOAT might explain the increase of only the unacylated form.
of circulating ghrelin in Myh6/Ghrl mice. This is in agreement with other tissue-specific Ghrl Tg mice featuring high UnAG circulating levels in the absence of significant changes of AG (43–45).

The observations that Myh6/Ghrl mice did not feature any change in circulating and muscular IGF-1 or in tibial or whole body length, along with the lack of skeletal muscle hypertrophy, further indicate that the GH/IGF-1 axis is not activated in these mice. Finally, the finding that both AG and UnAG impaired skeletal muscle atrophy in Ghsr–/– mice indicated that their antiatrophic activity is mediated by a receptor distinct from GHSR-1α. In these mice, AG exerted antiatrophic activity in the skeletal muscle independent of its role in modulating GH release and energy balance. Nevertheless, these data do not exclude the possibility that in WT animals, GHSR-1α may contribute to the antiatrophic activity of AG by also regulating the GH/IGF-1 axis and positive energy balance. For instance, AG has been suggested to prevent downregulation of muscular IGF-1 expression in an experimental model of cachexia through an indirect mechanism involving GHSR-1α activity on positive energy balance (20).

The hypothesis that AG/UnAG impairs muscle atrophy in vivo by acting directly on the skeletal muscle is further supported by our finding that UnAG administration rapidly stimulated anti-atrophic signaling in the gastrocnemius. Moreover, AG/UnAG activated antiatrophic signaling in cultures of C2C12 myotubes, which do not express GHSR-1α, protecting them from dexamethasone-induced atrophy and atrogene upregulation. Although AG has previously been reported to fail in reducing dexamethasone-induced Atrogin-1 expression in C2C12 myotubes (20), the 10-fold lower dexamethasone concentration used in that study and the considerably weaker Atrogin-1 induction may explain the different results. Conversely, Sheriff et al. showed that UnAG reduces TNF-α/IFN-γ-induced cachexia in C2C12 myotubes in a PI3K/mTOR-dependent manner (51). The results of our present study not only confirmed the involvement of PI3K/mTOR pathways in AG/UnAG activity on skeletal muscle, but also showed the specific contribution of the mTORC2 over the mTORC1-mediated signaling pathway, which may explain, at least in part, the ability of AG/UnAG to protect from skeletal muscle atrophy without a concomitant induction of hypertrophy.

Indeed, the molecular mechanisms underlying AG/UnAG anti-atrophic activity in the skeletal muscle involved the activation of mTORC2-mediated signaling pathways, leading to phosphorylation of AktS473 and of its substrate FoxO3aT32, which eventually impaired Atrogin-1 expression and muscle protein degradation. At

Figure 4
Myh6/Ghrl mice are protected from denervation-induced skeletal muscle atrophy induced by sciatic nerve resection. (A and B) Mean percentage of weight loss (A) and CSA reduction (B) of denervated gastrocnemius at 7 and 14 days after denervation, compared with the unperturbed side. (C and D) Frequency distribution of gastrocnemius CSA at 7 and 14 days after denervation in Myh6/Ghrl and WT mice. (E–H) CSA reduction and fiber area distribution of (E and F) EDL and (G and H) TA muscles at 7 days after denervation. (I and J) Atrogin-1 and MuRF1 expression, determined by real-time RT-PCR, in denervated gastrocnemius at 7 days after denervation, compared with the unperturbed side. **P < 0.01, *P < 0.05 vs. WT. n = 6 (WT); 5 (Myh6/Ghrl); 3 (CSA loss and distribution, WT and Myh6/Ghrl).
the same time, in C2C12 myotubes, AG/UnAG failed to stimulate mTORC1-mediated phosphorylation of S6K\(^{T389}\) and S6\(^{235/236}\), protein synthesis, and hypertrophy. Consistently, chronic upregulation of circulating UnAG in \(\text{Myh6/Ghrl}\) mice did not induce muscle hypertrophy. This finding highlights a remarkable difference between the antiatrophic activities of AG/UnAG and IGF-1 in the skeletal muscle, as IGF-1 stimulates both mTORC2-mediated impairment of protein degradation and mTORC1-dependent stimulation of protein synthesis and hypertrophy (23–26). Consistently, in TNF-\(\alpha\)/IFN-\(\gamma\)–treated C2C12 myotubes, UnAG inhibited protein catabolism and impaired the induction of Atrogin-1 and MuRF1. Moreover, UnAG restored the basal phosphorylation state of proteins of mTORC1 and mTORC2 pathways, although the lack of UnAG-induced increase in Akt\(^{S473}\) phosphorylation observed herein may depend on receptor desensitization, given the higher concentration of UnAG used and the protracted treatment (51).

The finding that downregulation of the mTORC1-specific component raptor did not affect the antiatrophic activity of AG/UnAG, while impairing IGF-1 antiatrophic activity, further supports the conclusion that AG/UnAG antiatrophic activity does not involve mTORC1-mediated stimulation of protein synthesis. On the other hand, the finding that AG/UnAG antiatrophic activity was sensitive to downregulation of rictor, the specific component of mTORC2, demonstrated the key role of mTORC2 in mediating AG/UnAG antiatrophic activity. The finding that ghrelin-induced phosphorylation of Akt\(^{S473}\) was uncoupled from the activation of mTORC1-mediated pathways and hypertrophy may appear controversial, as IGF-1–induced phosphorylation of Akt\(^{S473}\) is associated with the activity of both mTOR complexes (29), and overexpression of constitutive active Akt in the skeletal muscle prevents denervation-induced atrophy and induces hypertrophy (22, 52). The lack of muscle hypertrophy observed in \(\text{Myh6/Ghrl}\) mice may depend on weaker stimulation of the PI3K/Akt pathway by UnAG. Indeed, although tissue-specific expression of constitutive active Akt in Tg mice induces strong phosphorylation of Akt and of its substrates (53), phosphorylation of Akt was not detectable in

Figure 5
UnAG pharmacological treatment protects skeletal muscle from fasting- and denervation-induced atrophy in WT mice. (A–C) Phosphorylation of Akt\(^{S473}\), FoxO3a\(^{T32}\), and p38\(^{T180/Y182}\) in gastrocnemii of WT mice treated with 100 \(\mu\)g/kg UnAG or saline. At the indicated time points, gastrocnemii were removed and processed for Western blot analysis. Shown are representative blots and densitometric analysis of 3 independent experiments, normalized to untreated animals (not shown). (D–F) Mean percent weight loss (D), CSA reduction (E), and CSA frequency distribution (F) of gastrocnemii from fed or 48-hour fasted mice treated twice daily with 100 \(\mu\)g/kg UnAG or saline (\(n = 5\) per group). Frequency distribution was measured in 3 mice per group. In D and E, percent reduction shown is between fasted and fed mice. (G–I) Mean percent weight loss (G), CSA reduction (H), and CSA frequency distribution (I) of gastrocnemii from mice treated with 100 \(\mu\)g/kg UnAG or saline twice daily for 7 days after sciatic nerve resection (\(n = 5\) per group). Frequency distribution was measured in 3 mice per group. In G and H, percent reduction shown is between denervated gastrocnemii and gastrocnemii from the unperturbed side. *\(P < 0.05\), **\(P < 0.01\) vs. saline treatment.
molecules of Myh6/Ghr1 mice (data not shown). Indeed, we found that 2 distinct PI3K isoforms, namely PI3Kβ and PI3Kα, mediated the antiatrophic activity of AG/UnAG and IGF-1, respectively. This observation, along with the ability of a Gαi2-uncoupling drug to abolish the antiatrophic activity of AG/UnAG, but not IGF-1, is consistent with the hypothesis that the unknown receptor mediating the common activities of AG/UnAG is a GPCR (9). Moreover, these data further serve to rule out the hypothesis that AG/UnAG acts on myotubes by stimulating the autocrine release of IGF-1.

The inability of AG and UnAG to stimulate protein synthesis and hypertrophy in the skeletal muscle is consistent with their key role in the adaptive response to fasting and negative energy balance (13). The molecular mechanisms underlying the uncoupling of mTORC2 from mTORC1 remain to be investigated. AG and UnAG, which are released during fasting, might shift muscle metabolism toward amino acid oxidation, thereby decreasing the intracellular pool of free amino acids essential for mTORC1 activity (29). Alternatively, activation of PI3Kα, whose enzymatic activity is lower than that of PI3Kβ (54), may result in weaker activation of Akt. Finally, AMPK, which negatively regulates mTORC1 in skeletal muscle (55), may contribute to mTORC1 uncoupling, although AG was reported to be unable to stimulate AMPK in rat gastrocnemius (56).

The finding that p38 was required for AG/UnAG antiatrophic activity is consistent with previous findings that p38 cooperates with PI3K/Akt pathways to induce C2C12 differentiation (16, 57). However, the role of p38 in regulating muscle atrophy is complex, as its activation mediates muscle atrophy induced by oxidative stress and inflammatory cytokines (34, 36, 58). The role of p38 in signaling is determined by its association in distinct signaling complexes with different regulators and substrates and by its localization (35). Our findings are consistent with evidence indicating that, in myotubes, decreased p38 phosphorylation is associated with dexamethasone-induced atrophy, and that p38 mediates β-hydroxy-β-methylbutyrate protection from dexamethasone-induced protein degradation (59, 60). Moreover, p38 activity can regulate cytoplasmic localization of FoxO3a independently of Akt, thereby impairing its transcriptional activity and Atrogin-1 induction (33, 61). Furthermore, activation of p38 stabilizes and activates the transcriptional coactivator PGC1α, which represses FoxO3a activity (62, 63). Although IGF-1 activated p38, this was dispensable for IGF-1 antiatrophic activity. In addition, IGF-1 and AG/UnAG antiatrophic activities differed in the inability of AG/UnAG to downregulate myostatin, a TGF-β-like inhibitor of muscle growth, which further supports the hypothesis that AG/UnAG and IGF-1 counteract muscle atrophy through distinct molecular mechanisms.

The data presented herein unveiled a novel component of the complex role of AG/UnAG, i.e., the direct activation of antiaatrophic pathways in the skeletal muscle, eventually leading to reduced muscle wasting. This effect adds to the well-known capabilities of AG to stimulate appetite, regulate lipid metabolism, and release GH. Although the identity of the novel AG/UnAG receptor is yet unknown, these findings may have important biological and therapeutic implications, since they provide proof that UnAG has a strong and specific potential for the prevention or treatment of muscle atrophy, avoiding the diabetogenic side effects of AG (47) and the cancer risk associated with IGF-1 treatment (64).

**Methods**

*Reagents.* AG1–28 and UnAG1–28 were purchased from PolyPeptide Laboratories. The PI3K p110α inhibitor PIK-75 hydrochloride was purchased from Axon Medchem, and the PI3K p110β inhibitor TGX-221 was a gift from U. Galli (Synthetic Medicinal Chemistry group, Università del Piemonte Orientale, Novara, Italy). Water-soluble dexamethasone and all other reagents, unless otherwise stated, were from Sigma-Aldrich. Anti-phospho-AktS473, anti-Akt, anti–phospho-FoxO3aT32, anti-FoxO3a, anti–phospho-S6KT389, anti-S6K, anti–phospho-S6S235/236, anti-S6, anti-p38T180/Y182, anti-p38, anti-raptor, and anti-rictor antibodies were from Cell Signaling Technology; anti-actin antibody was from Santa Cruz Biotechnology. AG and UnAG and IGF-1 were purchased from Axon Medchem.

*Cell cultures and myotube analysis.* C2C12 myoblasts were differentiated in myotubes as previously described (16). For measurement of myotube diameters, mice of Myh6/Ghr1 mice (data not shown). Indeed, we found that 2 distinct PI3K isoforms, namely PI3Kβ and PI3Kα, mediated the antiatrophic activity of AG/UnAG and IGF-1, respectively. This observation, along with the ability of a Gαi2-uncoupling drug to abolish the antiatrophic activity of AG/UnAG, but not IGF-1, is consistent with the hypothesis that the unknown receptor mediating the common activities of AG/UnAG is a GPCR (9). Moreover, these data further serve to rule out the hypothesis that AG/UnAG acts on myotubes by stimulating the autocrine release of IGF-1.

The inability of AG and UnAG to stimulate protein synthesis and hypertrophy in the skeletal muscle is consistent with their key role in the adaptive response to fasting and negative energy balance (13). The molecular mechanisms underlying the uncoupling of mTORC2 from mTORC1 remain to be investigated. AG and UnAG, which are released during fasting, might shift muscle metabolism toward amino acid oxidation, thereby decreasing the intracellular pool of free amino acids essential for mTORC1 activity (29). Alternatively, activation of PI3Kβ, whose enzymatic activity is lower than that of PI3Kα (54), may result in weaker activation of Akt. Finally, AMPK, which negatively regulates mTORC1 in skeletal muscle (55), may contribute to mTORC1 uncoupling, although AG was reported to be unable to stimulate AMPK in rat gastrocnemius (56).

The finding that p38 was required for AG/UnAG antiatrophic activity is consistent with previous findings that p38 cooperates with PI3K/Akt pathways to induce C2C12 differentiation (16, 57). However, the role of p38 in regulating muscle atrophy is complex, as its activation mediates muscle atrophy induced by oxidative stress and inflammatory cytokines (34, 36, 58). The role of p38 in signaling is determined by its association in distinct signaling complexes with different regulators and substrates and by its localization (35). Our findings are consistent with evidence indicating that, in myotubes, decreased p38 phosphorylation is associated with dexamethasone-induced atrophy, and that p38 mediates β-hydroxy-β-methylbutyrate protection from dexamethasone-induced protein degradation (59, 60). Moreover, p38 activity can regulate cytoplasmic localization of FoxO3a independently of Akt, thereby impairing its transcriptional activity and Atrogin-1 induction (33, 61). Furthermore, activation of p38 stabilizes and activates the transcriptional coactivator PGC1α, which represses FoxO3a activity (62, 63). Although IGF-1 activated p38, this was dispensable for IGF-1 antiatrophic activity. In addition, IGF-1 and AG/UnAG antiatrophic activities differed in the inability of AG/UnAG to downregulate myostatin, a TGF-β-like inhibitor of muscle growth, which further supports the hypothesis that AG/UnAG and IGF-1 counteract muscle atrophy through distinct molecular mechanisms.

The data presented herein unveiled a novel component of the complex role of AG/UnAG, i.e., the direct activation of antiaatrophic pathways in the skeletal muscle, eventually leading to reduced muscle wasting. This effect adds to the well-known capabilities of AG to stimulate appetite, regulate lipid metabolism, and release GH. Although the identity of the novel AG/UnAG receptor is yet unknown, these findings may have important biological and therapeutic implications, since they provide proof that UnAG has a strong and specific potential for the prevention or treatment of muscle atrophy, avoiding the diabetogenic side effects of AG (47) and the cancer risk associated with IGF-1 treatment (64).
eters, myotubes were fixed, and diameters were quantified by measuring a total of >100 myobase diameters from 5 random fields in 3 replicates at ×40 magnification using Image-Pro Plus software (Media Cybernetics) as described previously (24).

Raptor and rictor silencing. Raptor siRNA (MISSION pre-designed siRNA SASI_Mm01_00055293; Sigma-Aldrich), rictor siRNA (SASI_Mm01_00137731; Sigma-Aldrich), Block-iT, or siRNA negative control sequence (Invitrogen) were transfected with Lipofectamine2000 (Invitrogen) in C2C12 myotubes. Transfection efficiency was evaluated by the fluorescent siRNA negative control Block-iT, and silencing was verified by Western blot.

[^H]-leucine incorporation assay. C2C12 myotubes were maintained for 24 hours with or without 10 nM AG or UnAG in differentiation medium supplemented with 2 μCi/ml[^H]-leucine (Perkin Elmer) to evaluate the induction of protein synthesis. At the end of treatments, cells were washed with PBS, treated with 5% trichloroacetic acid, and lysed with 0.5 M NaOH and 0.5% SDS. The amount of incorporated[^H]-leucine was evaluated with PBS, treated with 5% trichloroacetic acid, and lysed with 0.5 M NaOH and 0.5% SDS. The amount of incorporated[^H]-leucine was evaluated by β counter (Tri-Carb 2800TR; Perkin Elmer) analysis. Data are the average of 4 replicates.

Western blot. C2C12 myotubes were serum starved overnight and then treated as indicated in the figure legends. Western blot was performed as previously described (16). Unless otherwise specified, after use of anti–phospho-specific antibodies, membranes were stripped with Re-Blot Plus (Chemicon, Millipore) and rebolted with the corresponding total protein antibodies.

Muscles of mice fasted for 6 hours were s.c. injected with 100 μg/kg UnAG or AG or with saline solution. At the indicated time points, gastrocnemii were removed, homogenized at 4°C in RIPA buffer (1% Triton X-100; 1% sodium deoxycholate; 0.1% SDS; 1 mM EDTA; 1 mM EGTA; 50 mM NaF; 160 mM NaCl; and 20 mM Tris-HCl, pH 7.4) containing 1 mM DTT, protease inhibitor cocktail, and 1 mM Na3VO4. Homogenates were then processed as above.

TG animal generation and treatment. All experiments were conducted on young adult male FVB1 WT, FVB1 Myh6/Ghrl, and C57BL/6J Ghr−/− mice (50), matched for age and weight. TG animals were obtained by cloning the murine ghrelin gene (Ghrl) under control of the cardiac promoter sequences of the β myosin heavy chain 3′ UTR and the first 3 exons of the α isoform Myh6 (65). Transgene integration and expression were confirmed by PCR and real-time RT-PCR, respectively. Phenotypical characterization and experiments were carried out on hemizygote animals and littermate controls.

AG, UnAG, and IGF-1 plasmatic levels were measured by EIA kits (SPIbio Bertin Pharma for AG and UnAG; R&D Systems for IGF-1); insulin plasmatic levels were quantified with the Insulin (mouse) ELISA kit (ALPCO Diagnostics); and glycerol and free fatty acid plasmatic levels were evaluated by enzymatic assay kits (Cayman).

BMI was calculated as animal weight divided by the square of the nasoanalog length.

Fasting-induced atrophy was achieved by 48 hours of food removal (63), while denervation-induced muscle atrophy was obtained by resection of the sciatic nerve under anesthesia with sevoflurane (Baxter) and evaluated 7 and 14 days later (66). Muscles were collected, weighed, and normalized for tibial length and processed either for RNA extraction or for histology.

Skeletal muscle force was assessed using the BS-GRIP (2Biological Instruments) as previously described (68). Each animal was tested 3 times, and the average value of the maximum weight that the animal managed to hold was recorded and normalized to the mouse’s weight.

Statistics. Data are presented as mean ± SEM. Variation among groups was evaluated using nonparametric Wilcoxon and Mann-Whitney U tests. Statistical significance was assumed for P values less than 0.05. All statistical analyses were performed with SPSS for Windows version 17.0.

Study approval. All animal experimental procedures were approved by the Institutional Animal Care and Use Committee at Università del Piemonte Orientale "Amedeo Avogadro."

Acknowledgments

We are grateful to Riccarda Granata and Cristina Grande for insulin measurements and to Thi-en Thi Nguyen, Christian Zurlo, Laura Badà, and Giulia Bettas Ardisson for technical assistance. This work was supported by Telethon (grant no. GGP030386 to A. Graziani), Regione Piemonte CIPE (to A. Graziani, S. Geuna, and I. Perroteau), Regione Piemonte Ricerca Sanitaria (to A. Graziani), Italian Ministry for University and Research (PRIN grant to A. Graziani, S. Geuna, and I. Perroteau), and Opera Pia Eletto Lualdi.

Received for publication December 23, 2011, and accepted in revised form November 1, 2012.

Address correspondence to: Nicoletta Filigheddu, Department of Translational Medicine, Università del Piemonte Orientale “Amedeo Avogadro,” Via Solaroli 17, 28100 Novara, Italy. Phone: 39.032160529; Fax: 39.0321620421; E-mail: nicoletta.filigheddu@med.unipmn.it.

Paolo E. Porporato’s present address is: Unit of Pharmacology and Therapeutics, Université Catholique de Louvain, Brussels, Belgium.

Viola F. Gnocchi’s present address is: Research Center for Genetic Medicine, Children’s National Medical Center, Washington, DC, USA.

Federica Chianale’s present address is: Oncological Sciences Department, Systems Biology Unit, IRCC, Candiolo (TO), Italy.

The Journal of Clinical Investigation  Volume 123  Number 2  February 2013


